## High Performance Fortran Language Specification

High Performance Fortran Forum

January 31, 1997 Version 2.0 The High Performance Fortran Forum (HPFF), with participation from over 40 organizations, met from March 1992 to March 1993 to define a set of extensions to Fortran called High Performance Fortran (HPF). Our goal was to address the problems of writing data parallel programs for architectures where the distribution of data impacts performance. While we hope that the HPF extensions will become widely available, HPFF is not sanctioned or supported by any official standards organization. The HPFF had a second series of meetings from April 1994 to October 1994 to consider requests for corrections, clarifications, and interpretations to the Version 1.0 HPF document and also to develop user requirements for possible future changes to HPF. A third set of meetings took place From January 1995 through December 1996 to incorporate features recommended to meet user needs identified in the 1994 meetings.

This document contains all the technical features proposed for the version of the language known as HPF Version 2.0. This copy of the draft was processed by IATEX on January 31, 1997.

HPFF encourages requests for interpretation of this document, and comments on the language defined here. We will give our best effort to answering interpretation questions, and general comments will be considered in future HPFF language specifications.

Please send interpretation requests to hpff-interpret@cs.rice.edu. Your request is archived and forwarded to a group of HPFF committee members who attempt to respond to it.

The text of interpretation requests becomes the property of Rice University.

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The High Performance Fortran Forum (HPFF) is a coalition of industrial and academic groups working to suggest a set of standard extensions to Fortran that provide support for high performance programming on a wide variety of machines, including massively parallel SIMD and MIMD systems and vector processors. From its beginning, HPFF has included most vendors delivering parallel machines, a number of government laboratories, and university research groups. Public input has been encouraged. This document defining HPF 2.0 is the third in a series of documents resulting from the HPFF. HPF 2.0 is intended to be a language portable from workstations to massively parallel supercomputers while being able to express the algorithms needed to achieve high performance on specific architectures. HPF 2.0 builds on the efforts of the previous HPFF meetings, primarily in 1992 and 1994. Specific acknowledgments for the many people who contributed to the previous versions of HPF are included in Annex D.

## HPFF 2 Acknowledgments

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- Chuck Koelbel, Editor, assisted by multiple committee members (names later).

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# Part I

# Introduction

This major section describes the organization of the document as a whole. It also defines terms and concepts that are common to High Performance Fortran version 2.0 (described in Part II) and the HPF Approved Extensions (described in Part III). Therefore, it provides necessary background for the succeeding sections.

## Section 1

## Overview

This document specifies the form and establishes the interpretation of programs expressed in the High Performance Fortran (HPF) language. It is designed as a set of extensions and modifications to the established International Standard for Fortran. At the time of publication of this document, the version of the standard used as a base is informally referred to as "Fortran 95" (ISO/IEC 1539:1997). References to that document are made as follows: Section 13.11.6 in that document is referred to here as F95:13.11.6.

In this overview Section of the document, we outline the goals and scope of the language, introduce the HPF language model, highlight the main features of the language, describe the changes between HPF 1.1 and HPF 2.0, and provide a guide to the rest of this document.

### 

## 1.1 Goals and Scope of High Performance Fortran

The primary goals behind the development of the HPF language include:

- Support for data parallel programming (single threaded, global name space, and loosely synchronous parallel computation);
- Portability across different architectures;
- High performance on parallel computers with non-uniform memory access costs (while not impeding performance on other machines);
- Use of Standard Fortran (currently Fortran 95) as a base;
  - Open interfaces and interoperability with other languages (e.g., C) and other programming paradigms (e.g., message passing using MPI).
- Secondary goals include:
- Implementation feasibility within a limited time span;
- Provision of input to future standards activities for Fortran and C;
- Provision of an evolutionary path for adding advanced features to the language in a consistent manner.

The first version of the language definition, HPF 1.0 was released in May 1993. A number of language features that were defined in HPF 1.0 have now been absorbed into

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the Fortran 95 language standard (e.g., the FORALL statement and construct, and PURE procedures). These features are therefore no longer detailed in the definition of HPF 2.0. Information about the evolution of the HPF language (through versions 1.0, 1.1, and 2.0) and an enumeration of the differences between HPF 2.0 from HPF 1.1 may be found in subsection 1.4.

## 1.2 HPF Language Model

An important goal of HPF is to achieve code portability across a variety of parallel machines. This requires not only that HPF programs compile on all target machines, but also that a highly-efficient HPF program on one parallel machine be able to achieve reasonably high efficiency on another parallel machine with a comparable number of processors. Otherwise, the effort spent by a programmer to achieve high performance on one machine would be wasted when the HPF code is ported to another machine. Although shared-memory machines and distributed-memory machines may use different low-level primitives, there is broad similarity with respect to the fundamental factors that affect the performance of parallel programs on these machines. Thus, achieving high efficiency across different parallel machines with the same high level HPF program is a feasible goal. Some of the fundamental factors affecting the performance of a parallel program are the degree of available parallelism, exploitation of data locality, and choice of appropriate task granularity. HPF provides mechanisms for the programmer to guide the compiler with respect to these factors.

The first versions of HPF were defined to extend Fortran 90. HPF 2.0 is defined as an extension to the current Fortran Standard (Fortran 95). Future revisions of HPF will include and be consistent with advances in the Fortran standards, as they are approved by ISO.

Building on Fortran, HPF language features fall into four categories:

- HPF directives;
- New language syntax;
- New library routines; and
- Language changes and restrictions.

HPF directives appear as structured comments that suggest implementation strategies or assert facts about a program to the compiler. When properly used, they affect only the efficiency of the computation performed, but do not change the value computed by the program. The form of the HPF directives has been chosen so that a future Fortran standard may choose to include these features as full statements in the language by deleting the initial comment header.

A few new language features have been defined as direct extensions to Fortran syntax 42 and interpretation. The new HPF language features differ from HPF directives in that they 43 are first-class language constructs and can directly affect the result computed by a program. 44

The HPF library of computational functions defines a standard interface to routines that have proven valuable for high performance computing. These additional functions include those for mapping inquiry, bit manipulation, array reduction, array combining scatter, prefix and suffix, and sorting.

A small number of changes and restrictions to Fortran 95 have also been defined. The most significant restrictions are those imposed on the use of sequence and storage association, since they are not compatible with the data distribution features of HPF. It is however possible to retain sequence and storage association semantics in a program by use of certain explicit HPF directives.

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## 1.2.1 Data Mapping Directives

<sup>9</sup> The fundamental model of parallelism in HPF is that of single-threaded data-parallel ex-<sup>10</sup> ecution with a globally shared address space. Fortran array statements and the FORALL <sup>11</sup> statement are natural ways of specifying data parallel computation. In addition, HPF pro-<sup>12</sup> vides the INDEPENDENT directive. It can be used to assert that certain loops do not carry <sup>13</sup> any dependences and therefore may be executed in parallel.

Exploitation of data locality is critical to achieving good performance on a high-14 performance computer, whether a uniprocessor workstation, a network of workstations, 15or a parallel computer. On a Non-Uniform-Memory-Access (NUMA) parallel computer, 16 the effective distribution of data among processor memories is very important in reducing 17data movement overheads. One of the key features of HPF is the facility for user specifica-18 tion of data mapping. HPF provides a logical view of the parallel machine as a rectilinear 19 arrangement of abstract processors in one or more dimensions. The programmer can specify 20 the relative alignment of elements of different program arrays, and the distribution of arrays 21 over the logical processor grid. Data mapping is specified using HPF directives that can 22 aid the compiler in optimizing parallel performance, but have no effect on the semantics of 23 the program. This is illustrated by the following simple example. 24

25

```
REAL A(1000,1000)
26
       !HPF$ PROCESSORS procs(4,4)
27
       !HPF$ DISTRIBUTE (BLOCK, BLOCK) ONTO procs :: A
28
             DO k = 1, num_iter
29
                 FORALL (i=2:999, j=2:999)
30
                   A(i,j) = (A(i,j-1) + A(i-1,j) + A(i,j+1) + A(i+1,j))/4
31
                 END FORALL
32
             END DO
33
```

34 35

The code fragment describes a simple Jacobi relaxation computation using a twodimensional floating-point array A. The HPF directives appear as structured comments. The PROCESSORS directive specifies a logical  $4 \times 4$  grid of processors proc. The DISTRIBUTE directive recommends that the compiler partition the array A into equal-sized blocks along each of its dimensions. This will result in a  $4 \times 4$  configuration of blocks each containing  $250 \times 250$  elements, one block per processor. The PROCESSORS and DISTRIBUTE directive are described in detail later in Section 3.

The outer DO k loop iterates over num\_iter Jacobi relaxation steps. The inner loop uses the Fortran 95 FORALL construct. It specifies the execution of the loop body for all values of i and j in the range 2 through 999. The semantics of the FORALL require that the right-hand-side expressions for all iterations (i.e., for all values of i and j between 2 and 999) be evaluated before any of the assignments to the left-hand-side variables are performed.

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When targeted for execution on a distributed-memory machine with 16 processors, 1 an HPF compiler generates SPMD code, with each processor locally containing a part 2 of the global array A. The outer  $\mathbf{k}$  loop is executed sequentially while the inner FORALL 3 is executed in parallel. Each processor will require some "boundary" elements of A that 4 reside in partitions mapped to the local memories of other processors. Primitives to achieve 5 the necessary inter-processor communication are inserted by the HPF compiler into the 6 generated SPMD code. The single-threaded data-parallel model with a global name-space 7makes it convenient for the programmer to specify the strategy for parallelization and data 8 partitioning at a higher level of abstraction. The tedious low-level details of translating 9 from an abstract global name space to the local memories of individual processors and the 10 management of explicit inter-processor communication are left to the compiler. 11

The following example illustrates some of the communication implications of scalar assignment statements. The purpose is to illustrate the implications of data distribution specifications on communication requirements for parallel execution. The explanations given do not necessarily reflect the actual compilation process.

Consider the following code fragment:

```
REAL a(1000), b(1000), c(1000), x(500), y(0:501)
      INTEGER inx(1000)
!HPF$ PROCESSORS procs(10)
!HPF$ DISTRIBUTE (BLOCK) ONTO procs :: a, b, inx
!HPF$ DISTRIBUTE (CYCLIC) ONTO procs :: c
!HPF$ ALIGN x(i) WITH y(i+1)
      . . .
      a(i) = b(i)
                                      ! Assignment 1
      x(i) = y(i+1)
                                      ! Assignment 2
      a(i) = c(i)
                                      ! Assignment 3
      a(i) = a(i-1) + a(i) + a(i+1)
                                      ! Assignment 4
      c(i) = c(i-1) + c(i) + c(i+1)
                                      ! Assignment 5
      x(i) = y(i)
                                      ! Assignment 6
      a(i) = a(inx(i)) + b(inx(i))
                                      ! Assignment 7
```

In this example, the PROCESSORS directive specifies a linear arrangement of 10 pro-34 cessors. The DISTRIBUTE directives recommend to the compiler that the arrays a, b, and 35 inx should be distributed among the 10 processors with blocks of 100 contiguous elements 36 per processor. The array c is to be cyclically distributed among the processors with c(1), 37  $c(11), \ldots, c(991)$  mapped onto processor  $procs(1); c(2), c(12), \ldots, c(992)$  mapped 38 onto processor procs(2); and so on. The complete mapping of arrays x and y onto the 39 processors is not specified, but their relative alignment is indicated by the ALIGN directive. 40 The ALIGN statement recommends that x(i) and y(i+1) be stored on the same processor 41for all values of i, regardless of the actual distribution chosen by the compiler for y(y(0))42 and y(1) are not aligned with any element of x). The PROCESSORS, DISTRIBUTE, and ALIGN 43 directives are discussed in detail in Section 3. 44

In Assignment 1 (a(i) = b(i)), the identical distribution of a and b specifies that for all i, corresponding elements of a(i) and b(i) should be mapped to the same processor. Therefore, execution of this statement requires no communication of data values between processors. 48 In Assignment 2 (x(i) = y(i+1)), there is no inherent communication. In this case, the relative alignment of the two arrays matches the assignment statement for any actual distribution of the arrays.

<sup>4</sup> Although Assignment 3 (a(i) = c(i)) looks very similar to the first assignment, the <sup>5</sup> communication requirements are very different due to the different distributions of a and c. <sup>6</sup> Array elements a(i) and c(i) are mapped to the same processor for only 10% of the possible <sup>7</sup> values of i. (This can be seen from the definitions of BLOCK and CYCLIC in Section 3.) The <sup>8</sup> elements are located on the same processor if and only if  $\lfloor (i - 1)/100 \rfloor = (i - 1) \mod 10$ . <sup>9</sup> For example, the assignment involves no inherent communication (i.e., both a(i) and c(i)<sup>10</sup> are on the same processor) if i = 1 or i = 102, but does require communication if i = 2.

In Assignment 4 (a(i) = a(i-1) + a(i) + a(i+1)), the references to array a are all on the same processor for about 98% of the possible values of i. The exceptions to this are i = 100 \* k for any k = 1, 2, ..., 9, (when a(i) and a(i-1) are on procs(k) and a(i+1) is on procs(k+1)) and i = 100 \* k + 1 for any k = 1, 2, ..., 9 (when a(i) and a(i+1) are on procs(k+1) and a(i-1) is on procs(k)). This statement requires communication. only for "boundary" elements on each processor,

Assignment 5, c(i) = c(i-1) + c(i) + c(i+1), while superficially similar to Assignment 4, has very different communication behavior. Because the distribution of c is CYCLIC rather than BLOCK, the three references c(i), c(i-1), and c(i+1) are mapped to three distinct processors for any value of i. Therefore, this statement requires communication for at least two of the right-hand side references, regardless of the implementation strategy.

The final two assignments have very limited information regarding the communica-23 tion requirements. In Assignment 6  $(\mathbf{x}(\mathbf{i}) = \mathbf{y}(\mathbf{i}))$  the only information available is that 24  $\mathbf{x}(\mathbf{i})$  and  $\mathbf{y}(\mathbf{i+1})$  are on the same processor; this has no logical consequences for the re-25lationship between  $\mathbf{x}(\mathbf{i})$  and  $\mathbf{y}(\mathbf{i})$ . Thus, nothing can be said regarding communication 26 required at runtime for the statement without further information. In Assignment 7 (a(i) 27 = a(inx(i)) + b(inx(i)), it can be proved that a(inx(i)) and b(inx(i)) are always 28 mapped to the same processor. Similarly, it is easy to deduce that **a(i)** and **inx(i)** are 29 mapped together. Without knowledge of the values stored in inx, however, the relation be-30 tween a(i) and a(inx(i)) is unknown, as is the relationship between a(i) and b(inx(i)). 31

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## 1.3 Overview of HPF 2.0 Language Features

The language defined in this document consists of two main parts:

• The HPF 2.0 Language (Part II)

• HPF 2.0 Approved Extensions (Part III)

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The HPF 2.0 language includes features that are expected to be implementable within a year of release of the language specification. These include basic data distribution features, data parallel features, intrinsic and library routines, and the extrinsic mechanism. The Approved Extensions include advanced features that meet specific needs, but are not likely to be supported in initial compiler implementations.

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## 1.3.1 HPF 2.0 Language Features

## 1.3.1.1 Data Distribution Features (Sections 3 and 4)

Most parallel and sequential architectures attain their highest speed when the data accessed exhibits locality of reference. The sequential storage order implied by Fortran standards often conflicts with the locality demanded by the architecture. To avoid this, HPF includes features that describe the co-location of data (ALIGN) and the partitioning of data among memory regions or abstract processors (DISTRIBUTE). Compilers may interpret these annotations to improve storage allocation for data, subject to the constraint that semantically every data object has a single value at any point in the program. Section 4 defines how the mapping features interact across subprogram boundaries.

While a goal of HPF is to maintain compatibility with Fortran, full support of Fortran sequence and storage association, however, is not compatible with the goal of high performance through distribution of data in HPF. Sections 3 and 4 describe restrictions and directives related to storage and sequence association.

### **1.3.1.2** Data Parallel Execution Features (Section 5)

To express parallel computation explicitly, HPF defines the INDEPENDENT directive. It asserts that the statements in a particular section of code do not exhibit any sequentializing dependences; when properly used, it does not change the semantics of the construct, but may provide more information to the language processor to allow optimizations. A REDUCTION clause can be used with the INDEPENDENT directive to identify variables that are updated by commutative and associative operations. This facilitates the utilization of parallelism with reduction operations, in the context of loops where the order of accumulation of updates to a variable is insignificant.

## **1.3.1.3** Extrinsic Program Units (Section 6)

Because HPF is designed as a high-level machine-independent language, there are certain operations that are difficult or impossible to express directly. For example, an application may benefit from finely-tuned systolic communications on certain machines; HPF's global address space does not express this well. HPF defines the Extrinsic mechanism to facilitate interfacing with procedures written in other paradigms, such as explicit message-passing subroutine libraries or in other languages, such as C.

### **1.3.1.4** Intrinsic Functions and Standard Library (Section 7)

Experience with massively parallel machines has identified many basic operations that are useful in parallel algorithm design. The Fortran array intrinsics address some of these. HPF adds several classes of parallel operations to the language definition as intrinsic functions and as standard library functions. In addition, several system inquiry functions useful for controlling parallel execution are provided in HPF.

#### HPF 2.0 Approved Extensions 1.3.21

#### 1.3.2.1Extensions for Data Mapping (Section 8)

4 The extended mapping features permit greater control over the mapping of data, includ-5ing facilities for dynamic realignment and redistribution of arrays at run-time (REALIGN, 6 **REDISTRIBUTE**, **DYNAMIC** directives), mapping of data among subsets of processors, mapping of pointers and components of derived types, and support for irregular distribution of 7data (GEN\_BLOCK and INDIRECT distributions). In addition, mechanisms are defined that 8 9 permit the programmer to provide information to the compiler about the range of possible distributions an array might take (RANGE directive) and the amount of buffering to be used 10  $1\,1$ with arrays involved in stencil-based nearest-neighbor computations (SHADOW).

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## 1.3.2.2

## Extensions for Data and Task Parallelism (Section 9)

The ON directive facilitates explicit computation partitioning. The site of recommended 15execution of a computation can be specified either as an explicitly identified subset of a 16processor arrangement, or indirectly as the set of processors onto which a data object or 17template is mapped. 18

In order to assist the compiler in generating efficient code, the **RESIDENT** directive is 19 defined, to be used in conjunction with an ON directive by the programmer. It can be used 20 to assert that all accesses to the specified object within the scope of the ON directive are to 21 be found locally on the executing processor. The TASK\_REGION directive allows the user to 22 specify the concurrent execution of different blocks of code on disjoint processor subsets. 23

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#### Extensions for Asynchronous I/O (Section 10) 1.3.2.3

26 In order to permit overlap of I/O with computation, an extension has been defined for 27 asynchronous READ/WRITE of direct, unformatted data. This is done through an additional 28 I/O control parameter in the Fortran READ/WRITE statement that specifies non-blocking 29 execution and a new statement (WAIT). 30

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#### Extensions to Intrinsic and Library Procedures (Section 12) 1.3.2.4

33 The approved extensions to the HPF intrinsics and library routines relate mostly to mapping 34 inquiry procedures. Some new inquiry routines are defined and other routines defined by 35 the HPF 2.0 language are extended to facilitate inquiry about extended mapping features, 36 such as mapping to processor subsets, GEN\_BLOCK, INDIRECT and DYNAMIC distributions. A 37 generalization of the Fortran TRANSPOSE intrinsic is also defined.

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#### Approved Extensions for HPF Extrinsics (Section 11) 1.3.2.5

A number of specific extrinsic interfaces are defined in Section 11 as approved HPF 2.0 4142 extensions. These include interfaces to facilitate interoperability with other languages (e.g., C and FORTRAN 77) as well as interfaces for different models of parallelism (LOCAL for 43 SPMD parallel, and SERIAL for single-process sequential). Library routines useful in the 44 extrinsic models are defined in Section 11.7. Additional extrinsic interfaces that are formally 45recognized by the HPF Forum, but not defined and maintained by the Forum, are included 46 in Annexes F and G. The policy and mechanism for formal recognition of such extrinsic 47interfaces is described in Annex E. 48

#### 1.4Changes from HPF 1.1 1 2 HPF 2.0 differs from HPF 1.1 in a number of ways: з 4 **Repartitioning of the Language:** The new document describes two components: the 5HPF 2.0 language (which is expected to be widely and relatively rapidly implemented) 6 and the set of Approved Extensions (which are not part of HPF 2.0 but may be 7included in future implementations in response to user demand, as the compilation 8 technology matures.) 9 Features Now in Standard Fortran: Fortran, instead of Fortran 90 is now defined as 10 the base language for extensions; this implies that HPF includes all features added to 11 Fortran at the 1995 revision. With this revision, a few HPF 1.1 features are now part 12 of the Fortran standard, and hence no longer appear as HPF extensions to Fortran. 13 14 Features Removed or Restricted in HPF 2.0: Some features of HPF 1.1, that have 15not been implemented to date, have been removed from the language because expe-16rience has shown that the simplicity gained by doing so outweight the advantage of 17the features. 18 Elimination of the HPF Subset: Unlike HPF 1.1, HPF 2.0 no longer has a recom-19 mended minimal subset for faster implementation (i.e. Subset HPF), although the 20 original HPF 1.1 Subset is documented in an annex. $^{21}$ 22 Features Moved to Approved Extensions: A few language features have been moved 23 from HPF 1.1 to the category of Approved Extensions. $^{24}$ $^{25}$ **New Features of HPF 2.0:** A few new features have been added to the base language. 26 New Approved Extensions: A number of further new features are defined as approved 27 extensions to the language. 28 29 **Recognized Externally-Supported HPF Extrinsics:** Finally, the document acknowl-30 edges a new category, HPF-related EXTRINSIC interfaces, that are recognized as meet-31 ing appropriate standards for such interfaces, but are not included as Approved Ex-32 tensions. Responsibility for the content of each such interface is assumed by the 33 organization proposing it rather than by the HPF Forum. 34 Each of these categories is summarized in the following subsections. 35 36 **Repartitioning of the Language** 1.4.137 38 The HPF Forum had two important goals that were sometimes in conflict: 39 • Providing advanced language capabilities that users had requested. 40 41 • Allowing fast compiler development by vendors. 42 One compromise made to satisfy both goals was to divide the language definition into two 43 parts. HPF 2.0 is very similar to HPF 1.1, and is expected to be efficiently implemented by 44 a number of vendors within approximately one year from the appearance of this document. 45Advanced features that require more implementation effort are collected as Approved Ex-46 tensions. Implementors are encouraged to support these features as rapidly as possible, and 47users are encouraged to speed this process by making their wishes known to the vendors.

1.4.2	Features Now in Standard Fortran
	llowing features, which formed part of HPF 1.1, have been removed from the docu- ecause they are now part of ISO Fortran:
• 1	'he FORALL statement and construct;
• 1	'he PURE attribute for procedures;
• E	Extensions to the MINLOC and MAXLOC intrinsics to include an optional DIM argument.
1.4.3	Features Removed or Restricted in HPF 2.0
The fo	lowing features have been removed from the language:
• S	equential arrays may no longer be explicitly mapped;
	n any procedure call in which distributed data may require redistribution, the pro- edure must now have an explicit interface;
	'he treatment of the INHERIT directive has been simplified in that it is no longer ossible to specify both INHERIT and DISTRIBUTE together.
• 1	'he treatment of pointers has been simplified.
1.4.4	Features Moved to Approved Extensions
	<b>'NAMIC</b> attribute and the <b>REDISTRIBUTE</b> and <b>REALIGN</b> statements have been moved Approved Extensions.
1.4.5	New Features of HPF 2.0
The fo	lowing new constructs have been introduced in HPF 2.0:
• 1	he REDUCTION clause for INDEPENDENT loops;
• 1	he new HPF_LIBRARY procedures SORT_DOWN, SORT_UP.
1.4.6	New Approved Extensions
The A <sub>j</sub>	pproved Extensions include the following features not part of HPF 1.1:
• 1	Apping of objects to processor subsets;
• E	Explicit mapping of pointers and components of derived types;
• 1	lew distribution formats: GEN_BLOCK and INDIRECT;
	lew directives: RANGE, SHADOW, ON, RESIDENT, TASK_REGION;
• A	Additional intrinsic procedures: ACTIVE_NUM_PROCS, ACTIVE_PROCS_SHAPE, and a gen- ralized TRANSPOSE intrinsic;
	New HPF_LIBRARY procedures: HPF_MAP_ARRAY and HPF_NUMBER_MAPPED; revision of rocedures HPF_ALIGNMENT, HPF_DISTRIBUTION and HPF_TEMPLATE;

• Support for asynchronous I/O with a new statement WAIT, and an additional I/O control parameter in the Fortran READ/WRITE statement;	1 2
• Extensions to the EXTRINSIC facilities to support interoperability with C and FOR- TRAN 77.	3 4 5
1.4.7 Recognized Externally-Supported HPF Extrinsics	6 7
Two externally supported extrinsic interfaces are recognized in this document:	8
• HPF_CRAFT: providing an SPMD paradigm with HPF features;	9 10
<ul> <li>The Fortran 77 Local Library: defining library support for calling Fortran 77 proce- dures in local mode.</li> </ul>	11 12 13
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## Section 2

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# Notation and Syntax

This chapter describes the notational conventions employed in this document and the syntax of HPF directives.

## 2.1 Notation

<sup>19</sup> This document uses the same notation as the Fortran 95 standard. In particular, the same <sup>20</sup> conventions are used for syntax rules. BNF descriptions of language features are given in <sup>21</sup> the style used in the Fortran standard. To distinguish HPF syntax rules from Fortran rules, <sup>22</sup> each HPF rule has an identifying number of the form Hsnn, where s corresponds to the <sup>23</sup> section number and nn is a two-digit sequence number. Nonterminals not defined in this <sup>24</sup> document are defined in the Fortran standard. Also note that certain technical terms such <sup>25</sup> as "storage unit" are defined by the Fortran standard.

As previously noted in Section 1, a reference of the form F95:2.4.7 in the text refers to
 Section 2.4.7 of the Fortran 95 standard.

Part III describes the approved extensions. In some cases this requires extending the syntax rules already introduced in an earlier section. In particular, the syntax rules here are often supersets of similar syntax rules in Part II; in these cases, the names of the nonterminals include the suffix *-extended*. Thus, when a non-terminal such as *name* is redefined it is referred to as *name-extended* under the proviso that any reference to *name* is to be replaced by *name-extended* in the rest of the syntax rules.

When a constraint or restriction in Part II is modified by an approved extension, this fact is noted, in the text, and a forward reference is provided. A downward-pointing double arrow is used in the margin (as here) to highlight such a forward reference.

<sup>37</sup> Each such modification (in Part III) contains a backward reference to the original <sup>38</sup> language in Part II that is modified. An upward-pointing double arrow is used in the <sup>39</sup> margin (as here) to highlight such a backward reference.

Rationale. Throughout this document, material explaining the rationale for including features, for choosing particular feature definitions, and for making other decisions, is set off in this format. Readers interested only in the language definition may wish to skip these sections, while readers interested in language design may want to read them more carefully. (End of rationale.)

Advice to users. Throughout this document, material that is primarily of interest to
 users (including most examples of syntax and interpretation) is set off in this format.

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Readers interested only in technical material may wish to skip these sections, while readers wanting a more tutorial approach may want to read them more carefully. (*End of advice to users.*)

Advice to implementors. Throughout this document, material that is primarily of interest to implementors is set off in this format. Readers interested only in the language definition may wish to skip these sections, while readers interested in compiler implementation may want to read them more carefully. (End of advice to implementors.)

## 2.2 Syntax of Directives

HPF directives are consistent with Fortran syntax in the following sense: if any HPF directive were to be adopted as part of a future Fortran standard, the only change necessary to convert an HPF program would be to replace the directive-origin with blanks.

H201	hpf-d	irective-line	$\mathbf{is}$	$directive$ - $origin\ hpf$ - $directive$	17		
					18		
H202	direc	ctive-origin	is	!HPF\$	19		
		0	or	CHPF\$	20		
			or	*HPF\$	21		
H203 hpf-directive		is	specification- $directive$	22			
11200	s npj-airective			executable-directive	23		
			or	executuole-arrective	24		
H204	speci	$\it fication$ - $\it directive$	$\mathbf{is}$	processors- $directive$	25		
			or	align-directive	26		
			or	distribute- $directive$	27		
			or	inherit- $directive$	28		
			or	template- $directive$	29		
			or	$combined\mspace{-}directive$	30		
			or	sequence- $directive$	31		
H205	execu	<i>stable-directive</i>	$\mathbf{is}$	independent- $directive$	32		
				-	33		
Const	raint:	An <i>hpf-directive-line</i> can the same line.	nnot	be commentary following another statement on	34 35		
a .	•				36		
Const	raint:	1 0	may	appear only where a <i>declaration-construct</i> may	37		
		appear.			38		
Const	raint:	An <i>executable-directive</i>	may appear only where an <i>executable-construct</i>				
Const	r arri e .	appear.	maj	appear only where an excession construct may	40		
		appear.					
Const	raint:	10		the rules of either Fortran free form $(F95:3.3.1.1)$	42		
			· ·	mment lines, depending on the source form of the	43		
		surrounding Fortran sou	rce fo	orm in that program unit. $(F95:3.3)$	44		
	1 0				45		
A	n hpf-	<i>directive</i> is case insensitiv	ve an	d conforms to the rules for blanks in free source	46		

An *hpf-directive* is case insensitive and conforms to the rules for blanks in free source form (3.3.1), even in an HPF program otherwise in fixed source form. However an HPF- 47 conforming language processor is not required to diagnose extra or missing blanks in an HPF 48

## 2.2. SYNTAX OF DIRECTIVES

directive. Note that, due to Fortran rules, the *directive-origin* in free source form must be 1 the characters !HPF\$. HPF directives may be continued, in which case each continued line 2 з also begins with a *directive-origin*. No statements may be interspersed within a continued HPF-directive. HPF directive lines must not appear within a continued statement. HPF 4 directive lines may include trailing commentary. 5 The blanks in the adjacent keywords END FORALL and NO SEQUENCE are optional, in 6 either source form. 7 An example of an HPF directive continuation in free source form is: 8 9 10 !HPF\$ ALIGN ANTIDISESTABLISHMENTARIANISM(I,J,K) & !HPF\$ 11WITH ORNITHORHYNCHUS\_ANATINUS(J,K,I) 1213 An example of an HPF directive continuation in fixed source form follows. Observe 14that column 6 must be blank, except when signifying continuation. 1516!HPF\$ ALIGN ANTIDISESTABLISHMENTARIANISM(I,J,K) 17!HPF\$\*WITH ORNITHORHYNCHUS\_ANATINUS(J,K,I) 18 19 This example shows an HPF directive continuation that is "universal" in that it can 20 be treated as either fixed source form or free source form. Note that the "&" in the first 21line is in column 73. 22 23 !HPF\$ ALIGN ANTIDISESTABLISHMENTARIANISM(I,J,K) & 24 !HPF\$&WITH ORNITHORHYNCHUS\_ANATINUS(J,K,I) 2526 Part III introduces new directives, both specifications and executable ones, for the 27approved extensions to HPF 2.0. These are included below: 28 29 H206 specification-directive-extended is processors-directive 30 **or** subset-directive 31 **or** align-directive 32 **or** *distribute-directive* 33 **or** *inherit-directive* 34 **or** *template-directive* 35 or combined-directive 36 **or** sequence-directive 37 **or** dynamic-directive 38 or range-directive 39 **or** shadow-directive 40 41 H207 executable-directive-extended *independent-directive* 42 is **or** realign-directive 43 **or** redistribute-directive 44 on-directive or 45 **or** resident-directive 46 47 The following rule extends rule R215 of Fortran 95: 48

H208	$executable\-construct\-extended$	is or	action-stmt case-construct	1 2
		or	do-construct	2
		or	if-construct	4
		or	where-construct	5
		or	on-construct	6
		or	resident-construct	7
		or	task-region-construct	8
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# Part II

# High Performance Fortran Language

This major section describes the syntax and semantics of features of the High Performance Fortran language, version 2.0. Some technical terms used herein are defined in Part I; otherwise this description is self-contained. Part III builds upon this material.

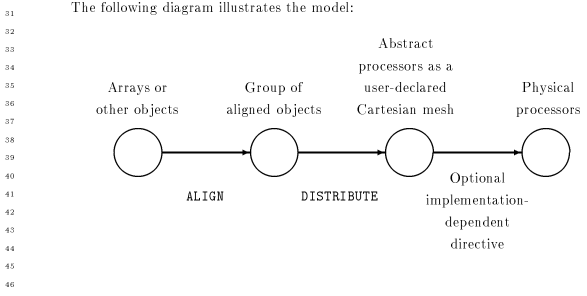
## Section 3

# Data Mapping

HPF data alignment and distribution directives allow the programmer to advise the compiler how to assign array elements to processor memories. This section discusses the basic data mapping features applicable, particularly those that are meaningful within a single scoping unit. Section 4 discusses features that apply when mapped variables appear as procedure arguments.

## 3.1 Model

HPF adds directives to Fortran to allow the user to advise the compiler on the allocation of data objects to processor memories. The model is that there is a two-level mapping of data objects to memory regions, referred to as "abstract processors." Data objects  $^{25}$ (typically array elements) are first *aligned* relative to one another; this group of arrays is then distributed onto a rectilinear arrangement of abstract processors. (The implementation then uses the same number, or perhaps some smaller number, of physical processors to implement these abstract processors. This mapping of abstract processors to physical processors is implementation-dependent.) 



The underlying assumptions are that an operation on two or more data objects is likely to be carried out much faster if they all reside in the same processor, and that it may

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be possible to carry out many such operations concurrently if they can be performed on different processors.

Fortran provides a number of features, notably array syntax, that make it easy for a compiler to determine that many operations may be carried out concurrently. The HPF directives provide a way to inform the compiler of the recommendation that certain data objects should reside in the same processor: if two data objects are mapped (via the twolevel mapping of alignment and distribution) to the same abstract processor, it is a strong recommendation to the implementation that they ought to reside in the same physical processor. There is also a provision for recommending that a data object be stored in multiple locations, which may complicate any updating of the object but makes it faster 10 for multiple processors to read the object. 11

There is a clear separation between directives that serve as specification statements and 12directives that serve as executable statements (in the sense of the Fortran standards). Spec-13 ification statements are carried out on entry to a program unit, as if all at once; only then 14 are executable statements carried out. (While it is often convenient to think of specification 15statements as being handled at compile time, some of them contain specification expres-16sions, which are permitted to depend on run-time quantities such as dummy arguments, 17and so the values of these expressions may not be available until run time, specifically the 18 very moment that program control enters the scoping unit.) 19

The basic concept is that every array (indeed, every object) is created with some 20 alignment to an entity, which in turn has some distribution onto some arrangement of  $^{21}$ abstract processors. If the specification statements contain explicit specification directives 22 specifying the alignment of an array A with respect to another array B, then the distribution 23 of A will be dictated by the distribution of B; otherwise, the distribution of A itself may be  $^{24}$ specified explicitly. In either case, any such explicit declarative information is used when  $^{25}$ the array is created. 26

Advice to implementors. This model gives a better picture of the actual amount of work that needs to be done than a model that says "the array is created in some default location, and then realigned and/or redistributed if there is an explicit directive." Using ALIGN and DISTRIBUTE specification directives doesn't have to cause any more work at run time than using the implementation defaults. (End of advice to *implementors.*)

In the case of an allocatable object, we say that the object is created whenever it is allocated. Specification directives for an allocatable object may appear in the *specificationpart* of a program unit, but take effect each time the object is created, rather than on entry to the scoping unit.

Alignment is considered an *attribute* (in the Fortran sense) of a data object. If an object 39 A is aligned with an object B, which in turn is already aligned to an object C, this is regarded 40 as an alignment of A with C directly, with B serving only as an intermediary at the time of 41specification. We say that A is *immediately aligned* with B but *ultimately aligned* with C. If 42 an object is not explicitly aligned with another object, we say that it is ultimately aligned 43 with itself. The alignment relationships form a tree with everything ultimately aligned to 44 the object at the root of the tree; however, the tree is always immediately "collapsed" so 45that every object is related directly to the root. 46

Every object that is the root of an alignment tree has an associated *template* or index 47space. Typically, this template has the same rank and size in each dimension as the object 48 associated with it. (The most important exception to this rule is dummy arguments with
 the INHERIT attribute, described in Section 4.4.2.) We often refer to "the template for an
 array," which means the template of the object to which the array is ultimately aligned.
 (When an explicit TEMPLATE (see section 3.7) is used, this may be simply the template to
 which the array is explicitly aligned.)

6 The *distribution* step of the HPF model technically applies to the template of an array, although because of the close relationship noted above we often speak loosely of 7 the distribution of an array. Distribution partitions the template among a set of abstract 8 9 processors according to a given pattern. The combination of alignment (from arrays to templates) and distribution (from templates to processors) thus determines the relationship 10  $1\,1$ of an array to the processors; we refer to this relationship as the *mapping* of the array. 12(These remarks also apply to a scalar, which may be regarded as having an index space whose sole position is indicated by an empty list of subscripts.) 13

14Every object is created as if according to some complete set of specification directives; 15if the program does not include complete specifications for the mapping of some object, 16the compiler provides defaults. By default an object is not aligned with any other object; 17it is ultimately aligned with itself. The default distribution is implementation-dependent, 18 but must be expressible as explicit directives for that implementation. Identically declared 19objects need not be provided with identical default distribution specifications; the compiler 20 may, for example, take into account the contexts in which objects are used in executable 21 code. The programmer may force identically declared objects to have identical distributions 22 by specifying such distributions explicitly. (On the other hand, identically declared pro-23 cessor arrangements are guaranteed to represent "the same processors arranged the same  $^{24}$ way." This is discussed in more detail in section 3.6.)

Sometimes it is desirable to consider a large index space with which several smaller arrays are to be aligned, but not to declare any array that spans the entire index space. HPF allows one to declare a TEMPLATE, which is like an array whose elements have no content and therefore occupy no storage; it is merely an abstract index space that can be distributed and with which arrays may be aligned.

An object is considered to be *explicitly mapped* if it appears in an HPF mapping directive within the scoping unit in which it is declared; otherwise it is *implicitly mapped*. A mapping directive is an ALIGN, or DISTRIBUTE, or INHERIT directive, or any directive that confers an alignment, a distribution, or the INHERIT attribute.

Note that we extend this model in Section 8 to allow dynamic redistribution and remapping of objects.

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## 3.2 Syntax of Data Alignment and Distribution Directives

Specification directives in HPF have two forms: specification statements, analogous to the
 DIMENSION and ALLOCATABLE statements of Fortran; and an attribute form analogous to
 type declaration statements in Fortran using the "::" punctuation.

The attribute form allows more than one attribute to be described in a single directive. HPF goes beyond Fortran in not requiring that the first attribute, or indeed any of them, be a type specifier.

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48 H301 combined-directive

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combined-attribute	is or or or or	ALIGN align-attribute-stuff DISTRIBUTE dist-attribute-stuff INHERIT TEMPLATE PROCESSORS DIMENSION ( explicit-shape-spec-list )						
combined-decl	is or	hpf-entity [ ( explicit-shape-spec-list ) ] object-name	7 8 9					
hpf-entity	is or	processors-name template-name	10 11					
he INHERIT attribute is related ion 4.	to s	ubroutine call conventions and will be discussed	12 13 14					
aint: The same kind of <i>combin</i> given <i>combined-directive</i> .	ned-	attribute must not appear more than once in a	15 16					
Constraint: If the <b>DIMENSION</b> attribute appears in a <i>combined-directive</i> , any entity to whic it applies must be declared with the HPF <b>TEMPLATE</b> or <b>PROCESSORS</b> type specifier.								
ves or in a <i>combined-directive</i> . the <b>DISTRIBUTE</b> attribute is pre- considered to be a <i>distributee</i> and the <b>ALIGN</b> attribute is present, ered to be an <i>alignee</i> and is sub- he HPF keywords <b>PROCESSORS</b> and be cessor arrangements and temp IT play the role of attributes. At or to entities with other types ( at having the type specifier appendent o entity may be given a particul imension information may be spec- a are present, the one after the	sent id is the ject id T lates trib such such ar a ecifie <i>obje</i>	, then every name declared in the <i>combined-decl</i> - subject to the constraints listed in section 3.3. on every name declared in the <i>entity-decl-list</i> is to the constraints listed in section 3.4. <b>EMPLATE</b> play the role of type specifiers in declar- s. The HPF keywords ALIGN, DISTRIBUTE, and utes referring to processor arrangements, to tem- a as REAL) may be combined in an HPF directive ttribute more than once. d after an <i>hpf-entity</i> or in a DIMENSION attribute. <i>ct-name</i> overrides the DIMENSION attribute (this	21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36					
<pre>!HPF\$ TEMPLATE,DIMENSION(64,64) :: A,B,C(32,32),D</pre>								
irectives mapping a variable mu ed. a specification expression include ed in the same specification-part must be completely specified in	st b les a t, an 1 pr n th	e in the same scoping unit where the variable is reference to the value of an element of an array y explicit mapping or INHERIT attribute for the ior specification-directives. (This restriction is the Fortran standard, which states in part: If a	37 38 39 40 41 42 43 44					
	<pre>combined-decl hpf-entity ne INHERIT attribute is related ion 4. aint: The same kind of combin given combined-directive. aint: If the DIMENSION attribut- it applies must be declare ifier. ne following rules constrain the o ves or in a combined-directive. the DISTRIBUTE attribute is pre- considered to be a distributee ar the ALIGN attribute is present, ered to be an alignee and is sub ne HPF keywords PROCESSORS ar ocessor arrangements and temp T play the role of attributes. At or to entities with other types ( t having the type specifier apped be entity may be given a particul mension information may be specifier apped to the TempLATE, DIMENSION (64,64 and D are 64 × 64 templates; C is rectives mapping a variable mu d. a specification expression include ad in the same specification-part must be completely specified in d by and extends F95:7.1.6.2 i </pre>	or or or combined-decl is or hpf-entity is or he INHERIT attribute is related to st ion 4. aint: The same kind of combined- given combined-directive. aint: If the DIMENSION attribute app it applies must be declared wi ifier. the following rules constrain the decla we sor in a combined-directive. the DISTRIBUTE attribute is present considered to be a distribute and is the ALIGN attribute is present, the ered to be an alignee and is subject the HPF keywords PROCESSORS and T ocessor arrangements and templates T play the role of attributes. Attrib- or to entities with other types (such t having the type specifier appear. b entity may be given a particular a imension information may be specifie are present, the one after the object istent with the Fortran standard). If T TEMPLATE, DIMENSION(64,64) : a specification expression includes a din the same specification-part, an must be completely specified in pr d by and extends F95:7.1.6.2 in the	or DISTRIBUTE dist-attribute-stuff or INHERIT or TEMPLATE or TEMPLATE or DIMENSION ( explicit-shape-spec-list ) is hpf-entity [ ( explicit-shape-spec-list ) ] or object-name hpf-entity is processors-name or template-name e INHERIT attribute is related to subroutine call conventions and will be discussed ion 4. aint: The same kind of combined-attribute must not appear more than once in a given combined-directive. aint: If the DIMENSION attribute appears in a combined-directive, any entity to which it applies must be declared with the HPF TEMPLATE or PROCESSORS type spec- ifier. the DISTRIBUTE attribute is present, then every name declared in the combined-decl- ionsidered to be a distribute and is subject to the constraints listed in section 3.3. the ALIGN attribute is present, then every name declared in the entity-decl-list is red to be an dispose and TEMPLATE play the role of type specifier in declar- coressor arrangements and templates. The HPF keywords ALIGN, DISTRIBUTE, and CT play the role of attributes. Attributes referring to processor arrangements, to tem- or to entities with other types (such as REAL) may be combined in an HPF directive t having the type specified appear. o entity may be given a particular attribute more than once. mension information may be specified after an hpf-entity or in a DIMENSION attribute. are present, the one after the object-name overrides the DIMENSION attribute. (the target the fortan standard). For example, in: * TEMPLATE, DIMENSION(64,64) :: A, B, C(32, 32), D and D are 64 × 64 templates; C is 32 × 32. rectives mapping a variable must be in the same scoping unit where the variable is d. a specification expression includes a reference to the value of an element of an array ed in the same specified in prior specification-directives. (This restriction is d by and extends F95:7.1.6.2 in the Fortran standard, which states in part: If a					

A comment on asterisks: The asterisk character "\*" appears in the syntax rules for 47 HPF alignment and distribution directives in three distinct roles: 48 • When a lone asterisk appears as a member of a parenthesized list, it indicates either a collapsed mapping, wherein many elements of an array may be mapped to the same abstract processor, or a replicated mapping, wherein each element of an array may be mapped to many abstract processors. See the syntax rules for *align-source* and *align-subscript* (see section 3.4) and for *dist-format* (see section 3.3).

• An asterisk appearing in an *align-subscript-use* expression represents the usual integer multiplication operator.

• When an asterisk appears before a left parenthesis "(" or after the keyword WITH or ONTO, it indicates a descriptive or transcriptive mapping for dummy arguments of subprograms (see Section 4) and for mapping of pointers under the approved extensions (see section 8.8).

An asterisk can also be used in the PASS\_BY attribute in an interface block to describe dummy arguments passed by reference to an extrinsic routine written in C (see Section 11.2).

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## 3.3 The DISTRIBUTE Directive

The DISTRIBUTE directive specifies a mapping of data objects to abstract processors in a processor arrangement. For example,

### REAL SALAMI(10000)

### !HPF\$ DISTRIBUTE SALAMI(BLOCK)

specifies that the array SALAMI should be distributed across some set of abstract processors
by slicing it uniformly into blocks of contiguous elements. If there are 50 processors, the
directive implies that the array should be divided into groups of [10000/50] = 200 elements,
with SALAMI (1:200) mapped to the first processor, SALAMI (201:400) mapped to the second
processor, and so on. If there is only one processor, the entire array is mapped to that
processor as a single block of 10000 elements.

The block size may be specified explicitly:

REAL WEISSWURST(10000) !HPF\$ DISTRIBUTE WEISSWURST(BLOCK(256))

This specifies that groups of exactly 256 elements should be mapped to successive abstract processors. (There must be at least [10000/256] = 40 abstract processors if the directive is to be satisfied. The fortieth processor will contain a partial block of only 16 elements, namely WEISSWURST(9985:10000).)

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HPF also provides a cyclic distribution format:

### REAL DECK\_OF\_CARDS(52) !HPF\$ DISTRIBUTE DECK\_OF\_CARDS(CYCLIC)

If there are 4 abstract processors, the first processor will contain DECK\_OF\_CARDS(1:49:4),
 the second processor will contain DECK\_OF\_CARDS(2:50:4), the third processor will contain
 DECK\_OF\_CARDS(3:51:4), and the fourth processor will contain DECK\_OF\_CARDS(4:52:4).
 Successive array elements are dealt out to successive abstract processors in round-robin
 fashion.

47 Distributions are specified independently for each dimension of a multidimensional
 48 array:

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INTEGER CHESS_BOARD(8, HPF\$ DISTRIBUTE CHESS_BOARD						
!HPF\$ DISTRIBUTE GO_BOARD(CY						
be distributed onto a two-dimension array will have its rows distributed cy processors. (The "*" specifies that G	ed up into contiguous rectangular patches, which will al arrangement of abstract processors. The GO_BOARD clically over a one-dimensional arrangement of abstract O_BOARD is not to be distributed along its second axis; as one object. This is sometimes called "on-processor" 9					
	appear only in the <i>specification-part</i> of a scoping unit $expr$ as the argument to a BLOCK or CYCLIC option. $12$					
H305 distribute-directive	is DISTRIBUTE distributee dist-directive-stuff					
H306 dist-directive-stuff	is dist-format-clause [ dist-onto-clause ]					
H307 dist-attribute-stuff	is dist-directive-stuff or dist-onto-clause					
H308 distributee	is object-name 20 or template-name 21					
H309 dist-format-clause	is (dist-format-list) or * (dist-format-list) or * 22 23 24 24 24					
H310 dist-format	is BLOCK [ ( scalar-int-expr ) ] 25 or CYCLIC [ ( scalar-int-expr ) ] 27 or * 28					
H311 dist-onto-clause	is ONTO dist-target 29					
H312 dist-target	is processors-name 31 or * processors-name 32 or * 33					
The full syntax is given here for only in Section 4. These "interproce	completeness. However, some of the forms are discussed dural" forms are:					
• The last two options of rule H:	$\begin{array}{c} 309 \ (\text{containing the } \star \text{ form}) \end{array} \right. \begin{array}{c} 37 \\ 38 \end{array}$					
• The last two options of rule H						
Constraint: An <i>object-name</i> mentio subobject designator or	40 ned as a <i>distributee</i> must be a simple name and not a a <i>component-name</i> . 42					
Constraint: An object-name mentioned as a distributee may not appear as an alignee.						
Constraint: An <i>object-name</i> mention tribute.	pned as a <i>distributee</i> may not have the POINTER at- $45$					
Constraint: An object-name mention	ned as a <i>distributee</i> may not have the <b>TARGET</b> attribute. 48					

1 2 3	Constraint:	If the <i>distributee</i> is scalar, the <i>dist-format-list</i> (and its surrounding parentheses) must not appear. In this case, the statement form of the directive is allowed only if a <i>dist-format-clause</i> of "*" is present.					
4 5 6	Constraint:	If a <i>dist-format-list</i> is specified, its length must equal the rank of each <i>distribu-</i> <i>tee</i> to which it applies.					
7 8 9 10	Constraint:	If both a <i>dist-format-list</i> and a <i>dist-target</i> appear, the number of elements of the <i>dist-format-list</i> that are not "*" must equal the rank of the specified processor arrangement.					
11 12	Constraint:	If a <i>dist-target</i> appears but not a <i>dist-format-list</i> , the rank of each <i>distributee</i> must equal the rank of the specified processor arrangement.					
13 14 15	Constraint:	If either the <i>dist-format-clause</i> or the <i>dist-target</i> in a <b>DISTRIBUTE</b> directive begins with "*" then every <i>distributee</i> must be a dummy argument.					
16 17	Constraint:	Any <i>scalar-int-expr</i> appearing in a <i>dist-format</i> of a <b>DISTRIBUTE</b> directive must be a <i>specification-expr</i> .					
18 19 20 21 22	extens 1), and	e to users. Some of the above constraints are relaxed under the approved ions (see Section 8): mapping of derived type components (relaxes constraint d mapping of pointers and targets (relaxes constraints 3, 4, and 9). (End of to users.)					
23 24	Note th	at the possibility of a DISTRIBUTE directive of the form					
25	!HPF\$ DI	$\texttt{STRIBUTE} \ dist-attribute-stuff :: distributee-list$					
26 27 28	is covered by Exampl	y syntax rule H301 for a <i>combined-directive</i> . les:					
29 30 31		STRIBUTE D1(BLOCK) STRIBUTE (BLOCK,*,BLOCK) ONTO SQUARE:: D2,D3,D4					
32 33 34 35 36 37 38 39 40 41 42 43 44 45 46 47 48	The meanings of the alternatives for $dist-format$ are given below. Define the ceiling division function $CD(J,K) = (J+K-1)/K$ (using Fortran integer arithmetic with truncation toward zero.) Define the ceiling remainder function $CR(J,K) = J-K*CD(J,K)$ . The dimensions of a processor arrangement appearing as a <i>dist-target</i> are said to correspond in left-to-right order with those dimensions of a <i>distributee</i> for which the corresponding <i>dist-format</i> is not *. In the example above, processor arrangement SQUARE must be two-dimensional; its first dimension corresponds to the first dimensions of D2, D3, and D4 and its second dimension corresponds to the third dimensions of D2, D3, and D4. Let d be the size of a <i>distributee</i> in a certain dimension and let p be the size of the processor arrangement in the corresponding dimension. For simplicity, assume all dimension have a lower bound of 1. Then BLOCK(m) means that a <i>distributee</i> position whose indet along that dimension is j is mapped to an abstract processor whose index along the corresponding dimension of the processor arrangement is $CD(j,m)$ (note that $m \times p \ge d$ must be true), and is position number $m+CR(j,m)$ among positions mapped to that abstract processor k along that axis is position number $1+m*(k-1)$ .						

 $\Downarrow$ 

The block size m must be a positive integer. BLOCK by definition means the same as BLOCK(CD(d, p)). CYCLIC(m) means that a *distributee* position whose index along that dimension is j is mapped to an abstract processor whose index along the corresponding dimension of the processor arrangement is 1+MODULO(CD(j,m)-1,p). The first distributee position in abstract processor k along that axis is position number 1+m\*(k-1). The block size m must be a positive integer. CYCLIC by definition means the same as CYCLIC(1). CYCLIC(m) and BLOCK(m) imply the same distribution when  $m \times p \ge d$ , but BLOCK(m) additionally asserts that the distribution will not wrap around in a cyclic manner, which a compiler cannot determine at compile time if m is not constant. Note that CYCLIC and **BLOCK** (without argument expressions) do not imply the same distribution unless p > d, a degenerate case in which the block size is 1 and the distribution does not wrap around. Suppose that we have 16 abstract processors and an array of length 100: !HPF\$ PROCESSORS SEDECIM(16) REAL CENTURY (100) Distributing the array BLOCK (which in this case would mean the same as BLOCK(7)): !HPF\$ DISTRIBUTE CENTURY(BLOCK) ONTO SEDECIM results in this mapping of array elements onto abstract processors:  $^{24}$  $^{25}$ Distributing the array BLOCK(8): !HPF\$ DISTRIBUTE CENTURY(BLOCK(8)) ONTO SEDECIM results in this mapping of array elements onto abstract processors:  $\mathbf{2}$ 

1	9	17	25	33	41	49	57	65	73	81	89	97		
2	10	18	26	34	42	50	58	66	74	82	90	98		
3	11	19	27	35	43	51	59	67	75	83	91	99		
4	12	20	28	36	44	52	60	68	76	84	92	100		
5	13	21	29	37	45	53	61	69	77	85	93			
6	14	22	30	38	46	54	62	70	78	86	94			
7	15	23	31	39	47	55	63	71	79	87	95			
8	16	24	32	40	48	56	64	72	80	88	96			

#### 3.3. THE DISTRIBUTE DIRECTIVE

Distributing the array BLOCK(6) is not HPF-conforming because  $6 \times 16 < 100$ . Distributing the array CYCLIC (which means exactly the same as CYCLIC(1)):

#### !HPF\$ DISTRIBUTE CENTURY(CYCLIC) ONTO SEDECIM

results in this mapping of array elements onto abstract processors:

1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16
1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16
17	18	19	20	21	22	23	24	25	26	27	28	29	30	31	32
33	34	35	36	37	38	39	40	41	42	43	44	45	46	47	48
49	50	51	52	53	54	55	56	57	58	59	60	61	62	63	64
65	66	67	68	69	70	71	72	73	74	75	76	77	78	79	80
81	82	83	84	85	86	87	88	89	90	91	92	93	94	95	96
97	98	99	100												

Distributing the array CYCLIC(3):

#### !HPF\$ DISTRIBUTE CENTURY(CYCLIC(3)) ONTO SEDECIM

results in this mapping of array elements onto abstract processors:

1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16
1	4	7	10	13	16	19	22	25	28	31	34	37	40	43	46
2	5	8	11	14	17	20	23	26	29	32	35	38	41	44	47
3	6	9	12	15	18	21	24	27	30	33	36	39	42	45	48
49	52	55	58	61	64	67	70	73	76	79	82	85	88	91	94
50	53	56	59	62	65	68	71	74	77	80	83	86	89	92	95
51	54	57	60	63	66	69	72	75	78	81	84	87	90	93	96
97	100														
98															
99															

Note that it is perfectly permissible for an array to be distributed so that some processors have no elements. Indeed, an array may be "distributed" so that all elements reside on one processor. For example,

#### !HPF\$ DISTRIBUTE CENTURY(BLOCK(256)) ONTO SEDECIM

results in having only one non-empty block—a partially-filled one at that, having only 100 elements—on processor 1, with processors 2 through 16 having no elements of the array.

The statement form of a **DISTRIBUTE** directive may be considered an abbreviation for an attributed form that happens to mention only one *distributee*; for example,

!HPF\$ DISTRIBUTE distributee ( dist-format-list ) ONTO dist-target

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is equivalent to

#### **!HPF\$ DISTRIBUTE (** *dist-format-list* **) ONTO** *dist-target* :: *distributee*

Note that, to prevent syntactic ambiguity, the *dist-format-clause* must be present in the statement form, so in general the statement form of the directive may not be used to specify the mapping of scalars.

If the *dist-format-clause* is omitted from the attributed form, then the language processor may make an arbitrary choice of distribution formats for each template or array. So the directive

!HPF\$ DISTRIBUTE ONTO P :: D1,D2,D3

means the same as

!HPF\$ DISTRIBUTE ONTO P :: D1
!HPF\$ DISTRIBUTE ONTO P :: D2
!HPF\$ DISTRIBUTE ONTO P :: D3

to which a compiler, perhaps taking into account patterns of use of D1, D2, and D3 within the code, might choose to supply three distinct distributions such as, for example,

!HPF\$ DISTRIBUTE D1(BLOCK, BLOCK) ONTO P !HPF\$ DISTRIBUTE D2(CYCLIC, BLOCK) ONTO P !HPF\$ DISTRIBUTE D3(BLOCK(43),CYCLIC) ONTO P

Then again, the compiler might happen to choose the same distribution for all three arrays.

In either the statement form or the attributed form, if the ONTO clause is present, it specifies the processor arrangement that is the target of the distribution. If the ONTO clause is omitted, then a implementation-dependent processor arrangement is chosen arbitrarily for each *distributee*. So, for example,

```
REAL, DIMENSION(1000) :: ARTHUR, ARNOLD, LINUS, LUCY

!HPF$ PROCESSORS EXCALIBUR(32)

!HPF$ DISTRIBUTE (BLOCK) ONTO EXCALIBUR :: ARTHUR, ARNOLD

!HPF$ DISTRIBUTE (BLOCK) :: LINUS, LUCY
```

causes the arrays ARTHUR and ARNOLD to have the same mapping, so that corresponding elements reside in the same abstract processor, because they are the same size and distributed in the same way (BLOCK) onto the same processor arrangement (EXCALIBUR). However, LUCY and LINUS do not necessarily have the same mapping because they might, depending on the implementation, be distributed onto differently chosen processor arrangements; so corresponding elements of LUCY and LINUS might not reside on the same abstract processor. (The ALIGN directive provides a way to ensure that two arrays have the same mapping without having to specify an explicit processor arrangement.)

In a given environment, for some distributions, there may be no appropriate processor arrangement.

1	<b>3.4</b> The	ALIGN Directive		
2 3 4 5 6 7 8 9 10 11 12 13	same way as to be more e (because tw processor). ' mappings fo cases throug frequently m The AL can contain	s certain other data object efficient than operations be to objects that are aligned The ALIGN directive is des or all the elements of an a gh careful use of matching nore convenient. IGN directive may appear	s. O etwe d ar igne rray g DI only us a	at certain data objects are to be mapped in the perations between aligned data objects are likely en data objects that are not known to be aligned e intended to be mapped to the same abstract d to make it particularly easy to specify explicit at once. While objects can be aligned in some STRIBUTE directives, ALIGN is more general and y in the <i>specification-part</i> of a scoping unit and <i>subscript</i> or in a <i>subscript-triplet</i> .
14 15	H313 align-	-directive	is	ALIGN alignee align-directive-stuff
16	H314 align	-directive-stuff	is	( $align-source-list$ ) $align-with-clause$
17 18	H315 align-	-attribute-stuff	is	[ ( align-source-list ) ] align-with-clause
19	H316 align	ee	is	object-name
20 21	H317 align-	-source	is	:
22			or	*
23			or	align-dummy
24 25	H318 align-	-dummy	is	scalar- $int$ - $variable$
26 27	Constraint:	An <i>object-name</i> mention subobject designator or a		s an <i>alignee</i> must be a simple name and not a <i>nponent-name</i> .
28 29	Constraint:	An object-name mentione	ed as	s an <i>alignee</i> may not appear as a <i>distributee</i> .
30 31	Constraint:	An object-name mentione	ed as	s an <i>alignee</i> may not have the <b>POINTER</b> attribute.
32 33	Constraint:	An object-name mentione	ed as	s an <i>alignee</i> may not have the <b>TARGET</b> attribute.
34 35	Constraint:			<i>ign-source-list</i> (and its surrounding parentheses) the statement form of the directive is not allowed.
36 37 38	Constraint:	If the <i>align-source-list</i> is p to which it applies.	prese	ent, its length must equal the rank of each <i>alignee</i>
39 40	Constraint:	An align-dummy must be	e a n	amed variable.
41 42	Constraint:	An object may not have	both	the INHERIT attribute and the ALIGN attribute.
43 44 45 46	extens	ions (see Section 8): map mapping of pointers and	ping	ove constraints are relaxed under the approved of derived type components (relaxes constraint gets (relaxes constraints 3 and 4). ( <i>End of advice</i>
47 48	Note th	at the possibility of an AL	IGN	directive of the form

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!HPF\$ ALIGN align-attribute-stuff :: alignee-list	1
is covered by syntax rule H301 for a <i>combined-directive</i> .	2
The statement form of an ALIGN directive may be considered an abbreviation of an	3 4
attributed form that happens to mention only one <i>alignee</i> :	4 5
!HPF\$ ALIGN $alignee$ ( $align-source-list$ ) WITH $align-spec$	6 7
is equivalent to	8 9
	10
!HPF\$ ALIGN ( <i>align-source-list</i> ) WITH <i>align-spec</i> :: <i>alignee</i>	11
If the <i>align-source-list</i> is omitted from the attributed form and the <i>alignees</i> are not	12
scalar, the <i>align-source-list</i> is assumed to consist of a parenthesized list of ":" entries,	13
equal in number to the rank of the <i>alignees</i> . Similarly, if the <i>align-subscript-list</i> is omitted	14
from the <i>align-spec</i> in either form, it is assumed to consist of a parenthesized list of ":"	15
entries, equal in number to the rank of the <i>align-target</i> . So the directive	16
	17
!HPF\$ ALIGN WITH B :: A1, A2, A3	18
	19
means	20
!HPF\$ ALIGN (:,:) WITH B(:,:) :: A1, A2, A3	21
mry Align (.,.) with D(.,.) Ar, AZ, AS	22 23
which in turn means the same as	24
	25
<pre>!HPF\$ ALIGN A1(:,:) WITH B(:,:)</pre>	26
<pre>!HPF\$ ALIGN A2(:,:) WITH B(:,:) !UDE4 ALIGN A2()</pre>	27
<pre>!HPF\$ ALIGN A3(:,:) WITH B(:,:)</pre>	28
because an attributed-form directive that mentions more than one <i>alignee</i> is equivalent to	29
a series of identical directives, one for each <i>alignee</i> ; all <i>alignee</i> s must have the same rank.	30
With this understanding, we will assume below, for the sake of simplifying the description,	31
that an ALIGN directive has a single <i>alignee</i> .	32
Each <i>align-source</i> corresponds to one axis of the <i>alignee</i> , and is specified as either ":"	33
or "*" or a dummy variable:	34
	35 36
• If it is ":", then positions along that axis will be spread out across the matching axis	37
of the <i>align-spec</i> (see below).	38
• If it is "*", then that axis is <i>collapsed</i> : positions along that axis make no difference	39
in determining the corresponding position within the <i>align-target</i> . (Replacing the "*"	40
with a dummy variable name not used anywhere else in the directive would have the	41
same effect; "*" is merely a convenience that saves the trouble of inventing a variable	42
name and makes it clear that no dependence on that dimension is intended.)	43
	44
• A dummy variable is considered to range over all valid index values for that dimension	45
of the <i>alignee</i> .	46
	47
The WITH clause of an ALIGN has the following syntax:	48

1	H319	a lign-with-clause	is	WITH align-spec
2 3 4	H320	align-spec	is or	align-target [ ( align-subscript-list ) ] * align-target [ ( align-subscript-list ) ]
5 6	H321	align-target	is or	object-name template-name
7 8 9 10	H322	a lign-subscript	is or or or	int-expr align-subscript-use subscript-triplet <b>*</b>
11 12 13 14	H323	align-subscript-use	is or	[ [ int-level-two-expr ] add-op ] align-add-operand align-subscript-use add-op int-add-operand
15 16	H324	align-add-operand	is or	[ int-add-operand * ] align-primary align-add-operand * int-mult-operand
17 18 19	H325	align-primary	is or	align-dummy ( align-subscript-use )
20	H326	int- $add$ - $operand$	is	add-operand
21	H327	int-mult-operand	is	mult-operand
22 23	H328	int-level-two-expr	is	level-2-expr
24 25 26 27 28	only i rule H	n Section 4. These "interproce [320 (containing the * form).	edura	eteness. However, some of the forms are discussed al" forms are those using the second option of

Constraint: An object-name mentioned as an align-target must be a simple name and not 29 a subobject designator or a *component-name*. 30

31 Constraint: An *align-target* may not have the OPTIONAL attribute.

40

32 Constraint: If the align-spec in an ALIGN directive begins with "\*" then every alignee must 33 be a dummy argument. 34

35 Constraint: In an align-directive any int-expr, int-level-two-expr, int-add-operand or int-36 mult-operand must be a specification expression. 37

Constraint: Any subscript or stride in a subscript-triplet that is an align-subscript in an 38 align-directive must be a specification expression. 39

Constraint: Each align-dummy may appear at most once in an align-subscript-list. 41

42 Constraint: An *align-subscript-use* expression may contain at most one occurrence of an 43 align-dummy. 44

Constraint: A scalar-int-variable that is used as an align-dummy may not appear any-45where in the *align-spec* except where explicitly permitted to appear by virtue 46 of the grammar shown above. Paraphrased, one may construct an *align*-47subscript-use only by starting with an align-dummy and then doing additive 48

and multiplicative things to it with integer specification expressions that contain no *align-dummy*.

- Constraint: A *subscript* within an *align-subscript* may not contain occurrences of any *align- dummy*.
- Constraint: An *int-add-operand*, *int-mult-operand*, or *int-level-two-expr* must be of type integer.

Advice to users. Some of the above constraints are relaxed under the approved extensions (see Section 8): mapping of derived type components (relaxes constraint 1), mapping of pointers (relaxes constraint 3) and remapping of data objects (relaxes constraints 4 and 5). (End of advice to users.)

The syntax rules for an *align-subscript-use* take account of operator precedence issues, but the basic idea is simple: an *align-subscript-use* is intended to be a linear (more precisely: affine) function of a single occurrence of an *align-dummy*.

For example, the following *align-subscript-use* expressions are valid, assuming that each of J, K, and M is an *align-dummy* and N is not an *align-dummy*:

J	J+1	З-К	2*M	N*M	100-3*M
-J	+ J	-K+3	M+2**3	M+N	-(4*7+IOR(6,9))*K-(13-5/3)
M*2	N*(M-N)	2*(J+1)	5-K <b>+</b> 3	10000-M*3	2*(3*(K-1)+13)-100

The following expressions are not valid *align-subscript-use* expressions:

J+J	J-J	3*K-2*K	M*(N-M)	2*J-3*J+J	2*(3*(K-1)+13)-K
J*J	J+K	3/K	2**M	M*K	K-3*M
K-J	IOR(J,1)	-K/3	M*(2+M)	M*(M-N)	2**(2*J-3*J+J)

The *align-spec* must contain exactly as many *subscript-triplets* as the number of colons (":") appearing in the *align-source-list*. These are matched up in corresponding left-to-right order, ignoring, for this purpose, any *align-source* that is not a colon and any *align-subscript* that is not a *subscript-triplet*. Consider a dimension of the *alignee* for which a colon appears as an *align-source* and let the lower and upper bounds of that dimension be LA and UA. Let the corresponding subscript triplet be LT:UT:ST or its equivalent. Then the colon could be replaced by a new, as-yet-unused dummy variable, say J, and the subscript triplet by the expression (J-LA)\*ST+LT without affecting the mapping specified by the directive. However, the colon form additionally requires that the axes must conform, which means that 

$$\max(0, UA - LA + 1) = \max(0, [(UT - LT + 1)/ST])$$

must be true. (This is entirely analogous to the treatment of array assignment.)

To simplify the remainder of the discussion, we assume that every colon in the *align-source-list* has been replaced by new dummy variables in exactly the fashion just described, and that every "\*" in the *align-source-list* has likewise been replaced by an otherwise unused dummy variable. For example,

!HPF\$ ALIGN A(:,\*,K,:,:,\*) WITH B(31:,:,K+3,20:100:3)

may be transformed into its equivalent

∜

!HPF\$ ALIGN A(I,J,K,L,M,N) WITH B(I-LBOUND(A,1)+31, 1 *&*: !HPF\$ L-LBOUND(A,4)+LBOUND(B,2),K+3,(M-LBOUND(A,5))\*3+20)2 3 with the attached requirements 4 5 SIZE(A,1) .EQ. UBOUND(B,1)-30 6 SIZE(A,4) .EQ. SIZE(B,2)7 SIZE(A,5) .EQ. (100-20+3)/3 8 Thus we need consider further only the case where every *align-source* is a dummy variable 9 and no *align-subscript* is a *subscript-triplet*. 10 Each dummy variable is considered to range over all valid index values for the cor-11 responding dimension of the *alignee*. Every combination of possible values for the index 12variables selects an element of the *alignee*. The *align-spec* indicates a corresponding element 13 (or section) of the *align-target* with which that element of the *alignee* should be aligned; 14 this indication may be a function of the index values, but the nature of this function is 15 syntactically restricted (as discussed above) to linear (precisely: affine) functions in order 16 to limit the complexity of the implementation. Each *align-dummy* variable may appear at 17most once in the *align-spec* and only in certain rigidly prescribed contexts. The result is 18 that each *align-subscript* expression may contain at most one *align-dummy* variable and the 19 expression is constrained to be a linear function of that variable. (Therefore skew alignments 20 are not possible.) 21 An asterisk "\*" as an *align-subscript* indicates a replicated representation. Each ele-22 ment of the *aliqnee* is aligned with every position along that axis of the *aliqn-target*. 23 24 Rationale. It may seem strange to use "\*" to mean both collapsing and replication; 25the rationale is that "\*" always stands conceptually for a dummy variable that appears 26 nowhere else in the statement and ranges over the set of indices for the indicated 27 dimension. Thus, for example, 28 29 !HPF\$ ALIGN A(:) WITH D(:,\*) 30 31 means that a copy of A is aligned with every column of D, because it is conceptually 32 equivalent to 33 34 for every legitimate index j, align A(:) with D(:,j)35 36 just as 37 38 !HPF\$ ALIGN A(:,\*) WITH D(:) 39 40 is conceptually equivalent to 4142 for every legitimate index j, align A(:,j) with D(:)43 44 Note, however, that while HPF syntax allows 45 !HPF\$ ALIGN A(:,\*) WITH D(:) 46 47 to be written in the alternate form 48

!HPF\$ ALIGN A(:,J) WITH D(:)

it does not allow

!HPF\$ ALIGN A(:) WITH D(:,\*)

to be written in the alternate form

!HPF\$ ALIGN A(:) WITH D(:,J)

because that has another meaning (only a variable appearing in the *align-source-list* following the *alignee* is understood to be an *align-dummy*, so the current value of the variable J is used, thus aligning A with a single column of D).

Replication allows an optimizing compiler to arrange to read whichever copy is closest. (Of course, when a replicated data object is written, all copies must be updated, not just one copy. Replicated representations are very useful for small lookup tables, where it is much faster to have a copy in each physical processor but without giving it an extra dimension that is logically unnecessary to the algorithm. (*End of rationale.*)

By applying the transformations given above, all cases of an *align-subscript* may be conceptually reduced to either an *int-expr* (not involving an *aliqn-dummy*) or an *aliqn*- $^{21}$ subscript-use, and the align-source-list may be reduced to a list of index variables with no "\*" or ":". An align-subscript-list may then be evaluated for any specific combination of values for the *aliqn-dummy* variables simply by evaluating each *aliqn-subscript* as an  $^{24}$ expression. The resulting subscript values must be legitimate subscripts for the *align-target*.  $^{25}$ (This implies that the *alignee* is not allowed to "wrap around" or "extend past the edges" of an *align-target*.) The selected element of the *alignee* is then considered to be aligned with the indicated element of the *align-target*; more precisely, the selected element of the alignee is considered to be ultimately aligned with the same object with which the indicated element of the *align-target* is currently ultimately aligned (possibly itself). 

More examples of ALIGN directives:

	INTEG	ER D1(N)
	LOGIC	AL D2(N,N)
	REAL,	DIMENSION(N,N):: X,A,B,C,AR1,AR2A,P,Q,R,S
!HPF\$	ALIGN	X(:,*) WITH D1(:)
!HPF\$	ALIGN	(:,*) WITH D1:: A,B,C,AR1,AR2A
!HPF\$	ALIGN	WITH D2:: P,Q,R,S

Note that, in a *alignee-list*, the alignees must all have the same rank but need not all have the same shape; the extents need match only for dimensions that correspond to colons in the *align-source-list*. This turns out to be an extremely important convenience; one of the most common cases in current practice is aligning arrays that match in distributed ("parallel") dimensions but may differ in collapsed ("on-processor") dimensions:

	REAL A(3,N), $B(4,N)$ , C	C(43,N), Q(N)
!HPF\$	DISTRIBUTE Q(BLOCK)	
!HPF\$	ALIGN (*,:) WITH Q:: A	A,B,C

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Here there are processors (perhaps  $\mathbb{N}$  of them) and arrays of different sizes (3, 4, 43) within 1 each processor are required. As far as HPF is concerned, the numbers 3, 4, and 43 may be  $^{2}$ different, because those axes will be collapsed. Thus array elements with indices differing 3 only along that axis will all be aligned with the same element of Q (and thus be specified 4 as residing in the same processor). 5

In the following examples, each directive in a group means the same thing, assuming 6 that corresponding axis upper and lower bounds match:

```
8
       !Second axis of X is collapsed
9
       !HPF$ ALIGN X(:,*) WITH D1(:)
10
       !HPF$ ALIGN X(J,*) WITH D1(J)
11
       !HPF$ ALIGN X(J,K) WITH D1(J)
12
13
       !Replicated representation along second axis of D3
14
       !HPF$ ALIGN X(:,:) WITH D3(:,*,:)
15
       !HPF$ ALIGN X(J,K) WITH D3(J,*,K)
16
17
       !Transposing two axes
18
       !HPF$ ALIGN X(J,K) WITH D2(K,J)
19
       !HPF$ ALIGN X(J,:) WITH D2(:,J)
20
       !HPF$ ALIGN X(:,K) WITH D2(K,:)
21
       !But there isn't any way to get rid of *both* index variables;
22
       ! the subscript-triplet syntax alone cannot express transposition.
23
^{24}
       !Reversing both axes
^{25}
       !HPF$ ALIGN X(J,K) WITH D2(M-J+1,N-K+1)
26
       !HPF$ ALIGN X(:,:) WITH D2(M:1:-1,N:1:-1)
27
28
       !Simple case
29
       !HPF$ ALIGN X(J,K) WITH D2(J,K)
30
       !HPF$ ALIGN X(:,:) WITH D2(:,:)
31
       !HPF$ ALIGN (J,K) WITH D2(J,K):: X
32
       !HPF$ ALIGN (:,:) WITH D2(:,:):: X
33
       !HPF$ ALIGN WITH D2:: X
34
```

#### 3.5**Allocatable Arrays and Pointers**

A variable with the ALLOCATABLE attribute may appear as an *alignee* in an ALIGN directive or as a *distributee* in a **DISTRIBUTE** directive. Such directives do not take effect immediately, however; they take effect each time the array is allocated by an ALLOCATE statement, rather than on entry to the scoping unit. The values of all specification expressions in such a directive are determined once on entry to the scoping unit and may be used multiple times (or not at all). For example:

```
SUBROUTINE MILLARD_FILLMORE(N,M)
44
             REAL, ALLOCATABLE, DIMENSION(:) :: A, B
45
       !HPF$ ALIGN B(I) WITH A(I+N)
46
       !HPF$ DISTRIBUTE A(BLOCK(M*2))
47
             N = 43
48
```

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 $1\,1$ 

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M = 91 ALLOCATE(A(27)) ALLOCATE(B(13))

The values of the expressions N and M\*2 on entry to the subprogram are conceptually retained by the ALIGN and DISTRIBUTE directives for later use at allocation time. When the array A is allocated, it is distributed with a block size equal to the retained value of M\*2, not the value 182. When the array B is allocated, it is aligned relative to A according to the retained value of N, not its new value 43.

Note that it would have been incorrect in the MILLARD\_FILLMORE example to perform the two ALLOCATE statements in the opposite order. In general, when an object X is created it may be aligned to another object Y only if Y has already been created or allocated. The following example illustrates several related cases.

	SUBROUTINE WARREN_HARDING(P,Q)	
	REAL P(:)	
	REAL Q(:)	
	REAL R(SIZE(Q))	
	REAL, ALLOCATABLE :: S(:),T(:)	
!HPF\$	ALIGN P(I) WITH T(I)	!Nonconforming
!HPF\$	ALIGN Q(I) WITH *T(I)	!Nonconforming
!HPF\$	ALIGN R(I) WITH T(I)	!Nonconforming
!HPF\$	ALIGN S(I) WITH T(I)	
	ALLOCATE(S(SIZE(Q)))	!Nonconforming
	ALLOCATE(T(SIZE(Q)))	

Three ALIGN directives are not HPF-conforming because the array T has not yet been 26 allocated at the time that the various alignments must take place. The four cases differ 27slightly in their details. The arrays P and Q already exist on entry to the subroutine, but 28 because T is not yet allocated, one cannot correctly prescribe the alignment of P or describe 29 the alignment of Q relative to T. (See Section 4 for a discussion of prescriptive and descriptive 30 directives.) The array R is created on subroutine entry and its size can correctly depend 31 on the SIZE of Q, but the alignment of R cannot be specified in terms of the alignment 32 of T any more than its size can be specified in terms of the size of T. It is permitted to 33 have an alignment directive for S in terms of T, because the alignment action does not take 34 place until S is allocated; however, the first ALLOCATE statement is nonconforming because 35 S needs to be aligned but at that point in time T is still unallocated. 36

When an array is allocated, it will be aligned to an existing object or template if there <sup>37</sup> is an explicit ALIGN directive for the allocatable variable. If there is no explicit ALIGN <sup>38</sup> directive, then the array will be ultimately aligned with itself. It is forbidden for any other <sup>39</sup> object to be ultimately aligned to an array at the time the array becomes undefined by <sup>40</sup> reason of deallocation. All this applies regardless of whether the name originally used in <sup>41</sup> the ALLOCATE statement when the array was created had the ALLOCATABLE attribute or the <sup>42</sup> POINTER attribute. <sup>43</sup>

Pointers cannot be explicitly mapped in HPF and thus can only be associated with objects which are not explicitly mapped. When used for allocation, the compiler may choose any arbitrary mapping for data allocated through the pointer. Explicit mapping of pointers is allowed under the approved extensions (see section 8.8). Also, the relationship of pointers and sequence attributes is described in section 3.8.

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#### 3.6 The PROCESSORS Directive

The PROCESSORS directive declares one or more rectilinear processor arrangements, specifying for each one its name, its rank (number of dimensions), and the extent in each dimension. It may appear only in the *specification-part* of a scoping unit. Every dimension of a processor arrangement must have nonzero extent; therefore a processor arrangement cannot be empty.

In the language of F95:14.1.2 in the Fortran standard, processor arrangements are local entities of class (1); therefore a processor arrangement may not have the same name as a variable, named constant, internal procedure, etc., in the same scoping unit. Names of processor arrangements obey the same rules for host and use association as other names in the long list in F95:12.1.2.2.1 in the Fortran standard.

A processor arrangement declared in a module has the default accessibility of the module.

Rationale. Because the name of a processor arrangement is not a first-class entity in HPF, but must appear only in directives, it cannot appear in an access-stmt (PRIVATE or PUBLIC). If directives ever become full-fledged Fortran statements rather than structured comments, then it would be appropriate to allow the accessibility of a processor arrangement to be controlled by listing its name in an access-stmt. (End of rationale.)

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If two processor arrangements have the same shape, then corresponding elements of the two arrangements are understood to refer to the same abstract processor. (It is anticipated that implementation-dependent directives provided by some HPF implementations could overrule the default correspondence of processor arrangements that have the same shape.)

If directives collectively specify that two objects be mapped to the same abstract pro cessor at a given instant during the program execution, the intent is that the two objects
 be mapped to the same physical processor at that instant.

The intrinsic functions NUMBER\_OF\_PROCESSORS and PROCESSORS\_SHAPE may be used to inquire about the total number of actual physical processors used to execute the program. This information may then be used to calculate appropriate sizes for the declared abstract processor arrangements.

35	H329 pr	rocessors-dire	ctive is	PROCESSORS pr	ocessors-decl-list
36 37	Н330 <i>рт</i>	rocessors-decl	is	processors-nam	e shape-spec-list ) ]
38 39					mape spee not , ]
40	Exa	mples:			
41					
42	!HPF\$	PROCESSORS	P(N)		
43	!HPF\$	PROCESSORS	Q(NUMBER_OF_PROC	CESSORS()),	&
44	!HPF\$		R(8,NUMBER_OF_PF	ROCESSORS()/8)	
45	!HPF\$	PROCESSORS	BIZARRO(1972:199	97,-20:17)	
46	!HPF\$	PROCESSORS	SCALARPROC		
47					

<sup>48</sup> If no shape is specified, then the declared processor arrangement is conceptually scalar.

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A scalar processor arrangement may be useful as a way of indicating Rationale. 1 that certain scalar data should be kept together but need not interact strongly with 2 distributed data. Depending on the implementation architecture, data distributed 3 onto such a processor arrangement may reside in a single "control" or "host" processor 4 (if the machine has one), or may reside in an arbitrarily chosen processor, or may be 5replicated over all processors. For target architectures that have a set of computational 6 processors and a separate scalar host computer, a natural implementation is to map 7every scalar processor arrangement onto the host processor. For target architectures 8 that have a set of computational processors but no separate scalar "host" computer, 9 data mapped to a scalar processor arrangement might be mapped to some arbitrarily 10 chosen computational processor or replicated onto all computational processors. (End 11 of rationale.) 12

An HPF compiler is required to accept any PROCESSORS declaration in which the product of the extents of each declared processor arrangement is equal to the number of physical processors that would be returned by the call NUMBER\_OF\_PROCESSORS(). It must also accept all declarations of scalar PROCESSOR arrangements. Other cases may be handled as well, depending on the implementation.

For compatibility with the Fortran attribute syntax, an optional "::" may be inserted. The shape may also be specified with the **DIMENSION** attribute:

!HPF\$ PROCESSORS :: RUBIK(3,3,3)
!HPF\$ PROCESSORS, DIMENSION(3,3,3) :: RUBIK

As in Fortran, an *explicit-shape-spec-list* in a *processors-decl* will override an explicit **DIMENSION** attribute:

!HPF\$ PROCESSORS, DIMENSION(3,3,3) :: &
!HPF\$ RUBIK, RUBIKS\_REVENGE(4,4,4), SOMA

Here RUBIKS\_REVENCE is  $4 \times 4 \times 4$  while RUBIK and SOMA are each  $3 \times 3 \times 3$ . (By the rules enunciated above, however, such a statement may not be completely portable because no HPF language processor is required to handle shapes of total sizes 27 and 64 simultaneously.)

Returning from a subprogram causes all processor arrangements declared local to that subprogram to become undefined. It is not HPF-conforming for any array or template to be distributed onto a processor arrangement at the time the processor arrangement becomes undefined unless at least one of two conditions holds:

- The array or template itself becomes undefined at the same time by virtue of returning from the subprogram.
- Whenever the subprogram is called, the processor arrangement is always locally defined in the same way, with identical lower bounds and identical upper bounds.

Rationale. Note that this second condition is slightly less stringent than requiring all expressions to be constant. This allows calls to NUMBER\_OF\_PROCESSORS or PROCESSORS\_SHAPE to appear without violating the condition. (End of rationale.)

Variables in COMMON or having the SAVE attribute may be mapped to a locally declared 47 processor arrangement, but because the first condition cannot hold for such variables (they 48

don't become undefined), the second condition must be observed. This allows COMMON
 variables to work properly through the customary strategy of putting identical declarations
 in each scoping unit that needs to use them, while allowing the processor arrangements to
 which they may be mapped to depend on the value returned by NUMBER\_OF\_PROCESSORS.
 (See section 3.8 for further information on mapping common variables.)

Advice to implementors. It may be desirable to have a way for the user to specify at compile time the number of physical processors on which the program is to be executed. This might be specified either by an implementation-dependent directive, for example, or through the programming environment (for example, as a UNIX command-line argument). Such facilities are beyond the scope of the HPF specification, but as food for thought we offer the following illustrative hypothetical examples:

```
14!Declaration for multiprocessor by ABC Corporation15!ABC$ PHYSICAL PROCESSORS(8)16!Declaration for mpp by XYZ Incorporated17!XYZ$ PHYSICAL PROCESSORS(65536)
```

```
19!PDQ$ PHYSICAL PROCESSORS(2,2,2,2,2,2,2,2,2,2)20!Declaration for two-dimensional grid machine by TLA GmbH21!TLA$ PHYSICAL PROCESSORS(128,64)22!One of the preceding might affect the following:
```

!Declaration for hypercube machine by PDQ Limited

```
23 !HPF$ PROCESSORS P(NUMBER_OF_PROCESSORS())
```

It may furthermore be desirable to have a way for the user to specify the precise mapping of the processor arrangement declared in a **PROCESSORS** statement to the physical processors of the executing hardware. Again, this might be specified either by a implementation-dependent directive or through the programming environment (for example, as a UNIX command-line argument); such facilities are beyond the scope of the HPF specification, but as food for thought we offer the following illustrative hypothetical example:

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!PDQ$ PHYSICAL PROCESSORS(2,2,2,2,2,2,2,2,2,2,2,2,2)
!HPF$ PROCESSORS G(8,64,16)
!PDQ$ MACHINE LAYOUT G(:GRAY(0:2),:GRAY(6:11),:BINARY(3:5,12))
```

This might specify that the first dimension of G should use hypercube axes 0, 1, 2 with a Gray-code ordering; the second dimension should use hypercube axes 6 through 11 with a Gray-code ordering; and the third dimension should use hypercube axes 3, 4, 5, and 12 with a binary ordering. (End of advice to implementors.)

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## 3.7 The TEMPLATE Directive

The TEMPLATE directive declares one or more templates, specifying for each the name, the rank (number of dimensions), and the extent in each dimension. It must appear in the *specification-part* of a scoping unit.

In the language of F95:14.1.2 in the Fortran standard, templates are local entities of class (1); therefore a template may not have the same name as a variable, named constant,

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internal procedure, etc., in the same scoping unit. Template names obey the rules for host and use association as other names in the list in F95:12.1.2.2.1 in the Fortran standard.

A template declared in a module has the default accessibility of the module.

*Rationale.* Because the name of a template is not a first-class entity in HPF, but must appear only in directives, it cannot appear in an *access-stmt* (PRIVATE or PUBLIC). If directives ever become full-fledged Fortran statements rather than structured comments, then it would be appropriate to allow the accessibility of a template to be controlled by listing its name in an *access-stmt*. (End of rationale.)

A template is simply an abstract space of indexed positions; it can be considered as an "array of nothings" (as compared to an "array of integers," say). A template may be used as an abstract *align-target* that may then be distributed.

H331 template-directive	is	TEMPLATE template-decl-list	14
1	is	template-name [ ( explicit-shape-spec-list ) ]	15
H332 template-decl	IS	iemplaie-name [ ( explicit-shape-spec-list ) ]	16
			17
Examples:			18
Examples.			19
!HPF\$ TEMPLATE A(N)			20
!HPF\$ TEMPLATE B(N,N), C(N,2*N			21
<pre>!HPF\$ TEMPLATE DOPEY(100,100),</pre>	,SNE	EZY(24),GRUMPY(17,3,5)	22
If the "::" syntax is used, then the de	clare	ed templates may optionally be distributed in the	23
		emplates declared by the directive must have the	24 25
		te will be meaningful. The <b>DIMENSION</b> attribute	25 26
may also be used.		C C	20
			28
<pre>!HPF\$ TEMPLATE, DISTRIBUTE(BLC !HPF\$</pre>	JCK,	*) :: & WHINEY(64,64),MOPEY(128,128)	29
:HPF\$ TEMPLATE, DIMENSION(91,9	11)		30
INFIQ IEMPLATE, DIMENSION (91,3	,1)	DORED, WHEEZI, FERRI	31
Templates are useful in the parti	icula	r situation where one must align several arrays	32
		ed to declare a single array that spans the entire	33
		night want four $N \times N$ arrays aligned to the four	34
corners of a template of size $(N + 1)$ >	$\times (N)$	(+1):	35
!HPF\$ TEMPLATE, DISTRIBUTE(BLC	DCK.	BLOCK) :: EARTH(N+1.N+1)	36
REAL, DIMENSION(N,N) ::	-	-	37
HPF\$ ALIGN NW(I,J) WITH EARTH			38
HPF\$ ALIGN NE(I,J) WITH EARTH		-	39
!HPF\$ ALIGN SW(I,J) WITH EARTH		-	40
!HPF\$ ALIGN SE(I,J) WITH EARTH	H(I+	1,J+1)	41
			42
Templates may also be useful in makin	g as:	sertions about the mapping of dummy arguments	43

(see Section 4). Unlike arrays, templates cannot be in COMMON. So two templates declared in different 45scoping units will always be distinct, even if they are given the same name. The only way  $^{46}$ for two program units to refer to the same template is to declare the template in a module 47that is then used by the two program units. 48 Templates are not passed through the subprogram argument interface. The template to which a dummy argument is aligned is always distinct from the template to which the actual argument is aligned, though it may be a copy (see section 4.4.2). On exit from a subprogram, an HPF implementation arranges that the actual argument is aligned with the same template with which it was aligned before the call.

Returning from a subprogram causes all templates declared local to that subprogram to become undefined. It is not HPF-conforming for any variable to be aligned to a template at the time the template becomes undefined unless at least one of two conditions holds:

- The variable itself becomes undefined at the same time by virtue of returning from the subprogram.
- Whenever the subprogram is called, the template is always locally defined in the same way, with identical lower bounds, identical upper bounds, and identical distribution information (if any) onto identically defined processor arrangements (see section 3.6).

*Rationale.* Note that this second condition is slightly less stringent than requiring all expressions to be constant. This allows calls to NUMBER\_OF\_PROCESSORS or PROCESSORS\_SHAPE to appear without violating the condition. (*End of rationale.*)

Variables in COMMON or having the SAVE attribute may be mapped to a locally declared template, but because the first condition cannot hold for such variable (they don't become undefined), the second condition must be observed.

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## 3.8 Storage and Sequence Association

HPF allows the mapping of data objects across multiple processors in order to improve
 parallel performance. Fortran specifies relationships between the storage for data objects
 associated through COMMON and EQUIVALENCE statements, and the order of array elements
 during association at procedure boundaries between actual arguments and dummy arguments. Otherwise, the location of data is not constrained by the language.

COMMON and EQUIVALENCE statements constrain the alignment of different data items based on the underlying model of storage units and storage sequences:

> Storage association is the association of two or more data objects that occurs when two or more storage sequences share or are aligned with one or more storage units.

— Fortran Standard (F95:14.6.3.1)

The model of storage association is a single linearly addressed memory, based on the traditional single address space, single memory unit architecture. This model can cause severe inefficiencies on architectures where storage for data objects is mapped.

43 Sequence association refers to the order of array elements that Fortran requires when
 44 an array expression or array element is associated with a dummy array argument:

The rank and shape of the actual argument need not agree with the rank and shape of the dummy argument, ...

— Fortran Standard (F95:12.4.1.4)

As with storage association, sequence association is a natural concept only in systems with 1 a linearly addressed memory. 2 As an aid to porting FORTRAN 77 codes, HPF allows codes that rely on sequence and 3 storage association to be valid in HPF. Some modification to existing FORTRAN 77 codes 4 may nevertheless be necessary. This section explains the relationship between HPF data 5mapping and sequence and storage association. 6 7 **Storage Association** 3.8.18 9 3.8.1.1Definitions 10  $1\,1$ 1. COMMON blocks are either sequential or nonsequential, as determined by either explicit 12directive or compiler default. A sequential COMMON block has a single common block storage sequence (F95:5.5.2.1). 13 14 2. An aggregate variable group is a collection of variables whose individual storage se-15quences are parts of a single storage sequence. 16 Variables associated by EQUIVALENCE statements or by a combination of EQUIVALENCE 1718 and COMMON statements form an aggregate variable group. The variables of a sequential 19 COMMON block form a single aggregate variable group. 20 3. The *size* of an aggregate variable group is the number of storage units in the group's 21 storage sequence (F95:14.6.3.1). 22 23 4. Data objects are either sequential or nonsequential. A data object is sequential if and  $^{24}$ only if any of the following holds:  $^{25}$ 26 (a) it appears in a sequential COMMON block; 27 (b) it is a member of an aggregate variable group; 28 (c) it is an assumed-size array; 29 (d) its type is a sequence type; 30 31 (e) it is a subobject of a sequential data object; or 32 (f) it is declared to be sequential in an HPF SEQUENCE directive. 33 34 A sequential object can be storage associated or sequence associated; nonsequential 35 objects cannot. 36 37 5. A COMMON block contains a sequence of *components*. Each component is either an 38 aggregate variable group, or a variable that is not a member of any aggregate variable 39 group. A sequential COMMON block contains a single component. A nonsequential 40 COMMON block may contain several components each of which may be a sequential 41 variable, an aggregate variable group, or a nonsequential variable. 42 43 Examples of Definitions 3.8.1.244 !Example 1: 45 COMMON /FOO/ A(100), B(100), C(100), D(100), E(100) 46 DIMENSION X(100), Y(150), Z(200) 47 EQUIVALENCE (A(1), Z(1))48

```
!Four components: (A, B), C, D, E
1
       !Sizes are: 200, 100, 100, 100
2
3
       !Example 2:
4
             COMMON /FOO/ A(100), B(100), C(100), D(100), E(100)
5
             DIMENSION X(100), Y(150), Z(200)
6
             EQUIVALENCE ( A(51), X(1) ) ( B(100), Y(1) )
7
       !Two components (A, B, C, D), E
8
       !Sizes are: 400, 100
9
10
       !Example 3:
11
             COMMON /FOO/ A(100), B(100), C(100), D(100), E(100)
12
             DIMENSION X(100), Y(150), Z(200)
13
       !HPF$ SEQUENCE /FOO/
14
       !The COMMON has one component, (A, B, C, D, E)
15
16
       !Size is 500
17
```

The COMMON block /FOO/ is nonsequential in Examples 1 and 2. Aggregate variable groups are shown as components in parentheses.

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## 3.8.2 The SEQUENCE Directive

22 A SEQUENCE directive is defined to allow a user to declare explicitly that data objects or 23 COMMON blocks are to be treated by the compiler as sequential. (COMMON blocks are by 24 default nonsequential. Data objects are nonsequential unless Definition 4 of Section 3.8  $^{25}$ applies.) Some implementations may supply an optional compilation environment where 26 the **SEQUENCE** directive is applied by default. For completeness in such an environment, HPF 27 defines a NO SEQUENCE directive to allow a user to establish that the usual nonsequential 28 default should apply to a scoping unit or to selected data objects and COMMON blocks within 29 the scoping unit. 30

31 32 33	H333	sequence- $directive$	SEQUENCE [ [ :: ] association-name-list ] NO SEQUENCE [ [ :: ] association-name-list ]
34 35	H334	association-name	object-name / [ common-block-name ] /
36			

Constraint: An object name or COMMON block name may appear at most once in a *sequence-directive* within any scoping unit.

Constraint: Only one sequence directive with no association-name-list is permitted in the same scoping unit.

A sequential pointer can be associated only with sequential objects. A nonsequential pointer can be associated only with nonsequential objects.

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## 3.8.2.1 Storage Association Rules

 A sequence-directive with an empty association-name-list is treated as if it contained the names of all implicitly mapped objects and COMMON blocks in the scoping unit that

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cannot otherwise be determined to be sequential or nonsequential by their language context.

- 2. A sequential object may not be explicitly mapped.
- 3. No explicit mapping may be given for a component of a derived type having the Fortran SEQUENCE attribute. Note that this rule is applicable only under the approved extensions since components of derived types cannot be explicitly mapped in HPF.
- 4. If a COMMON block is nonsequential, then all of the following must hold:
  - (a) Every occurrence of the COMMON block has exactly the same number of components with each corresponding component having a storage sequence of exactly the same size;
  - (b) If a component is a nonsequential variable in *any* occurrence of the COMMON block, then it must be nonsequential with identical type, shape, and mapping attributes in *every* occurrence of the COMMON block; and
  - (c) Every occurrence of the COMMON block must be nonsequential.

#### 3.8.2.2 Storage Association Discussion

Advice to users. Under these rules, variables in a COMMON block can be mapped as long as the components of the COMMON block are the same in every scoping unit that declares the COMMON block.

Correct Fortran programs will not necessarily be correct without modification in HPF. The use of EQUIVALENCE with COMMON blocks can impact the mappability of data objects in subtle ways. To allow maximum optimization for performance, the HPF default for data objects is to consider them mappable. In order to get correct separate compilation for subprograms that use COMMON blocks with different aggregate variable groups in different scoping units, it will be necessary to insert the HPF SEQUENCE directive. 30

As a check-list for a user to determine the status of a data object or COMMON block, the following questions can be applied, in order:

- Does the object appear in some explicit language context which dictates that the object be sequential (e.g. EQUIVALENCE) or nonsequential?
- If not, does the object appear in an explicit mapping directive?
- If not, does the object or COMMON block name appear in the list of names on a SEQUENCE or NO SEQUENCE directive?
- If not, does the scoping unit contain a nameless SEQUENCE or NO SEQUENCE?
- If not, is the compilation affected by some special implementation-dependent environment which dictates that names default to SEQUENCE?
- If not, then the compiler will consider the object or COMMON block name nonsequential and is free to apply data mapping optimizations that disregard Fortran sequence and storage association.

(End of advice to users.)

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Advice to implementors. In order to protect the user and to facilitate portability of older codes, two implementation options are strongly recommended. First, every implementation should supply some mechanism to verify that the type and shape of every mappable array and the sizes of aggregate variable groups in COMMON blocks are the same in every scoping unit unless the COMMON blocks are declared to be sequential. This same check should also verify that identical mappings have been selected for the variables in COMMON blocks. Implementations without interprocedural information can use a link-time check. The second implementation option recommended is a mechanism to declare that data objects and COMMON blocks for a given compilation should be considered sequential unless declared otherwise. The purpose of this feature is to permit compilation of large old libraries or subprograms where storage association is known to exist without requiring that the code be modified to apply the HPF SEQUENCE directive to every COMMON block. (End of advice to implementors.)

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## Section 4

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# Data Mapping in Subprogram Interfaces

In this Section, phrases such as "the caller must pass..." are constraints on the implementation (i.e., on the generated code produced by the compiler), not on the source code produced by the programmer.

#### 4.1 Introduction

This introduction gives an overview of the ways in which mapping directives interact with argument passing to subprograms. The language used here, however, is not definitive; the subsequent subsections of this Section contain the authoritative rules.

In addition to the data mapping features described in Section 3, HPF allows a number of options for describing the mapping of dummy arguments.

The mapping of each such dummy argument may be related to the mapping of its associated actual argument in the calling main program or procedure (the "caller") in several different ways. To allow for this, mapping directives applied to dummy arguments can have three different syntactic forms: *prescriptive*, *descriptive*, and *transcriptive*.

HPF provides these three forms to allow the programmer either to specify that the data is to be left in place, or to specify that during the execution of the call the data must be automatically remapped into a new and presumably more efficient mapping for the duration of the execution of the called subprogram.

The meaning of these forms is as follows:

**prescriptive** The directive describes the mapping of the dummy argument. However, the actual argument need not have this mapping. *If it does not*, it is the responsibility of the compiler to generate code to remap the argument as specified, and to restore the original mapping on exit. This code may be generated either in the caller or in the called subprogram; the requirements for explicit interfaces in Section 4.6 insure that the necessary information will be available at compile time to perform the mapping in either place.

Prescriptive directives are syntactically identical to directives occurring elsewhere in the program. For instance, if **A** is a dummy argument,

HPF\$ DISTRIBUTE A (BLOCK, CYCLIC)

is a prescriptive directive.

**descriptive** Descriptive syntax has exactly the same meaning as prescriptive syntax, except that in addition it amounts to a weak assertion by the programmer that the actual argument requires no remapping.

The assertion is characterized as "weak" because if it is false, the program is still standard-conforming. In such a case, the compiler must generate the appropriate remapping.

If the compiler can prove that the assertion is false, or if the compiler cannot verify that it is true, it may issue a warning or informational diagnostic message.

Advice to users. The purpose of descriptive, as opposed to prescriptive, directives is simply to provide a possible way for the compiler to report information to the programmer that may be useful in program development and debugging. Note that any diagnostic message that may be produced as a result of the use of descriptive directives is not a portable feature of this language. In particular, there are instances in which no remapping is needed but where this fact would be impossible or highly non-trivial for a compiler to ascertain. Different compilers may well emit messages in different circumstances; and there is no requirement that any such messages be emitted at all. (*End of advice to users.*)

Descriptive directives look like prescriptive directives, except that an asterisk precedes the description. For instance,

#### !HPF\$ DISTRIBUTE A \*(BLOCK, CYCLIC)

is a descriptive directive.

transcriptive The mapping is unspecified. The called subprogram must accept the mapping of the argument as it is passed. Of course this means that (the implementation of) the caller must pass this mapping information at run-time.

Transcriptive directives are written with a single asterisk for distributions and processor arrangements; for instance

!HPF\$ DISTRIBUTE A \*
!HPF\$ DISTRIBUTE B \* ONTO \*

are transcriptive directives. The INHERIT directive (see Section 4.4.2) is used to specify a transcriptive alignment.

Both distribution formats and processor arrangements can be specified prescriptively, descriptively, or transcriptively. Alignment is more complicated, because of the need to specify the template with which the dummy is aligned. This template may be unspecified (in this case of course there is no ALIGN directive), in which case it is the *natural template* of the dummy. ("Natural template" is defined in Section 4.4.1 below.) Otherwise, one of the following disjoint possibilities must be true:

• The template is explicitly specified by a prescriptive ALIGN directive.

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• The template is explicitly specified by a descriptive ALIGN directive.

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	• The template is <i>inherited</i> . This is specified by giving the dummy the INHERIT attribute
3	(described in Section 4.4.2 below). This implicitly specifies the template to be a copy
4	,
5	of the template with which the corresponding actual argument is ultimately aligned;
6	further, the alignment of the dummy with that template is the same as that of the
7	corresponding actual. This is, in effect, a transcriptive form of alignment.
8	
9	This is restated more precisely in Section 4.4.1 below.
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11	Advice to users. Although it is possible to write some combinations of mapping
12	directives that are partially prescriptive and partially transcriptive, for instance, there
13	is probably no virtue in doing so. The point of these directives is to enable the
14	compiler to handle any necessary remapping correctly and efficiently. Now remapping
15	can happen for one or more of the following reasons:
16	can happen for one of more of the fonoting reasons.
17	• to make the alignment of the actual and the dummy agree;
18	
19	• to make the distribution of the actual and the dummy agree;
20	ullet to make the processor arrangement of the actual and the dummy agree.
21	
22	For most machines, there is no real difference in the cost of remapping for any of these
23	reasons. It is therefore a better practice (for readability, at least) to make a mapping
24	either purely transcriptive, purely prescriptive, or purely descriptive.
25	While transprintive mappings can be useful in writing libraries, they impose a run
26	While transcriptive mappings can be useful in writing libraries, they impose a run-
	time cost on the subprogram. They should therefore be avoided in normal user code. $(End ef nduise to use no)$
27	(End of advice to users.)
28	
29	4.2 What Remapping Is Required and Who Does It
30	
0.1	
	If there is an explicit interface for the called subprogram and that interface contains pre-
32	
32 33	scriptive or descriptive mapping directives for a dummy argument, and if a remapping of
32 33	scriptive or descriptive mapping directives for a dummy argument, and if a remapping of the corresponding actual argument is necessary, the call should proceed as if the data were
32 33 34	scriptive or descriptive mapping directives for a dummy argument, and if a remapping of the corresponding actual argument is necessary, the call should proceed as if the data were copied to a temporary variable to match the mapping of the dummy argument as expressed
32 33 34 35	scriptive or descriptive mapping directives for a dummy argument, and if a remapping of the corresponding actual argument is necessary, the call should proceed as if the data were copied to a temporary variable to match the mapping of the dummy argument as expressed by the directives in the explicit interface. The template of the dummy will then be as
32 33 34 35 36	scriptive or descriptive mapping directives for a dummy argument, and if a remapping of the corresponding actual argument is necessary, the call should proceed as if the data were copied to a temporary variable to match the mapping of the dummy argument as expressed by the directives in the explicit interface. The template of the dummy will then be as declared in the interface.
31 32 33 34 35 36 37 38	<ul> <li>scriptive or descriptive mapping directives for a dummy argument, and if a remapping of the corresponding actual argument is necessary, the call should proceed as if the data were copied to a temporary variable to match the mapping of the dummy argument as expressed by the directives in the explicit interface. The template of the dummy will then be as declared in the interface.</li> <li>If there is no explicit interface, then no remapping will be necessary; this is a conse-</li> </ul>
32 33 34 35 36 37	<ul> <li>scriptive or descriptive mapping directives for a dummy argument, and if a remapping of the corresponding actual argument is necessary, the call should proceed as if the data were copied to a temporary variable to match the mapping of the dummy argument as expressed by the directives in the explicit interface. The template of the dummy will then be as declared in the interface.</li> <li>If there is no explicit interface, then no remapping will be necessary; this is a consequence of the requirements in Section 4.6.</li> </ul>
32 33 34 35 36 37 38	<ul> <li>scriptive or descriptive mapping directives for a dummy argument, and if a remapping of the corresponding actual argument is necessary, the call should proceed as if the data were copied to a temporary variable to match the mapping of the dummy argument as expressed by the directives in the explicit interface. The template of the dummy will then be as declared in the interface.</li> <li>If there is no explicit interface, then no remapping will be necessary; this is a consequence of the requirements in Section 4.6.</li> <li>The reader should note that for reasons of brevity, not all such required explicit inter-</li> </ul>
32 33 34 35 36 37 38 39	<ul> <li>scriptive or descriptive mapping directives for a dummy argument, and if a remapping of the corresponding actual argument is necessary, the call should proceed as if the data were copied to a temporary variable to match the mapping of the dummy argument as expressed by the directives in the explicit interface. The template of the dummy will then be as declared in the interface.</li> <li>If there is no explicit interface, then no remapping will be necessary; this is a consequence of the requirements in Section 4.6.</li> <li>The reader should note that for reasons of brevity, not all such required explicit interfaces are included in the code fragments in this Section.</li> </ul>
32 33 34 35 36 37 38 39 40 41	<ul> <li>scriptive or descriptive mapping directives for a dummy argument, and if a remapping of the corresponding actual argument is necessary, the call should proceed as if the data were copied to a temporary variable to match the mapping of the dummy argument as expressed by the directives in the explicit interface. The template of the dummy will then be as declared in the interface.</li> <li>If there is no explicit interface, then no remapping will be necessary; this is a consequence of the requirements in Section 4.6.</li> <li>The reader should note that for reasons of brevity, not all such required explicit interfaces are included in the code fragments in this Section.</li> <li>An overriding principle is that any remapping of arguments is not visible to the caller.</li> </ul>
32 33 34 35 36 37 38 39 40 41 41	<ul> <li>scriptive or descriptive mapping directives for a dummy argument, and if a remapping of the corresponding actual argument is necessary, the call should proceed as if the data were copied to a temporary variable to match the mapping of the dummy argument as expressed by the directives in the explicit interface. The template of the dummy will then be as declared in the interface.</li> <li>If there is no explicit interface, then no remapping will be necessary; this is a consequence of the requirements in Section 4.6.</li> <li>The reader should note that for reasons of brevity, not all such required explicit interfaces are included in the code fragments in this Section.</li> <li>An overriding principle is that any remapping of arguments is not visible to the caller.</li> </ul>
32 33 34 35 36 37 38 39 40	<ul> <li>scriptive or descriptive mapping directives for a dummy argument, and if a remapping of the corresponding actual argument is necessary, the call should proceed as if the data were copied to a temporary variable to match the mapping of the dummy argument as expressed by the directives in the explicit interface. The template of the dummy will then be as declared in the interface.</li> <li>If there is no explicit interface, then no remapping will be necessary; this is a consequence of the requirements in Section 4.6.</li> <li>The reader should note that for reasons of brevity, not all such required explicit interfaces are included in the code fragments in this Section.</li> <li>An overriding principle is that any remapping of arguments is not visible to the caller.</li> <li>That is, when the subprogram returns and the caller resumes execution, all objects accessible to the caller after the call are mapped exactly as they were before the call. It is not possible</li> </ul>
32 33 34 35 36 37 38 39 40 41 41 42	<ul> <li>scriptive or descriptive mapping directives for a dummy argument, and if a remapping of the corresponding actual argument is necessary, the call should proceed as if the data were copied to a temporary variable to match the mapping of the dummy argument as expressed by the directives in the explicit interface. The template of the dummy will then be as declared in the interface.</li> <li>If there is no explicit interface, then no remapping will be necessary; this is a consequence of the requirements in Section 4.6.</li> <li>The reader should note that for reasons of brevity, not all such required explicit interfaces are included in the code fragments in this Section.</li> <li>An overriding principle is that any remapping of arguments is not visible to the caller.</li> </ul>
32 33 34 35 36 37 38 39 40 41 41 42 43 44	<ul> <li>scriptive or descriptive mapping directives for a dummy argument, and if a remapping of the corresponding actual argument is necessary, the call should proceed as if the data were copied to a temporary variable to match the mapping of the dummy argument as expressed by the directives in the explicit interface. The template of the dummy will then be as declared in the interface.</li> <li>If there is no explicit interface, then no remapping will be necessary; this is a consequence of the requirements in Section 4.6.</li> <li>The reader should note that for reasons of brevity, not all such required explicit interfaces are included in the code fragments in this Section.</li> <li>An overriding principle is that any remapping of arguments is not visible to the caller.</li> <li>That is, when the subprogram returns and the caller resumes execution, all objects accessible to the caller after the call are mapped exactly as they were before the call. It is not possible</li> </ul>
32 33 34 35 36 37 38 39 40 41 42 43 44 45	<ul> <li>scriptive or descriptive mapping directives for a dummy argument, and if a remapping of the corresponding actual argument is necessary, the call should proceed as if the data were copied to a temporary variable to match the mapping of the dummy argument as expressed by the directives in the explicit interface. The template of the dummy will then be as declared in the interface.</li> <li>If there is no explicit interface, then no remapping will be necessary; this is a consequence of the requirements in Section 4.6.</li> <li>The reader should note that for reasons of brevity, not all such required explicit interfaces are included in the code fragments in this Section.</li> <li>An overriding principle is that any remapping of arguments is not visible to the caller.</li> <li>That is, when the subprogram returns and the caller resumes execution, all objects accessible to the caller after the call are mapped exactly as they were before the call. It is not possible</li> </ul>

#### 4.3 Distributions and Processor Arrangements

In a DISTRIBUTE directive where every *distributee* is a dummy argument, either the *dist-format-clause* or the *dist-target*, or both, may begin with, or consist of, an asterisk.

- Without an asterisk, a *dist-format-clause* or *dist-target* is prescriptive; the clause describes a distribution and constitutes a request of the language processor to make it so. This might require (the implementation of) either the caller or the called subprogram to remap or copy the actual argument on entry at run time in order to satisfy the requested distribution for the dummy.
- Starting with an asterisk, a *dist-format-clause* or *dist-target* is descriptive. Such a directive is equivalent in every respect to a prescriptive directive, except that if the compiler cannot verify that no remapping of the actual is required, it may issue a diagnostic message to that effect. See Section 4.1 for further information on this point.
- Consisting of only an asterisk, a *dist-format-clause* or *dist-target* is transcriptive; the clause says nothing about the distribution but constitutes a request to the language processor to copy that aspect of the distribution from that of the actual argument. (The intent is that if the argument is passed by reference, no movement of the data will be necessary at run time.)

It is possible that, in a single DISTRIBUTE directive, the *dist-format-clause* might have an asterisk but not the *dist-target*, or vice versa.

#### 4.3.1 Examples

These examples of **DISTRIBUTE** directives for dummy arguments illustrate the various combinations:

#### !HPF\$ DISTRIBUTE URANIA (CYCLIC) ONTO GALILEO

The language processor should do whatever it takes to cause URANIA to have a CYCLIC distribution on the processor arrangement GALILEO.

#### !HPF\$ DISTRIBUTE POLYHYMNIA \* ONTO ELVIS

The language processor should do whatever it takes to cause POLYHYMNIA to be distributed onto the processor arrangement ELVIS, using whatever distribution format it currently has (which might be on some other processor arrangement).

#### !HPF\$ DISTRIBUTE THALIA \*(CYCLIC) ONTO \*FLIP

The language processor should do whatever it takes to cause THALIA to have a CYCLIC distribution on the processor arrangement FLIP; the programmer believes that the actual is already distributed in this fashion and that no remapping is required.

!HPF\$ DISTRIBUTE EUTERPE (CYCLIC) ONTO \*

The language processor should do whatever it takes to cause EUTERPE to have a CYCLIC distribution onto whatever processor arrangement the actual was distributed onto.

#### !HPF\$ DISTRIBUTE ERATO \* ONTO \* 2 The mapping of **ERATO** should not be changed from that of the actual argument. З Note that **DISTRIBUTE ERATO \* ONTO \*** does not mean the same thing as 4 5 !HPF\$ DISTRIBUTE ERATO (\*) ONTO \* 6 This latter means: distribute ERATO \* (that is, on-processor) onto whatever processor ar-7 rangement the actual was distributed onto. The processor arrangement is necessarily scalar 8 in this case. 9 10 What Happens When a Clause Is Omitted 11 4.3.212One may omit either the *dist-format-clause* or the *dist-onto-clause* for a dummy argument. 13 This is understood as follows: 14 If the dummy argument has the INHERIT attribute (see Section 4.4.2), then no distri-15bution directive is allowed in any case: the distribution as well as the alignment is inherited 16 from the actual argument. 17In any other case in which distribution information is omitted, the compiler may choose 18 the distribution format or a target processor arrangement arbitrarily. 19 Here are two examples: 20 21 !HPF\$ DISTRIBUTE WHEEL\_OF\_FORTUNE \*(CYCLIC) 22 The programmer believes that the actual argument corresponding to the dummy argument 23 WHEEL\_OF\_FORTUNE is already distributed CYCLIC. The compiler should insure that the map-24 ping of the passed data is in fact CYCLIC, and remap it if necessary if it is not. It may in $^{25}$ addition be remapped onto some other processor arrangement, but there is no reason to; 26 most likely the programmer would be surprised if such a remapping occurred. 27 28 !HPF\$ DISTRIBUTE ONTO \*TV :: DAVID\_LETTERMAN 29 30 The programmer believes that the actual argument corresponding to the dummy argument 31 DAVID\_LETTERMAN is already distributed onto TV in some fashion. The compiler should 32 insure that this is so, and make it so if it is not. The distribution format may be changed 33 as long as DAVID\_LETTERMAN is kept on TV. (Note that this declaration must be made in 34 attributed form; the statement form 35 !HPF\$ DISTRIBUTE DAVID LETTERMAN ONTO \*TV !Nonconforming 36 37 does not conform to the syntax for a **DISTRIBUTE** directive.) 38 39 Alignment 4.4 40 41The Template of the Dummy Argument 4.4.142 Here we describe precisely how to determine the template with which the dummy argument 43 is ultimately aligned: 44

Templates are not passed through the subprogram argument interface. A dummy 45 argument and its corresponding actual argument may be aligned to the same template only 46 if that template is accessible in both the caller and the called subprogram either through 47host association or use association. In any other case, the template with which a dummy 48

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argument is aligned is always distinct from the template with which the actual argument 1 is aligned, though it may be a copy (see Section 4.4.2). On exit from a procedure, an HPF 2 implementation arranges that the actual argument is aligned with the same template with which it was aligned before the call.

The template of the dummy argument is arrived at in one of three ways:

- If the dummy argument appears explicitly as an *alignee* in an ALIGN directive, its template is the *align-target* if the *align-target* is a template; otherwise its template is the template with which the *align-target* is ultimately aligned.
- If the dummy argument is not explicitly aligned and does not have the INHERIT attribute (described in Section 4.4.2 below), then the template has the same shape and bounds as the dummy argument; this is called the *natural template* for the dummy.

(Thus, all the examples in Section 4.3 use the natural template.)

- If the dummy argument is not explicitly aligned and does have the INHERIT attribute, then the template is "inherited" from the actual argument according to the following rules:
  - If the actual argument is a whole array, the template of the dummy is a copy of the template with which the actual argument is ultimately aligned.
  - If the actual argument is an array section of array A where no subscript is a vector subscript, then the template of the dummy is a copy of the template with which A is ultimately aligned.
  - If the actual argument is any other expression, the shape and distribution of the template may be chosen arbitrarily by the language processor (and therefore the programmer cannot know anything *a priori* about its distribution).

In all of these cases, we say that the dummy has an *inherited template*.

#### The INHERIT Directive 4.4.2

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The INHERIT directive specifies that a dummy argument should be aligned to a copy of the template of the corresponding actual argument in the same way that the actual argument is aligned.

				0.
H401	inherit- $directive$	is	INHERIT inheritee-list	38
H402	inheritee	is	object-name	39
			0	40
Consti	raint: An <i>inheritee</i> must be a d	umn	nv argument.	41
-				42
Constr	raint: An <i>inheritee</i> must not be	an	alignee.	43
				44
Consti	raint: An <i>inheritee</i> must not be	a d	istributee.	43
				46
	Advice to years The first of the	co t	broa constraints is relayed for pointars under the	4.5

Advice to users. The first of these three constraints is relaxed for pointers under the approved extensions (see Section 8.8). (End of advice to users.)

The INHERIT directive causes the named subprogram dummy arguments to have the INHERIT attribute. Only dummy arguments may have the INHERIT attribute. An object must not have both the INHERIT attribute and the ALIGN attribute. An object must not have both the INHERIT attribute and the DISTRIBUTE attribute. The INHERIT directive may appear only in a *specification-part* of a scoping unit.

The INHERIT attribute specifies that the template for a dummy argument should be 6 inherited, by making a copy of the template of the actual argument. Moreover, no other 7 explicit mapping directive may appear for an argument with the INHERIT attribute: the 8 INHERIT attribute implies a distribution of DISTRIBUTE \* ONTO \* for the inherited tem-9 plate. Thus, the net effect is to tell the compiler to leave the data exactly where it is, 10 11 and not attempt to remap the actual argument. The dummy argument will be mapped in exactly the same manner as the actual argument; the subprogram must be compiled in 12such a way as to work correctly no matter how the actual argument may be mapped onto 13 abstract processors. 14

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Note that if A is an array dummy argument, the directive

17 **!HPF\$ INHERIT A** 

is more general than

```
<sup>20</sup> !HPF$ DISTRIBUTE A * ONTO *
```

for the following reason: The INHERIT directive states that the (inherited) template with which A is aligned is distributed \* ONTO \*, but that A may be aligned in some non-trivial manner with that template. On the other hand, the DISTRIBUTE directive states that A is aligned trivially with its natural template, which in turn is distributed \* ONTO \*.

For example, the following code is not permitted:

```
27
28
        !HPF$ PROCESSORS P(2)
29
              REAL, DIMENSION(100) :: A
30
        !HPF$ DISTRIBUTE (BLOCK) ONTO P :: A
31
32
              CALL FOO(A(1:50))
33
34
              . . .
35
36
              SUBROUTINE FOO(D)
37
              REAL, DIMENSION(50) :: D
38
       !HPF$ DISTRIBUTE D *
                                            ! Nonconforming
39
```

The transcriptive distribution for D is nonconforming because the natural template for D is
 not distributed BLOCK. On the other hand, it would be correct to replace the illegal directive
 by

44 !HPF\$ INHERIT D

45

47

43

<sup>46</sup> 4.4.2.1 Examples

<sup>48</sup> Here is a straightforward example of the use of INHERIT:

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 $1\,1$ 

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```
REAL DOUGH(100)

!HPF$ DISTRIBUTE DOUGH(BLOCK(10))

CALL PROBATE( DOUGH(7:23:2) )

...

SUBROUTINE PROBATE(BREAD)

REAL BREAD(9)

!HPF$ INHERIT BREAD
```

The inherited template of BREAD has shape [100]; element BREAD(I) is aligned with element 5 + 2\*I of the inherited template, and that template has a BLOCK(10) distribution.

More complicated examples can easily be constructed. It is important to bear in mind that the rank of the inherited template may be different from the rank of the dummy, and it might even be different from the rank of the actual. For instance, one might have a program containing the following:

```
REAL A(100,100)

!HPF$ TEMPLATE T(100,100,100)

!HPF$ DISTRIBUTE T(BLOCK,CYCLIC,*)

!HPF$ ALIGN A(I,J) with T(J,3,I)

CALL SUBR(A(:,7))

...

SUBROUTINE SUBR(D)

REAL D(100)

!HPF$ INHERIT D
```

In this case, the dummy D has rank 1. It corresponds to a 1-dimensional section of a 2-dimensional actual A, which in turn is aligned with a 2-dimensional section of a 3dimensional template T. The template of D is a copy of this three-dimensional template. D is aligned with the section (7, 3, :) of this inherited template. Thus, the "visible" dimension of the dummy D is distributed \*, although if the call statement had been

```
CALL SUBR(A(7,:))
```

for instance, the "visible" dimension of the dummy would be distributed BLOCK.

#### 4.4.3 Descriptive ALIGN Directives

The presence or absence of an asterisk at the start of an *align-spec* has the same meaning as in a *dist-format-clause*: it specifies whether the ALIGN directive is descriptive or prescriptive, respectively.

If an *align-spec* that does not begin with \* is applied to a dummy argument, the meaning is that the dummy argument will be forced to have the specified alignment on entry to the subprogram. This may require (the implementation of) either the caller or the subprogram to temporarily remap the data of the actual argument or a copy thereof.

Note that a dummy argument may also be used as an *align-target*.

SUBROUTINE NICHOLAS(TSAR,CZAR) REAL, DIMENSION(1918) :: TSAR,CZAR !HPF\$ INHERIT :: TSAR !HPF\$ ALIGN WITH TSAR :: CZAR

In this example the first dummy argument, TSAR, remains aligned with the correspond-1 ing actual argument, while the second dummy argument, CZAR, is forced to be aligned with 2 the first dummy argument. If the two actual arguments are already aligned, no remapping 3 of the data will be required at run time. If they are not, some remapping will take place. 4

If the *align-spec* begins with "\*", then the *alignee* must be a dummy argument. The "\*" 5 indicates that the programmer believes that the actual argument already has the specified 6 alignment, and that no action to remap it is required at run time. (As before, there is 7 no requirement that the programmer's belief is correct, and the compiler must generate a 8 remapping if one appears to be necessary, just as in the case of a prescriptive alignment.) 9 For example, if in the above example the alignment directive were changed to 10

```
11
12
```

13

```
!HPF$ ALIGN WITH *TSAR :: CZAR
```

LOGICAL FRUG(128)

!HPF\$ PROCESSORS DANCE\_FLOOR(16)

!HPF\$ DISTRIBUTE (BLOCK) ONTO DANCE\_FLOOR::FRUG

then the programmer is expressing a belief that no remapping of the actual argument 14 corresponding to TSAR will be necessary. 15

It is not permitted to say simply "ALIGN WITH \*"; an *align-target* must follow the 16 asterisk. (The proper way to say "accept any alignment" is INHERIT.) 17

If a dummy argument has no explicit ALIGN or DISTRIBUTE attribute, then the compiler 18 provides an implicit alignment and distribution specification, one that could have been 19 described explicitly without any "assertion asterisks". 20

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#### 4.4.3.1Example

If the INHERIT directive is not used, explicit alignment of a dummy argument may be necessary to insure that no remapping takes place at the subprogram boundary. Here is an example:

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CALL TERPSICHORE(FRUG(1:40:3)) The array section FRUG(1:40:3) is mapped onto abstract processors in the following manner:

1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16
1			25												
	10			34											
		19													
4			28												
	13			37											
		22													
7			31												
	16			40											

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Suppose first that the interface to the subroutine TERPSICHORE looks like this:

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 $^{25}$ 

SUBROUTINE TERPSICHORE(FOXTROT) LOGICAL FOXTROT(:) !HPF\$ INHERIT FOXTROT

The template of FOXTROT is a copy of the 128 element template of the whole array FRUG. The template is mapped like this:

1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16
1	9	17	25	33	41	49	57	65	73	81	89	97	105	113	121
2	10	18	26	34	42	50	58	66	74	82	90	98	106	114	122
3	11	19	27	35	43	51	59	67	75	83	91	99	107	115	123
4	12	20	28	36	44	52	60	68	76	84	92	100	108	116	124
5	13	21	29	37	45	53	61	69	77	85	93	101	109	117	125
6	14	22	30	38	46	54	62	70	78	86	94	102	110	118	126
7	15	23	31	39	47	55	63	71	79	87	95	103	111	119	127
8	16	24	32	40	48	56	64	72	80	88	96	104	112	120	128

FOXTROT(I) is aligned with element 3\*I-2 of the template.

Suppose, on the other hand, that the interface to  ${\tt TERPSICHORE}$  were to look like this instead:

#### SUBROUTINE TERPSICHORE(FOXTROT) LOGICAL FOXTROT(:) !HPF\$ DISTRIBUTE FOXTROT(BLOCK)

In this case, the template of FOXTROT is its natural template; it has the same size 14 as FOXTROT itself. The actual argument, FRUG(1:40:3) is mapped to the 16 processors in this manner:

Abstract	Elements
processor	of FRUG
1	1,2,3
2	4,5,6
3	7, 8
4	$9,\ 10,\ 11$
5	12,13,14
6 - 16	none

That is, the original positions (in the template of the actual argument) of the elements of the dummy are as follows:

1		1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16
2		1			9												
3			4			12											
4			-	7		12											
5 6				1													
7		2			10												
8			5			13											
9				8													
10		3			11												
11		-	6			14											
12 13			0			14											
14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31	wil to	rth, . l take Ren a dec. PHPF\$ PHPF\$ HPF\$ <i>Ac</i> it ; mo	e place nappi lared SUB LOG PRO TEM ALIC <i>lvice t</i> gives pre eff	canno e at t ng ca temp ROUTI ICAL CESSC PLATE GN FC the co ficient	t pro he cal n be late o INE Ti FOXT DRS D. C, DI DXTRO	perly ll. avoid f size ERPSI ROT( ANCE_ STRIE T(I) The a er mo	be d led w 128 CHOR :) FLOO BUTE( WITH dvant re inf <i>id of</i>	escrib ithou distri E (FO R (16 BLOC GUR age o orma <i>advic</i>	t usir buted XTRO: ) K) OI F(3*1 f this tion; e to i	s a BI ng IN BLOC F) Into D [-2) latte this in users.	.DCK ( HERIT CK: ANCE_ r tech nform	listrik by e FLOO nique ation	nution xplici R::G is th can o	uRF(1 at, w	erefo lignin 228) here i be use	re, re: g the t can	3 on the mapping dummy be used, generate
32 33 34 35 36 37 38 39 40 41 42 43 44 45 46	equ <i>spe</i> spe	cializ ecified ow in Ac if : gen Ex of The	nce. ation Sect lvice you al nerall ctensic advic	Roug of <i>P</i> that ion 4. to use lways y hap on, ex <i>e</i> to <i>v</i> ise de	hly spectrum $Q$ (i.e. $Q$ is $c$ $6$ , and $ers$ . Spectrum $provious$	peaki , "Q one of d also Since ide ex autor ed be ) on is	ng, if is be f the o in S these cplicit natic: low in as fol	<i>P</i> a clow	nd Q P" in ings 1 8.8. itions faces and if ion 8	are this that i s are (which you .8), th	two n ordet s cons comp ch, as don't hen ye	nappi ring) sisten lex to expla use m ou nee	ngs, is to t with state ained appe ed no	then say h P. e, it is below d poi t read	to say that This wort v, is c nters this	y tha P is notion th not juite o (an A	a certain t $Q$ is a partially n is used ting that easy and pproved on. ( <i>End</i>
47 48					tation		-				-		<u> </u>	-			

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	1
	2
$f BLOCK \equiv f BLOCK$ $CYCLIC \equiv CYCLIC$	3
$\begin{array}{rcl} \texttt{CYCLIC} &\equiv & \texttt{CYCLIC} \\ & * &\equiv & * \end{array}$	4
T = T BLOCK(n) = BLOCK(m) iff m and n have the same value	5
$CYCLIC(n) \equiv CYCLIC(m)$ iff m and n have the same value	6
$CYCLIC \equiv CYCLIC(1)$	7
	8
2. Other than this, no two lexically distinct <i>dist-format</i> specifications are equivalent.	9 10
This is an equivalence relation in the usual mathematical sense.	11
Now we define the partial order on mappings: Let S ("special") and G ("general") be	12
two data objects.	13 14
The mapping of <b>S</b> is a <i>specialization</i> of the mapping of <b>G</b> if and only if either:	15
	16
1. G has the INHERIT attribute, or	17
2. S does not have the INHERIT attribute, and the following constraints all hold:	18
2. 5 does not have the initial attribute, and the following constraints an hold.	19
(a) S is a named object, and	20
(b) The shapes of the ultimate align targets of S and G are the same, and	21
	22
(c) Corresponding dimensions of $S$ and $G$ are mapped to corresponding dimensions	23
of their respective ultimate align targets, and corresponding elements of S and G are aligned with corresponding elements of their respective ultimate align targets,	24
and	25 26
	20
(d) Either	28
i. The ultimate align targets of both ${\tt S}$ and ${\tt G}$ are not explicitly distributed, or	29
ii. The ultimate align targets of both ${\tt S}$ and ${\tt G}$ are explicitly distributed. In this	30
case, the distribution directive specified for the ultimate align target of ${\tt G}$	31
must satisfy one of the following conditions:	32
A. It has no <i>dist-onto-clause</i> , or	33
B. It has a <i>dist-onto-clause</i> of "ONTO *", or	34
C. It has a <i>dist-onto-clause</i> specifying a processor arrangement having the	35
same shape as that explicitly specified in a distribution directive for the	36
ultimate align target of S.	37
and must also satisfy one of the following conditions:	38 39
A. It has no <i>dist-format-clause</i> , or	40
B. It has a <i>dist-format-clause</i> of "*", or	41
C. Each <i>dist-format</i> is equivalent (in the sense defined above) to the <i>dist-</i>	42
format in the corresponding position of the dist-format-clause in an ex-	43
plicit distribution directive for the ultimate align target of S.	44
	45
With this definition,	46
	47
<ul> <li>Any mapping of a named object is a specialization of itself.</li> </ul>	48

mapping of A is a specialization of the mapping of C.

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 • If A, B, and C are named objects, and if the mapping of A is a specialization of the

mapping of B and the mapping of B is a specialization of the mapping of C, then the

5	That is, the specialization relation, as applied to mappings of named objects, is reflexive
6	and transitive, and it can therefore be applied to produce an equivalence relation on the
7	set of mappings of named objects: two such mappings can be said to be equivalent iff
8	each is a specialization of the other. With this definition, the specialization relation yields a
9	partial ordering on the set of mappings of named objects, modulo equivalence. The INHERIT
10	mapping is the unique maximal element in this partial order.
11	
12 13	4.6 Conditions for Omitting Explicit Interfaces
14	Under certain conditions, an explicit interface for a subprogram is not required. The condi-
15	tions in Fortran under which this is allowable are tightened considerably for HPF programs
16	that use mapping directives.
17	that use mapping directives.
18	Advice to users. These conditions are complex. The important thing to realize is
	that you don't have to read any of this if you have an explicit interface. So if there is
19	any doubt in your mind, just make sure you have an explicit interface. (End of advice
20	to users.)
21	
22	An explicit interface is required <i>except</i> when all of the following conditions hold:
23	r i r o
24	1. Fortran does not require one, and
25	
26 27	2. No dummy argument is distributed transcriptively or with the INHERIT attribute, and
28	3. For each pair of corresponding actual and dummy arguments, either:
29	
30	(a) They are both implicitly mapped, or
31	(b) They are both explicitly mapped and the mapping of the actual argument is a
32	specialization of the mapping of the dummy argument,
33	
34	and
35	
36	4. For each pair of corresponding actual and dummy arguments, either:
37	(a) Both and accuration
38	(a) Both are sequential, or
39	(b) Both are nonsequential.
40	
41	<i>Rationale.</i> This has the following consequences:
42	
43	• A plain Fortran program (i.e., with no HPF directives) will be HPF-conforming
44	without the need to add additional interfaces, at least in a compilation environ-
45	ment in which all variables are sequential by default. This is insured by items 1,
46	2, 3(a), and 4(a).
47	• If remapping is necessary, this fact will be visible to the caller. Thus the imple-
48	mentation may choose to have all remapping performed by the caller.

	(End of rationale.)	1
		2
	Advice to users. This requirement pushes the user strongly in the direction of	3
	always providing explicit interfaces. This is a good thing—explicit interfaces allow	4
	many errors to be caught at compile-time and greatly speed up the process of robust software development.	5 6
	Note that an explicit interface can be provided in three ways:	7
	Tote that an explicit interface can be provided in three ways.	8
	1. A module subprogram has an explicit interface.	9
	2. An internal subprogram has an explicit interface.	10 11
	3. An explicit interface may be provided by an interface block.	12
	In addition, an intrinsic procedure always has an explicit interface by definition.	13 14
	The idiomatic Fortran way of programming makes extensive use of modules; every subprogram, for instance, can be in a module. This provides explicit interfaces automatically, with no extra effort on the part of the programmer. It should very seldom be necessary to write an interface block. ( <i>End of advice to users.</i> )	15 16 17 18 19
4.7	Characteristics of Procedures	20 21
stan	characteristics of dummy data objects and function results as given in the Fortran dard (F95:12.2) are extended to include also the <i>HPF-characteristics</i> of such objects, h are defined recursively as follows:	22 23 24 25
•	A processor arrangement has one HPF-characteristic: its shape.	26 27
•	A template has up to three HPF-characteristics:	28 29
	1. its shape;	30
	2. its distribution, if explicitly stated;	31
	3. the HPF-characteristic (i.e., the shape) of the processor arrangement onto which it is distributed, if explicitly stated.	32 33 34
•	A dummy data object has the following HPF-characteristics:	35 36
	<ol> <li>its alignment, if explicitly stated, as well as all HPF-characteristics of its align target;</li> </ol>	37 38 39
	2. its distribution, if explicitly stated, as well as the HPF-characteristic (i.e., the shape) of the processor arrangement onto which it is distributed, if explicitly stated.	40 41 42
•	A function result has the same HPF-characteristics as a dummy data object. Specifically, it has the following HPF-characteristics:	43 44 45
	1. its alignment, if explicitly stated, as well as all HPF-characteristics of its align target;	46 47 48

2. its distribution, if explicitly stated, as well as the HPF-characteristic (i.e., the shape) of the processor arrangement onto which it is distributed, if explicitly stated.

*Rationale.* In case an explicit interface is given by an interface block, the Fortran standard specifies what information must be specified in that interface block; it does this using the concept of a Fortran *characteristic*. Characteristics of dummy data objects, for instance, include their types. Characteristics must be specified in interface blocks; F95:12.3.2.1 in the Fortran standard states

An interface body specifies all of the procedure's characteristics and these shall be consistent with those specified in the procedure definition ...

Normally, an interface block for a procedure is a textual copy of the appropriate declarations of that procedure. This Section simply says that such a textual copy must include any explicit mapping directives relevant to dummy arguments of the procedure. (*End of rationale.*)

#### 4.8 Argument Passing and Sequence Association

For actual arguments in a procedure call, Fortran allows an array element (scalar) to be associated with a dummy argument that is an array. It furthermore allows the shape of a dummy argument to differ from the shape of the corresponding actual array argument, in effect reshaping the actual argument via the procedure call. Storage sequence properties of Fortran are used to identify the values of the dummy argument. This feature, carried over from FORTRAN 77, has been widely used to pass starting addresses of subarrays, rows, or columns of a larger array, to procedures. For HPF arrays that are potentially mapped across processors, this feature is not fully supported. 

#### 

## 4.8.1 Sequence Association Rules

1. When an array element or the name of an assumed-size array is used as an actual argument, the associated dummy argument must be a scalar or specified to be a sequential array.

An array-element designator of a nonsequential array must not be associated with a dummy array argument.

- 2. When an actual argument is an array or array section and the corresponding dummy argument differs from the actual argument in shape, then the dummy argument must be declared sequential and the actual array argument must be sequential.
- 3. An object of type character (scalar or array) is nonsequential if it conforms to the requirements of Definition 4 of Section 3.8.1.1. If the length of an explicit-length character dummy argument differs from the length of the actual argument, then both the actual and dummy arguments must be sequential.
- 4. Without an explicit interface, a sequential actual may not be associated with a nonsequential dummy and a nonsequential actual may not be associated with a sequential dummy. (This item merely repeats part of Section 4.6).

4.8.2 Discussion of Sequence Association	1
When the shape of the dummy array argument and its associated actual array argument	2
differ, the actual argument must not be an expression. There is no HPF mechanism for	3
declaring that the value of an array-valued expression is sequential. In order to associate	4
such an expression as an actual argument with a dummy argument of different rank, the	5 6
actual argument must first be assigned to a named array variable that is forced to be	7
sequential according to Definition 4 of Section 3.8.1.1.	8
	9
4.8.3 Examples of Sequence Association	10
Given the following subroutine fragment:	11
SUBROUTINE HOME (X)	12
DIMENSION X (20,10)	13 14
	15
By rule 1	16
CALL HOME (ET (2,1))	17
is legal only if <b>X</b> is declared sequential in <b>HOME</b> and <b>ET</b> is sequential in the calling procedure.	18
Likewise, by rules 2 and 4	19 20
CALL HOME (ET)	21
CALL HOME (EI)	22
requires either that $ET$ and $X$ are both sequential arrays or that $ET$ and $X$ have the same	23
shape and (in the absence of an explicit interface) have the same sequence attribute.	24
Rule 3 addresses a special consideration for objects of type character. Change of the	25
length of character objects across a call, as in	26 27
CHARACTER (LEN=44) one_long_word	28
one_long_word = 'Chargoggagoggmanchaugagoggchaubunagungamaugg'	29
CALL webster(one_long_word)	30
-	31
SUBROUTINE webster(short_dictionary)	32
CHARACTER (LEN=4) short_dictionary (11)	33
!Note that short_dictionary(3) is 'agog', for example	34
is an exact welling to well in Franktion. In HDF, both the extend of an exact and down on a survey of	35
is conceptually legal in Fortran. In HPF, both the actual argument and dummy argument must be sequential. (Chargoggagoggmanchaugagoggchaubunagungamaugg is the original	36
Nipmuc name for what is now called Lake Webster in Massachusetts.)	37
inplified nume for what is now cance have webster in Mubbaenuberts.	38 39
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# Section 5

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# INDEPENDENT and Related Directives

16The HPF INDEPENDENT directive allows the programmer to give information to the compiler 17concerning opportunities for parallel execution. The user can assert that no data object 18 is defined by one iteration of a DO loop and used (read or written) by another; similar 19 information can be provided about the combinations of index values in a FORALL statement. 20 Such information is sometimes valuable to enable compiler optimization, but may require 21 knowledge of the application that is available only to the programmer. HPF therefore 22 allows a user to make these assertions, and the compiler may rely on them in its translation 23 process. If the assertion is true, the semantics of the program are not changed; if it is false,  $^{24}$ the program is not HPF-conforming and has no defined meaning.  $^{25}$ 

In contrast to HPF 1.0, the INDEPENDENT assertion of HPF 2.0 allows reductions to be performed in INDEPENDENT loops, provided the reduction operator is a built-in, associative and commutative Fortran operator (such as .AND.) or function (such as MAX). It is often the case that a data parallel computation cannot be expressed in HPF 1.0 as an INDEPENDENT loop because several loop iterations update one or more variables. In such cases parallelism may be possible and desirable because the order of updates is immaterial to the final result. This is most often the case with accumulations, such as the following loop:

<sup>33</sup> DO I = 1, 100000000 <sup>34</sup> X = X + COMPLICATED\_FUNCTION(I) <sup>35</sup> END DO <sup>36</sup>

This loop can run in parallel as long as its iterations make their modifications to the shared variable X in an atomic manner. Alternatively, the loop can be run in parallel by making updates to temporary local accumulator variables, with a (short) final phase to merge the values of these variables with the initial value of X. In either case, the computation is conceptually parallel, but it cannot be asserted to be INDEPENDENT by the strict definition found in HPF 1.0.

It is worth mentioning that Fortran now includes several means to express data parallel computation:

- Array assignments, including the WHERE statement.
- Elemental invocation of intrinsic and user-defined functions.

• The FORA tions.	LL statement and const	ruct	, including element-wise invocation of PURE func-	1 2	
• Transformational intrinsics such as SUM and TRANSPOSE.					
			from HPF version 1.0. As these are all now part	5 6	
of Fortran, they	v are not discussed sepa	irate	ely in this document.	7	
				8	
5.1 The IN	DEPENDENT Dir	ect	ive	9	
The INDEPENDE	INT directive can prece	de a	n indexed DO loop or FORALL statement. It as-	10	
	_		n the following $DO$ loop or the operations in the	11 12	
following FORAL	L may be executed ind	eper	ndently—that is, in any order, or interleaved, or	12	
-	without changing the se			14	
	-		ne DO loop or FORALL for which it asserts behavior,	15	
and is said to a	<i>pply</i> to that loop or FU	RALI	L. The syntax of the INDEPENDENT directive is	16	
H501 independ	lent-directive	is	INDEPENDENT [ , new-clause ]	17	
-			[ , reduction-clause ]	18	
H502 new-clau	se	is	NEW ( variable-name-list )	19 20	
H503 reduction	n-clause	is	<b>REDUCTION (</b> reduction-variable-list )	21	
				22	
H504 reduction	n-variable	is or	array-variable-name scalar-variable-name	23	
		or	structure-component	24	
				25 26	
Constraint: Th	ne first non-comment li	ne fo	ollowing an <i>independent-directive</i> must be a <i>do-</i>	20	
str	<i>nt</i> , <i>forall-stmt</i> , or a <i>for</i>	all-c	onstruct.	28	
Constraint: If	the first non-comment	line	following an <i>independent-directive</i> is a <i>do-stmt</i> ,	29	
			ntain a loop-control option containing a do-vari-	30	
ab	le.			31 32	
Constraint: If	aithar tha NEW clause or	• tho	REDUCTION clause is present, then the first non-	33	
			rective must be a <i>do-stmt</i> .	34	
	0			35	
			or the REDUCTION clause and any component or	36	
ele	ment thereof must not	:		37	
	• Be a dummy argume	ont•		38	
			T - 44-11 - 44-1	39 40	
	• Have the SAVE or TA			41	
	• Occur in a COMMON b	lock	;	42	
	0		th another object as a result of appearing in an	43	
	EQUIVALENCE statem	ient;		44	
	• Be use associated;			45	
	• Be host associated; o	or		46 47	
	• Be accessed in anoth	ner s	coping unit via host association.	47	

# 5.1. THE INDEPENDENT DIRECTIVE

1 2 3 4	Constraint: A variable that occurs as a <i>reduction-variable</i> may not appear in a in the same <i>independent-directive</i> , nor may it appear in either a <i>ne</i> a <i>reduction-clause</i> in the range (i.e., the lexical body) of the following <i>forall-stmt</i> , or <i>forall-construct</i> to which the <i>independent-directive</i> a	<i>w-clause</i> or ng <i>do-stmt</i> ,
5 6 7	Constraint: A <i>structure-component</i> in a <i>reduction-variable</i> may not contain a <i>section-list</i> .	. subscript-
8 9 10	Constraint: A variable that occurs as a $reduction$ -var must be of intrinsic type. be of type CHARACTER.	It may not
11 12 13 14 15	<i>Rationale.</i> The second constraint means that an INDEPENDENT directive applied to a WHILE loop or a simple DO loop (i.e., a "do forever"). An IN in such cases could only correctly describe a loop with zero or one trips; the confusion was felt to outweigh the possible benefits. ( <i>End of rationale.</i> )	DEPENDENT
16 17 18 19	When applied to a DO loop, an INDEPENDENT directive is an assertion by the p that no iteration can interfere with any other iteration, either directly or indir following operations define such interference:	0
20 21 22	• Any two operations that assign to the same atomic object interfere with (A data object is called <i>atomic</i> if it contains no subobjects.)	each other.
23 24 25	<ul> <li>Exception: If a variable appears in a NEW clause, then operations assig to it in separate iterations of the DO loop <i>do not</i> interfere. The reason explained in Section 5.1.2.</li> </ul>	-
26 27 28 29 30	- Exception: If a variable appears in a REDUCTION clause, then assign it by reduction statements in the range of the DO loop <i>do not</i> int assignments to it by other reduction statements in the same loop. for this is explained in Section 5.1.3.	erfere with
31	Operations that assign to objects include:	
32 33	- Assignment statements assign to their left-hand side and all its subo	bjects.
34 35	<ul> <li>ASSIGN statements assign to their integer variables.</li> <li>ALLOCATE and DEALLOCATE statements with the STAT= specifier assign</li> </ul>	to the STAT
36 37	variable.	
38 39	- DO statements assign to their indices. - I/O statements with the IOSTAT= specifier assign to the IOSTAT vari	able. Thev
40	may also assign to other objects, as described below.	asie. They
41 42	<ul> <li>Asynchronous READ and WRITE statements (as described in Section 10 their ID= variable.</li> </ul>	)) assign to
43 44 45 46	<ul> <li>READ statements assign to all variables in their input item list and an accessed at runtime through their NAMELIST. READ statements with specifier assign to the SIZE variable.</li> </ul>	e e
40 47 48	- INQUIRE statements assign to all variables in their specifier list, except and FILE specifiers.	ot the UNIT

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- Compound statements (e.g., IF statements) cause assignments to objects if their component statements do.
- Subprogram invocations cause assignments to objects if operations in the subprogram execution do.
- An operation that assigns to an atomic object interferes with any operation that uses the value of that object.
  - Exception: If a variable appears in a NEW clause, then operations assigning values to it in one iteration of the DO loop *do not* interfere with uses of the variable in other iterations. The reason for this is explained in Section 5.1.2.
  - Exception: If a variable appears in a **REDUCTION** clause, then assignments to it by reduction statements in the range of the **DO** loop *do not* interfere with the allowed uses of it by reduction statements in the same loop. The reason for this is explained in Section 5.1.3.

Any expression that computes the value of a variable uses that object. This includes uses on the right-hand side of assignment statements, uses in subscripts on the lefthand side of assignment statements, conditional expressions, specification lists for I/O statements, output lists for WRITE statements, allocation shape specifications in ALLOCATE statements, and similar situations.

Rationale. These are the classic Bernstein conditions to enable parallel ex-23 ecution. Note that two assignments of the same value to a variable interfere  $^{24}$ with each other and thus an INDEPENDENT loop with such assignments is not  $^{25}$ HPF-conforming. This is not allowed because such overlapping assignments are 26 difficult to support on some hardware, and because the given definition was felt 27to be conceptually clearer. Similarly, it is not HPF-conforming to assert that 28 assignment of multiple values to the same location is INDEPENDENT, even if the 29 program logically can accept any of the possible values. In this case, both the 30 "conceptually clearer" argument and the desire to avoid indeterminate behavior 31 favored the given solution. (*End of rationale.*) 32

• An ALLOCATE statement, DEALLOCATE statement, NULLIFY statement or pointer assignment statement interferes with any other access, pointer assignment, allocation, deallocation, or nullification of the same pointer. In addition, an ALLOCATE or DEALLOCATE statement interferes with any other use of or assignment to the object that is allocated by ALLOCATE or deallocated by DEALLOCATE.

*Rationale.* These constraints extend Bernstein's conditions to pointers. Because a Fortran pointer is an alias to an object or subobject rather than a first-class data type, a bit more care is needed than for other variables. (*End of rationale.*)

- Any transfer of control to a branch target statement outside the body of the loop interferes with all other operations in the loop.
- Any execution of an EXIT, STOP, or PAUSE statement interferes with all other operations in the loop.

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Rationale. Branching (by GOTO or ERR= branches in I/O statements) implies that some iterations of the loop are not executed, which is drastic interference with those computations. The same is true for EXIT and the other statements. Note that these conditions do not restrict procedure calls in INDEPENDENT loops, except to disallow taking alternate returns to statements outside the loop, executing a STOP, or executing a PAUSE. (End of rationale.)

• Any two file I/O operations except INQUIRE associated with the same file or unit interfere with each other. Two INQUIRE operations do not interfere with each other; however, an INQUIRE operation interferes with any other I/O operation associated with the same file.

Rationale. Because Fortran carefully defines the file position after a data transfer or file positioning statement, these operations affect the global state of a program. (Note that file position is defined even for direct access files.) Multiple non-advancing data transfer statements affect the file position in ways similar to multiple assignments of the same value to a variable, and is disallowed for the same reason. Multiple OPEN and CLOSE operations affect the status of files and units, which is another global side effect. INQUIRE does not affect the file status, and therefore does not affect other inquiries. However, other file operations may affect the properties reported by INQUIRE. (End of rationale.)

• Any data realignment or redistribution performed by subprogram invocation (see Section 4) interferes with any access to or any other remapping of the same data.

*Rationale.* Remapping may change the processor storing a particular array element, which interferes with any assignment or use of that element. This applies even though the remappings are "undone" when the call returns. During the execution of the call, the homes of the array elements have changed, thus interfering with accesses in the caller, accesses in other invocations of the same procedure, and remappings of the array due to another procedure call. (*End of rationale.*)

Advice to users. Data remapping performed by the **REALIGN** and **REDISTRIBUTE** aproved extensions also causes interference under this rule. See Chapter 8.5 for details. (*End of advice to users.*)

The interpretation of INDEPENDENT for FORALL is similar to that for DO: it asserts that no combination of the FORALL indices assigns to an atomic storage unit that is read by another combination. A DO and a FORALL with the same body are equivalent if they both have the INDEPENDENT directive. This is illustrated in Section 5.1.1.

<sup>40</sup> If a procedure is called from within an INDEPENDENT loop or FORALL, then any local <sup>41</sup> variables in that procedure are considered distinct on each call unless they have the SAVE <sup>42</sup> attribute. This is consistent with the Fortran standard. Therefore, uses of local variables <sup>43</sup> without the SAVE attribute in calls from different iterations do not cause interference as <sup>44</sup> defined above.

46 Advice to implementors. A conforming Fortran implementation can often avoid 47 creating distinct storage for locals on every call. The same is true for an HPF imple-48 mentation; however, such an implementation must still interpret INDEPENDENT in the

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same way. If locals are not allocated unique storage locations on every call, then the INDEPENDENT loop must be serialized to respect these semantics (or other techniques must be used to avoid conflicting accesses). (End of advice to implementors.)

Note that all these rules describe interfering behavior; they do not disallow specific syntax. Statements that appear to violate one or more of these restrictions are allowed in an INDEPENDENT loop, if they are not executed due to control flow. These restrictions allow an INDEPENDENT loop to be executed safely in parallel if computational resources are available. The directive is purely advisory and a compiler is free to ignore it if it cannot make use of the information.

Advice to implementors. Although the restrictions allow safe parallel implementation of INDEPENDENT loops, they do not imply that this will be profitable (or even possible) on all architectures or all programs. For example,

- An INDEPENDENT loop may call a routine with explicitly mapped local variables. The implementation must then either implement the mapping (which may require serializing the calls, under some implementation strategies) or override the explicit directives (which may surprise the user).
- An INDEPENDENT loop may have very different behavior on different iterations. For example,

```
!HPF$ INDEPENDENT
DO i = 1, 3
IF (i.EQ.1) CALL F(A)
IF (i.EQ.2) CALL G(B)
IF (i.EQ.3) CALL H(C)
END DO
```

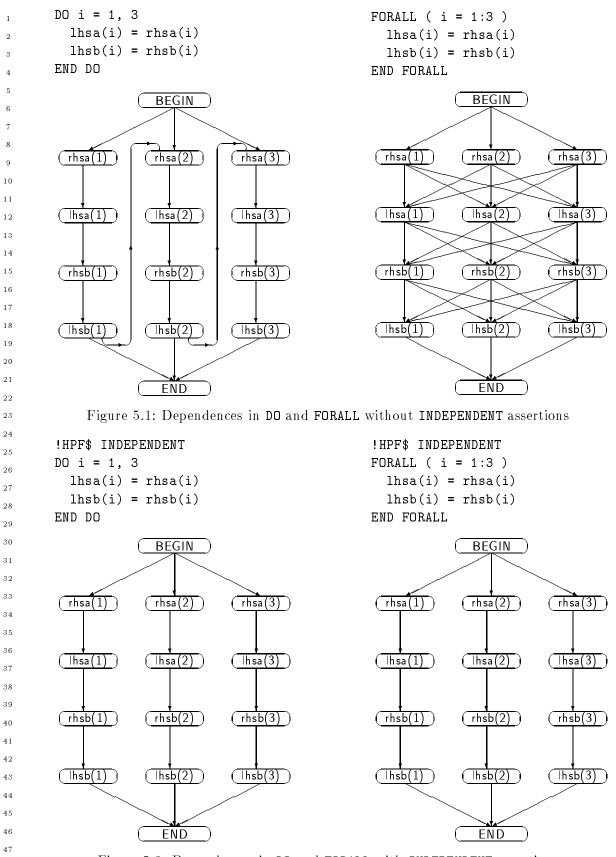
This poses obvious problems for implementations on SIMD machines.

• An INDEPENDENT loop may call a subroutine that accesses global mapped data. On distributed-memory machines, generating the communication to reference the data may be challenging, since there is in general no guarantee that the owners of the data will also call the subroutine.

In all cases, it is the implementation's responsibility to produce correct behavior, which may in turn limit optimization. It is recommended that implementations provide some feedback if an INDEPENDENT assertion may be ignored. (*End of advice to implementors.*)

# 5.1.1 Visualization of INDEPENDENT Directives

Graphically, the INDEPENDENT directive can be visualized as eliminating edges from a prece-41dence graph representing the program. Figure 5.1 shows some of the dependences that 42 may normally be present in a DO and a FORALL. (Most of the transitive dependences are not 43 shown.) An arrow from a left-hand-side node (for example, "lhsa(1)") to a right-hand-side 44 node ("rhsb(1)") means that the right-hand side computation might use values assigned 45in the left-hand-side node; thus the right-hand side must be computed after the left-hand 46 side completes its store. Similarly, an arrow from a right-hand-side node to a left-hand-side 47node means that the left-hand side may overwrite a value needed by the right-hand side 48





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computation, again forcing an ordering. Edges from the BEGIN and to the END nodes represent control dependences. The INDEPENDENT directive asserts that the only dependences that a compiler need enforce are those in Figure 5.2. That is, the programmer who uses INDEPENDENT is certifying that if the compiler enforces only these edges, then the resulting program will be equivalent to the one in which all the edges are present. Note that the set of asserted dependences is identical for INDEPENDENT DO and FORALL statements.

The compiler is justified in producing a warning if it can prove that one of these assertions is incorrect. It is not required to do so, however. A program containing any false assertion of this type is not HPF-conforming, thus is not defined by HPF, and the compiler may take any action it deems appropriate.

#### 5.1.1.1 Examples of INDEPENDENT

```
!HPF$ INDEPENDENT
DO i = 2, 99
A(I) = B(I-1) + B(I) + B(I+1)
END DO
```

This is one of the simplest examples of an INDEPENDENT loop. (For simplicity, all examples in this section assume there is no storage or sequence association between any variables used in the code.) Every iteration assigns to a different location in the **A** array, thus satisfying the first condition above. Since no elements of **A** are used on the right-hand side, no location that is assigned in the loop is also read, thus satisfying the second condition. Note, however, that many elements of **B** are used repeatedly; this is allowed by the definition of **INDEPENDENT**. This loop is **INDEPENDENT** regardless of the values of the variables involved.

```
!HPF$ INDEPENDENT
FORALL ( I=2:N ) A(I) = B(I-1) + B(I) + B(I+1)
```

This example is equivalent in all respects to the first example.

```
!HPF$ INDEPENDENT
DO I=1, 100
   A(P(I)) = B(I)
END DO
```

This INDEPENDENT directive asserts that the array P does not have any repeated entries (else they would cause interference when A was assigned). The DO loop is therefore equivalent to the Fortran statement

A(P(1:100)) = B(1:100)

### 5.1.2 NEW Variables

The NEW clause asserts that the named variables act as private variables to each iteration 42 of the INDEPENDENT loop. That is, there would be no interfering assignments and uses in 43 the loop, and thus no change in ghe behavior of the program, if new objects were created 44 for the NEW variables for each iteration of the DO loop and those objects were destroyed at 45 the end of each iteration. Thus, no values flow into NEW variables from execution before the 46 loop, no values flow from NEW variables to execution after the loop, and (most importantly) 47 no values flow from one iteration to another through NEW variables. 48 Advice to users. A pointer or allocatable variable may appear in a NEW clause. The interpretation of the paragraph above, in these cases, is that one should not rely on the value, the association status, or the allocation status of such a variable on entry to the loop; rather, such variables should be allocated or pointer assigned in the loop body before they are used. It would also be advisable to deallocate or nullify such a variable in the loop body after its last use as well. (End of advice to users.)

Rationale. NEW variables provide the means to declare temporaries in INDEPENDENT loops. Without this feature, many conceptually independent loops would need substantial rewriting (including expansion of scalars into arrays) to meet the rather strict requirements described above. Note that a temporary must be declared NEW only at the innermost lexical level at which it is assigned, since all enclosing INDEPENDENT assertions must take that NEW into account. Note also that index variables for nested DO loops must be declared NEW; the alternative was to limit the scope of an index variable to the loop itself, which changes Fortran semantics. FORALL indices, however, are restricted by the semantics of the FORALL; they require no NEW declarations. (End of rationale.)

<sup>19</sup> 5.1.2.1 Examples of NEW

This example would be correct either with or without the NEW clause; in either case, the compiler could confidently parallelize the assignments to array A. Additionally however, the NEW clause asserts that the loop index I is not used after the completion of the loop. Some compilers may be able to use this information to avoid updating replicated copies of I on other processors, or to enable other optimizations.

```
!HPF$ INDEPENDENT, NEW (I2)
31
       DO I1 = 1, N1
32
           !HPF$ INDEPENDENT, NEW (I3)
33
          DO I2 = 1, N2
34
              DO I3 = 2,N3
                              ! The inner loop is NOT independent!
35
                  A(I1,I2,I3) = A(I1,I2,I3) - A(I1,I2,I3-1)*B(I1,I2,I3)
36
              END DO
37
          END DO
38
       END DO
39
```

The inner loop is not independent because each element of A is computed from the preceding
 one. However, the two outer loops are independent because they access different elements
 of A. The NEW clauses are required, since the inner loop indices are assigned and used in
 different iterations of the outermost loops.

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# 5.1.3 **REDUCTION** Variables and Statements

The REDUCTION clause asserts that the named variables are updated in the INDEPENDENT loop by a series of operations that are associative and commutative. Furthermore, the

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intermediate values of the **REDUCTION** variables are not used within the loop (except, of course, in the updates themselves). Thus, the value of a **REDUCTION** variable after the loop may be computed as the result of a reduction tree.

Rationale. REDUCTION variables provide the means to accumulate values generated in an INDEPENDENT loop. Without this feature, the programmer must store update information in a temporary array whose size is equal to the number of loop iterations, and then use an intrinsic reduction function or XXX\_SCATTER library function after the loop. The problem with this approach is that the temporary array may be excessively large. (End of rationale.)

The semantics of reductions are discussed in detail in Section 5.1.4. This section defines correct syntax.

Any variable whose name occurs as a *reduction-variable* is said to be *protected* while the immediately following DO loop is active (i.e. being executed). It may not be referenced while the loop in which it is protected is active, with one exception. It may occur in special locations in assignment statements of a special form, and these statements must be in the range (i.e. the lexical body) of the loop. In particular, it may not occur in any HPF directive, including the variable list in a NEW clause. This includes any NEW clause in the same INDEPENDENT directive.

A reduction statement is an assignment statement of the following special form that occurs in the range of an independent DO loop for which the name of its reduction variable occurs in a reduction clause. This description is not part of the grammar of HPF; rather, it serves to define the restricted assignment statements in which reduction variables are allowed.

variable - variable mult on mult open and	26
variable = add-operand * variable	27
variable = variable add-op add-operand	28
variable = level-2-expr + variable	29
variable = variable and-op and-operand	30
variable = and-operand and-op variable	31
variable = variable or-op or-operand	32
variable = or-operand or-op variable	33
variable = variable equiv-op equiv-operand	34
variable = equiv-operand equiv-op variable	35
$\cdot$ variable = reduction-function ( variable , expr )	36
$\cdot$ variable = reduction-function ( expr , variable )	37
	38
MAX	39
MIN	40
IAND	
	41
	42
	43
	r variable = variable mail-op mail-operanta r variable = add-operand * variable r variable = variable add-op add-operand r variable = level-2-expr + variable r variable = variable and-op and-operand r variable = and-operand and-op variable r variable = variable or-op or-operand r variable = or-operand or-op variable r variable = variable equiv-op equiv-operand r variable = variable equiv-op variable r variable = reduction-function ( variable , expr ) r variable = reduction-function ( expr , variable ) MAX r MIN r IAND r IOR r IEOR

Constraint: The two occurances of *variable* in a *reduction-stmt* must be textually identical.

The first two assertions of Section 5.1 account for the fact that the occurrences of 46 reduction variables in their allowed positions in reduction statements do not cause interfer-47 ence between iterations of an INDEPENDENT loop. Any other assignment to or reference to 48

### 5.1. THE INDEPENDENT DIRECTIVE

a reduction variable *does* interfere with the reduction statement; this includes occurrences 1 in subprograms and in the *expr* part of a reduction statement.  $^{2}$ A variable that is updated by reduction statements in an independent loop must be 3 protected by explicit appearance in a reduction clause. This clause must appear in the 4 **INDEPENDENT** directive for the outermost independent loop that 5 6 • Contains the reduction statement; 7 8 • Does not have a NEW clause naming the reduction variable; and 9 10 • Lies within the innermost independent loop, if any, that contains the reduction state-11 ment and does have a NEW clause naming the reduction variable. 12If the same variable is updated by two or more reduction statements, then the operators in 13 14those statements must be in the same class (e.g. both must be an *add-op* if one is). 1516Advice to users. When a reduction statement is executed, some nest of DO loops 17will be active. If there are several nested INDEPENDENT DO loops surrounding the 18 reduction statements in which the variable is updated, which one is the right one to 19 get the reduction clause? The answer is the outermost one, subject to the constraint 20 that a reduction variable may not appear in a NEW clause for that loop or a contained 21 loop. Consider 22 23 !HPF\$ INDEPENDENT, NEW(J), REDUCTION(X) 24 DO I = 1, 10 25!HPF\$ INDEPENDENT 26 DO J = 1, 2027 X = X + J28 END DO 29 END DO 30 31 It would be incorrect to move the reduction clause to the inner **INDEPENDENT** directive. 32 Since X is updated by reduction operations (twenty times) for *each* iteration of the 33 outer loop, it does not have a well-defined value until the completion of the outer 34 loop. (End of advice to users.) 35 36 The *reduction-variable* reference may be an array element or array section. The two 37 references that occur in a reduction statement must be lexically identical. The Fortran rules 38 of operator precedence and the use of parentheses in the expression must ensure that the 39 reduction operator is the top-level operator (i.e. it is evaluated last) on the right-hand side.

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X = X \* A + 1

Therefore,

is not a correctly formed reduction statement.

<sup>45</sup> Note that the syntax of the INDEPENDENT directive does not allow an array element or
<sup>46</sup> array section to be designated as a reduction variable in the reduction clause. Even though
<sup>47</sup> such a subobject may occur in a reduction statement, it is the entire array or character
<sup>48</sup> variable that is treated as a reduction variable.

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The allowed reduction operators and functions are all associative (in their mathematical definitions, even though the usual implementations of the arithmetic operators by Fortran language processors and the underlying hardware are not).

In most cases, only one operator will be used in the reduction statements (if there are more than one) that update a given reduction variable. It is sensible, however, to use + and - together on the same reduction variable: mathematically, subtraction is just addition of the additive inverse. For example:

```
!HPF$ INDEPENDENT, REDUCTION(X)
DO I = 1, 100
    X(IDX(I,1)) = X(IDX(I,1)) + Y(I)
    X(IDX(I,2)) = X(IDX(I,2)) - Y(I)
    END DO
```

The same is true for multiplication (\*) and division (/). No other mixing of operators is allowed.

Advice to users. While it is true that

```
X = I + X
```

is permitted as a reduction statement, for most purposes

X = X + I

is stylistically cleaner. (End of advice to users.)

## 5.1.4 Semantics and Implementation of Reduction

HPF specifies an allowed parallel implementation of an INDEPENDENT DO loop with reduction statements, thereby specifying the semantics of such a loop.

Just as the result of the Fortran intrinsic function SUM is defined to be a implementation-dependent approximation to the sum of the elements of its argument array, the value of a reduction variable on exit from its INDEPENDENT DO loop is likewise not completely specified by HPF. One possible value is that which would have been computed by sequential execution of the loop, but other implementation-dependent approximations to this value may be produced. Any such implementation-dependent value is, however, an approximation to the value produced by sequential execution of the loop. If rounding error, underflow, and overflow do not occur, it will be identical to that value. 

Advice to users. If overflow, underflow, or rounding occur, this is one of the few places where an HPF directive in a conforming program may cause that program to produce different output. However, the same problems occur in other systems that attempt to parallelize these operations, for the same reasons. (*End of advice to users.*)

Since no reference to a protected reduction variable can occur except in a reduction statement, it is not necessary to define the values that these variables may have while protected.

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Advice to users. The following "advice to implementors" is useful for understanding the behavior of an INDEPENDENT loop with reduction statements. (End of advice to users.)

Advice to implementors. In the discussion in this section, the term "processor" means a single physical processor or a group of physical processors that together sequentially execute some or all of the iterations of an independent loop.

We describe a simple implementation mechanism that applies to commutative reduction operations. On entry to an independent loop, every executing processor allocates a private accumulator variable associated with each variable in the reduction clause on the INDEPENDENT directive, and initializes it to the identity element for the corresponding intrinsic reduction operator. The private accumulator variable has the same shape, type, and kind type parameter as the reduction variable.

<sup>14</sup> The identity elements for the intrinsic operators are defined in Table 5.1.

16	Operator	Identity Element			
17					
18	+	0			
19	-	0			
20	*	1			
21	/	1			
22	. AND .	.TRUE.			
23	. OR .	.FALSE.			
24	.EQV.	.TRUE.			
25	.NEQV.	.FALSE.			
26					
27	T-11- F 1. LI	antita alamanta fan inteineir ar lastian ananatara			
28	Table 5.1: Id	entity elements for intrinsic reduction operators.			
29					
30	Function	Identity element			
31		v			
32	IAND(I,J)	NOT(O) (all one-bits)			
33	IOR(I,J)	0			
34	IEOR(I,J)	0			
35	MIN(X,Y)	the positive number of largest absolute value			
36		that has the same type and kind type param-			
37		eter as the reduction variable			
38	MAX(X,Y)	the negative number of largest absolute value			
39		that has the same type and kind type param-			
40		eter as the reduction variable			
41					
42					
43	Table 5.2: Id	entity elements for intrinsic reduction functions.			
44	The intrincia function	a that may be used as reduction functions are listed together.			
45	with their identity ele	s that may be used as reduction functions are listed, together ments, in Table 5.2			
46					
47	Each processor performs a subset of the loop iterations; when it encounters a reduction				
48	statement, it updates	its own accumulator variable. A processor is free to perform its			

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loop iterations in any order; furthermore, it may start an iteration, suspend work on it, do some or all of the work of other iterations, and resume work on the suspended iteration. However, any update of a private accumulator variable occurs through the execution of a reduction statement, and reduction statements *are* executed atomically.

The final value of the reduction variable is computed by combining the private accumulator variables with the value of the reduction variable on entry to the loop, using the reduction operator. The ordering of this reduction is language-processor dependent, just as it is for the intrinsic reduction functions (SUM, etc.).

As an example, consider:

REAL Z

Z = 5.!HPF\$ INDEPENDENT, REDUCTION(Z) DO I = 1, 10 Z = Z + IEND DO

The final value of Z will be 5 + (1+2+3+4+5+6+7+8+9+10) = 60; the order in which the additions occur is not specified by HPF.

For a second example, here is a SUM\_SCATTER done as an independent loop:

```
!HPF$ INDEPENDENT, REDUCTION(X)
DO I = 1, N
        X(INDEX(I)) = X(INDEX(I)) - F(I)
END DO
```

The implementation will most likely make a private copy on every processor of an accumulator array XLOCAL of the same type and shape as X, and initialize it to zero. Each iteration will subtract the value of F(I) from its own XLOCAL(INDEX(I)). To create the final result, the implementation must combine all the private accumulator arrays with the initial value of X. The combining operator is the same as the reduction operator, namely addition, so that the result is the sum of the initial value of X and the accumulator arrays. The implementation has the option of using a sparse data structure to store only the updated elements of the local accumulator.

In an MPI based implementation, the MPI\_REDUCE function could be used for this task. (*End of advice to implementors.*)

## 5.2 Further Examples of INDEPENDENT Directives

If IOUNIT(I) evaluates to a different value for every value of I from 1 to 10, then the loop writes to a different I/O unit (and thus a different file) on every iteration. The loop is then properly described as independent. On the other hand, if IOUNIT(I)=5 for all I, then the assertion is in error and the directive is not HPF-conforming.

```
!HPF$ INDEPENDENT, NEW (J)
6
              DO I = 2, 100, 2
7
               !HPF$ INDEPENDENT, NEW(VL, VR, UL, UR)
8
                 DO J = 2, 100, 2
9
                   VL = P(I,J) - P(I-1,J)
10
                   VR = P(I+1,J) - P(I,J)
11
                   UL = P(I,J) - P(I,J-1)
12
                   UR = P(I, J+1) - P(I, J)
13
                   P(I,J) = F(I,J) + P(I,J) + 0.25 * (VR - VL + UR - UL)
14
                 END DO
15
              END DO
16
17
     Without the NEW clause on the J loop, neither loop would be independent, because an
18
     interleaved execution of loop iterations might cause other values of VL, VR, UL, and UR to
19
     be used in the assignment of P(I,J) than those computed in the same iteration of the
20
     loop. The NEW clause, however, specifies that this is not true if distinct storage units are
21
     used in each iteration of the loop. Using this implementation makes iterations of the loops
22
     independent of each other. Note that there is no interference due to accesses of the array
23
     P because of the stride of the DO loop (i.e. I and J are always even, therefore I-1, etc., are
24
     always odd.)
25
          When loops are nested, a reduction variable may need to be protected in an independent
26
     outer loop even though the reduction operations in which it occurs are nested inside an inner
27
     loop. Moreover, the inner loop and any intervening loops may or may not be independent.
28
29
        Ţ
            Nested Loop Example 1. Inner loop is sequential
30
31
              X = 10
32
       OUTER: DO WHILE (X < 1000)
                                         ! this loop is sequential
33
                  !HPF$ INDEPENDENT, NEW(J), REDUCTION(X)
34
                  DO I = 1, N
       MIDDLE:
35
        INNER:
                    DO J = 1, M
36
                       X = X + J
37
                         Note that it would be incorrect to refer to X
38
                       ! here, except in another reduction statement
39
                    END DO INNER
40
                  !
                     Note that it would be incorrect to refer to X
41
                  ! here, except in another reduction statement
42
                  END DO MIDDLE
43
                  PRINT *, X
44
                END DO OUTER
45
     Since the variable X occurs in a reduction clause for loop MIDDLE, it is a protected reduction
46
     variable throughout that loop, including inside the inner loop. If INNER had an INDEPENDENT
47
```

directive, it would be incorrect to include X in a REDUCTION or a NEW clause of that directive.

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The outermost loop is not independent, and so **X** need not and cannot be protected in that part of its range outside the middle loop.

A variable that occurs in a NEW clause must not be a reduction variable in the same or a containing loop, although it may be used as a reduction variable in a contained loop:

```
5
      Nested Loop Example 2. Outer loop NEW clause.
  i
                                                                                           6
  !HPF$ INDEPENDENT, NEW(I)
                                                                                           8
  OUTER: DO K = 1, 100
                                                                                           9
             !HPF$ INDEPENDENT, NEW (J,X)
                                                                                          10
  MIDDLE:
             DO I = 1, N
                                                                                          11
                X = 10
                                                                                          12
                !HPF$ INDEPENDENT, REDUCTION(X)
                                                                                          13
  INNER:
                 DO J = 1, M
                                                                                          14
                   X = X + J * * 2
                                                                                          15
                   ! Note that it would be incorrect to refer to X
                                                                                          16
                   ! here, except in another reduction statement
                                                                                          17
                 END DO INNER
                                                                                          18
                 Y(I) = X
                                                                                          19
             END DO MIDDLE
                                                                                          20
           END DO OUTER
                                                                                          21
Here, X is a protected reduction variable only in the inner loop.
                                                                                          22
                                                                                          23
  INTEGER, DIMENSION(M) :: VECTOR
                                                                                          ^{24}
                                                                                          ^{25}
  !HPF$ INDEPENDENT, REDUCTION(X, Y)
                                                                                          26
         DO I = 1, N-4
                                                                                          27
           X(I:I+4) = X(I:I+4) + A(I)
                                             ! As many as 5 updates
                                                                                          28
           Y(VECTOR) = Y(VECTOR) + B(I,1:N)
                                                                                          29
         END DO
                                                                                          30
                                                                                          31
Note that the compiler, if it distributes iterations of this loop in a block-wise manner, will
                                                                                          32
not need to make a private copy of the entire array X on each processor.
                                                                                          33
    If a statement that has the form of a reduction statement occurs while an independent
                                                                                          34
loop is active, but the updated variable is not a protected reduction variable, then the
                                                                                          35
programmer is guaranteeing that no two iterations of the independent loop will update the
                                                                                          36
same location. For example:
                                                                                          37
                                                                                          38
  !HPF$ INDEPENDENT
                                                                                          39
         DO I = 1, N
                                                                                          40
```

```
! X is NOT a reduction variable, but
! I know there are no repeated values in INDX(1:N)
! Updates will be written directly to X(INDX(I))
X(INDX(I)) = X(INDX(I)) + F(I)
! I also guarantee that the condition in the IF statement
! is true for at most one value of I.
IF (A(I) > B(I)) Y = Y + 1
END DO
```

Section 6

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# **Extrinsic Program Units**

14The HPF global model of computation extends (and restricts) Fortran to provide program-15mers with the Fortran model of computation implementable efficiently on a wide class of 16hardware architectures with, in general, multiple processors, multiple memories with non-17uniform access characteristics, and multiple interconnections. This model of computation 18 presents a single logical thread of control, including Fortran's data parallel features such 19as array syntax and the FORALL statement, and data visibility defined by the scoping rules 20 of Fortran. In particular, this model does not require the use of low-level features such as 21 threads libraries and explicit message passing to exploit such architectures. Programmers 22 expect their HPF compilers to generate efficient code by using HPF's features to assist in 23 mapping data and computation to the given hardware architecture.

 $^{24}$ This chapter defines the *extrinsic* mechanism by which HPF program units may use  $^{25}$ non-HPF program units that don't use the HPF global model. It describes how to write 26 an explicit interface for a non-HPF procedure and defines the caller's assumptions about 27 handling distributed and replicated data at the interface. This allows the programmer to 28 use non-HPF language facilities, for example, to descend to a lower level of abstraction to 29 handle problems that are not efficiently addressed by HPF, to hand-tune critical kernels, 30 or to call optimized libraries. Such an interface can also be used to interface HPF to other 31 languages, such as C. 32

6.1 Overview

An HPF program may need to call a procedure implemented in a different programming model or in a different programming language. A procedure's *programming model* might provide:

- a single logical thread-of-control where *one* copy of the procedure is conceptually executing and there is a single locus of control within the program text; this model is called *global* when the underlying target hardware has (potentially) multiple processors or memories and is called *serial* when the underlying target hardware is treated as a uniprocessor (or a single node in a multiprocessor),
- multiple threads-of-control, one per processor, each thread executing the same procedure; this model is called *local* or, more generally, SPMD (Single Program, Multiple Data), or

• some other model, not discussed here, such as multiple threads-of-control, perhaps with dynamic assignment of loop iterations to processors or explicit dynamic process forking, where there is, at least initially upon invocation, one copy of the procedure that is conceptually executing but that may spawn multiple loci of control, possibly 4 changing in number over time, within the program text. 5

A programming language provides a specific syntax (language features), semantics (meanings), and pragmatics (purposes). Examples of programming languages include Fortran (an ANSI and ISO standard—the most recent revision is expected to be approved by 1997), HPF (a specification of extensions and restrictions to Fortran), Fortran 77 (a 10 previous ANSI and ISO standard), C, C++, Java, Visual Basic, and COBOL. 11

A program unit's language and model, when taken together, constitute its *extrinsic* kind. This extrinsic kind may be specified explicitly by an extrinsic-prefix or implicitly by the selection of a compiler and its invocation with a particular set of compiler options. Thus, one might view the compiler as providing a *host scoping unit* as defined by Fortran. For example, a program unit compiled by an HPF compiler will be of extrinsic kind HPF. Alternatively, its extrinsic kind may be specified explicitly by an *extrinsic-prefix* such as EXTRINSIC(HPF) or EXTRINSIC(LANGUAGE='HPF', MODEL='GLOBAL') .

#### **Declaration of Extrinsic Program Units** 6.2

#### 6.2.1**Function and Subroutine Statements**

An extrinsic-prefix may appear in a function-stmt or subroutine-stmt (as defined in the Fortran standard) in the same place that the keywords RECURSIVE, PURE, and ELEMENTAL may appear. This is specified by an extension of rule R1219 for *prefix-spec* in the Fortran standard. Rules R1217 for function-stmt, R1218 for prefix, and R1222 for subroutine-stmt are not changed, but are restated here for reference.

H601 function-stmt	is	[ prefix ] FUNCTION function-name	29 30
		( [ dummy-arg-name-list ] ) [ RESULT ( result-name ) ]	31
			32
m H602~subroutine-stmt	is	[ prefix ] SUBROUTINE subroutine-name	33
		[ ( [ dummy-arg-list ] ) ]	34
H602 modin	:-	nnefin ence [ nnefin ence ]	35
H603 prefix	IS	prefix-spec [ prefix-spec ]	36
H604 prefix-spec	is	type- $spec$	37
	or	RECURSIVE	38
	or	PURE	39
	or	ELEMENTAL	40
	or	extrinsic-prefix	41
			42

Constraint: Within any HPF external-subprogram, every internal-subprogram must be of the same extrinsic kind as its host and any *internal-subprogram* whose extrinsic kind is not given explicitly is assumed to be of that extrinsic kind.

The definition of *characteristics of a procedure* as given in F95:12.2 is extended to include the procedure's extrinsic kind.

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# 6.2.2 Program, Module, and Block Data Statements

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An *extrinsic-prefix* may also appear at the beginning of a *program-stmt*, *module-stmt*, or *block-data-stmt*. The following syntax definition extends the Fortran 95 syntax rules R1102 for *program-stmt*, R1105 for *module-stmt*, and R1111 for *block-data-stmt*.

6 7	H605 progr	ram-stmt	is	[ extrinsic-prefix ] PROGRAM program-name
8	H606 modu	ıle-stmt	is	[ extrinsic-prefix ] MODULE module-name
9 10 11	H607 block	e-data-stmt	is	[ extrinsic-prefix ] BLOCK DATA [ block-data-name ]
12 13 14	Constraint:		тоа	any HPF <i>module</i> must be of the same extrinsic <i>lule-subprogram</i> whose extrinsic kind is not given that extrinsic kind.
15 16 17 18 19	Constraint:	must be of the same ext	rinsi	f any HPF main-program or module-subprogram ic kind as its host, and any internal-subprogram iven explicitly is assumed to be of that extrinsic
20 21	6.2.3 Th	e EXTRINSIC Prefix	x	
22 23	H608 extri	nsic-prefix	is	EXTRINSIC ( extrinsic-spec )
24 25	H609 $extri$	nsic-spec	is or	extrinsic-spec-arg-list extrinsic-kind-keyword
26 27 28	H610 extri	nsic-spec-arg	is or or	language model external-name
29 30 31	H611 langu	lage	is	[ LANGUAGE = ] scalar-char-initialization-expr
32 33	H612 mode	el	is	[ MODEL = ] scalar-char-initialization-expr
34 35 36	H613 exter	nal-name	is	[ EXTERNAL_NAME = ] scalar-char-initialization-expr
37 38 39	Constraint:			at least one of <i>language</i> , <i>model</i> , or <i>external-name</i> nay be specified more than once.
40 41 42 43 44 45 46	Constraint:	the <i>extrinsic-spec-arg-list</i> out LANGUAGE= must be th <i>extrinsic-spec-arg-list</i> . If	f. If <i>n</i> he fin <i>exte</i> GE=	ut LANGUAGE=, <i>language</i> must be the first item in <i>model</i> is specified without MODEL=, <i>language</i> witherst item and <i>model</i> must be the second item in the <i>rnal-name</i> is specified without EXTERNAL_NAME=, must be the first item and <i>model</i> without MODEL= he <i>extrinsic-spec-arg-list</i> .
47 48	Constraint:	The forms with LANGUAG order except as prohibite		MODEL=, and EXTERNAL_NAME= may appear in any pove.

		<i>insic-spec-arg-list</i> are as if EXTRINSIC were a pro- face with a <i>dummy-arg-list</i> of LANGUAGE, MODEL,	1 2					
	EXTERNAL_NAME, each of whi		3					
Constraint.	In language values of scalar	-char-initialization-expr may be:	4					
	In language, values of scalar	<i>char-initialization-expr</i> may be.	5 6					
	• 'HPF', referring to the I	IPF language; if a <i>model</i> is not explicitly specified,	7					
	the <i>model</i> is implied to	be 'GLOBAL';	8					
	, 0	b the ANSI/ISO standard Fortran language; if a specified, the <i>model</i> is implied to be 'SERIAL';	9 10					
	, 0	former ANSI/ISO standard FORTRAN 77 lan- t explicitly specified, the <i>model</i> is implied to be	11 12 13					
	, 0	ISI standard C programming language; if a <i>model</i> d, the <i>model</i> is implied to be 'SERIAL'; or	14 15 16					
	• an implementation-dep implied <i>model</i> .	endent value with an implementation-dependent	17 18 19					
	· · ·	entations, 'C' will only be allowed for <i>function</i> - occurring in an <i>interface-body</i> .	20 21					
Constraint:	If language is not specified i	is the same as that of the host scoping unit.	22 23					
Constraint:	In model, values of scalar-ch	ar-initialization-expr may be:	24 25					
	• 'GLOBAL', referring to	the global model,	26 27					
	• 'LOCAL', referring to the table $\bullet$	ne local model,	28					
	• 'SERIAL', referring to	the serial model, or	29					
	• an implementation-dep	endent value.	30 31					
Constraint:	If <i>model</i> is not specified or in	nplied by the specification of a language, it is the	32					
	same as that of the host sco		33 34					
Constraint.	All languages and models u	have named havin with the three latters UDE are	35					
Constraint.		hose names begin with the three letters HPF are definition by this specification and its successors.	36					
	reperted for present of future	definition by this specification and he successfull	37					
Constraint:		of <i>scalar-char-initialization-expr</i> is a character	38					
	_	string whose use is determined by the extrinsic kind. For example, an extrin-						
	sic kind may use the <i>external-name</i> to specify the name by which the procedure							
		would be known if it were referenced by a C procedure. In such an implementa-						
	tion, a user would expect the compiler to perform any transformations of that name that the C compiler would perform. If <i>external-name</i> is not specified,							
	-	its value is implementation-dependent.						
	r montoaction (		44 45					
H614 extri	nsic-kind-keyword is	HPF	46					
	01	HPF_LOCAL	47					
	01	HPF_SERIAL	48					

- Constraint: EXTRINSIC(HPF) is equivalent to EXTRINSIC('HPF', 'GLOBAL'). In the absence of an *extrinsic-prefix* an HPF compiler interprets a compilation unit
- sence of an *extrinsic-prefix* an HPF compiler interprets a compilation unit
   as if it were of extrinsic kind HPF. Thus, for an HPF compiler, specifying
   EXTRINSIC(HPF) or EXTRINSIC('HPF', 'GLOBAL') is redundant. Such explicit
   specification may, however, be required for use with a compiler that supports
   multiple extrinsic kinds.
- Constraint: EXTRINSIC(HPF\_LOCAL) is equivalent to EXTRINSIC('HPF', 'LOCAL'). A main-program whose extrinsic kind is HPF\_LOCAL behaves as if it were a subroutine of extrinsic kind HPF\_LOCAL that is called with no arguments from a main program of extrinsic kind HPF whose executable part consists solely of that call.
- Constraint: EXTRINSIC(HPF\_SERIAL) is equivalent to EXTRINSIC('HPF', 'SERIAL'). A main-program whose extrinsic kind is HPF\_SERIAL behaves as if it were a subroutine of extrinsic kind HPF\_SERIAL that is called with no arguments from a main program of extrinsic kind HPF whose executable part consists solely of that call.
- <sup>19</sup> Constraint: All *extrinsic-kind-keywords* whose names begin with the three letters HPF are <sup>20</sup> reserved for present or future definition by this specification and its successors.
  - Advice to implementors.

Other *languages* or *models* may be defined and provided by compiler vendors. Although not part of this HPF specification, they are expected to conform to the rules and spirit of HPF extrinsic kinds.

An implementation may place certain restrictions on the programmer; moreover, each
 extrinsic kind may call for a different set of restrictions.

For example, an implementation on a parallel processor may find it convenient to replicate scalar arguments so as to provide a copy on every processor. This is permitted so long as this process is invisible to the caller. One way to achieve this is to place a restriction on the programmer: on return from the subprogram, all the copies of this scalar argument must have the same value. This implies that if the dummy argument has INTENT (OUT), then all copies must have been updated consistently by the time of subprogram return.

- (End of advice to implementors.)
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# 6.3 Calling HPF Extrinsic Subprograms

A call to an extrinsic procedure behaves, as observed by a calling program coded in HPF, exactly as if the subprogram were coded in HPF. If a function or subroutine called from a program unit of an HPF extrinsic kind does not have an explicit interface visible in the caller, it is assumed to have the same extrinsic kind as the caller.

In order to call a subprogram of an extrinsic kind other than that of the caller, that subprogram must have an explicit interface visible in the caller, and the subprogram is expected to behave, as observed by the caller, roughly as if it had been written as code of the same extrinsic kind as the caller. Some of the responsibility for meeting this requirement

			Exte	rinsic	kind	l of	the us	ed m	odu	e
			HPF		HPF_SERIAL		HPF_LOCAL		CAL	
Extrinsic kind	HPF	Т	Ρ	D	Т	Ρ		Т	Ρ	
of the using	HPF_SERIAL	Т			Т	Ρ	D	Т		
program unit	HPF_LOCAL	Т			Т			Т	Р	D

 $\mathbf{T} =$ derived type definitions

 $\mathbf{P} =$ procedures and procedure interfaces

 $\mathbf{D} = \text{data objects}$ 

Table 6.1: Entities that a using program unit is entitled to access from a module, according to the HPF extrinsic kind of each.

may rest on the compiler and some on the programmer. This interface defines the "HPF view" of the extrinsic procedure.

A called procedure that is written in a model or language other than HPF, whether or not it uses the local procedure execution model, should be declared **EXTRINSIC** within an HPF program that calls it. The **EXTRINSIC** prefix declares what sort of interface should be used when calling indicated subprograms. If there is no extrinsic specification, then the users must assume full responsibility for correctness of the implementation-dependent interface.

A function-stmt or subroutine-stmt that appears within an interface-block within a program unit of an HPF extrinsic kind may have an extrinsic prefix mentioning any extrinsic in kind supported by the language implementation. If no extrinsic-prefix appears in such a function-stmt or subroutine-stmt, then it is assumed to be of the same HPF extrinsic kind as the program unit in which the interface block appears. 27

The procedure characteristics defined by an *interface-body* must be consistent with the procedure's definition.

The definition and rules for a procedure with an extrinsic interface lies outside the scope of HPF. However, explicit interfaces to such procedures must conform to HPF. Note that any particular HPF implementation is free to support any selection of extrinsic kinds, or none at all except for HPF itself, which clearly must be supported by an HPF implementation.

### 6.3.1 Access to Types, Procedures, and Data

In general, program units of a given extrinsic kind may use names of types, procedures, or data of another program unit of the same extrinsic kind, subject to the scoping rules of Fortran.

Use of names of types, procedures, or data of another program unit of a different extrinsic kind are subject to additional restrictions summarized in Table 6.1 and described below.

Note that, if a module X of one HPF extrinsic kind is used by a program unit Y of another HPF extrinsic kind, then only names of items in X that Y is entitled to use or invoke may be use associated; that is, either X must make private all items that Y is not entitled to use, or the USE statement in Y must have an ONLY option that lists only names of items it is entitled to use.

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# 6.3.1.1 Types

Derived type definitions without explicitly mapped components may be thought of as "extrinsic kind neutral"; a program unit of any HPF extrinsic kind may use derived type definitions from a module of any HPF extrinsic kind. Note that an Approved Extension permits the mapping of components of derived types.

# 6.3.1.2 Procedures

An HPF global program or procedure may call other HPF procedures that are global, local, or serial.

An HPF local program or procedure may call only other HPF local procedures and not HPF global or serial procedures.

An HPF serial program or procedure may call only other HPF serial procedures and not HPF global or local procedures.

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# 6.3.1.3 Data

A named COMMON block in any program unit of an HPF kind will be associated with the COMMON block, if any, of that same name in every other program unit of that same extrinsic kind; similarly for unnamed COMMON. (Such COMMON storage behaves like other declared data objects within program units of that extrinsic kind; in particular, for HPF\_LOCAL code there will be one copy of the COMMON block on each processor.)

It is not permitted for any given COMMON block name to be used in program units of different HPF kinds within a single program; similarly, it is not permitted for unnamed COMMON to be used in program units of different HPF kinds within a single program.

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# 6.3.2 The Effect of a Call

A call to an extrinsic procedure must be semantically equivalent to a call of an ordinary HPF procedure that does not remap its arguments. Thus a call to an extrinsic procedure must behave *as if* the following actions occur. The HPF technical term *as if* means that the described actions should appear to a user as if they occurred, in the order specified; an implementation may carry out any actions in any order that provide the correct user-visible effects.

- 1. All actions of the caller preceding the subprogram invocation should be completed before any action of the subprogram is executed; and all actions of the subprogram should be completed before any action of the caller following the subprogram invocation is executed.
- 2. Each actual argument is remapped, if necessary, according to the directives (explicit or implicit) in the declared interface for the extrinsic procedure. Thus, HPF mapping directives appearing in the interface are binding—the compiler must obey these directives in calling local extrinsic procedures. As in the case of non-extrinsic subprograms, actual arguments may be mapped in any way; if necessary, they are copied automatically to correctly mapped temporaries before invocation of—and copied back to the actual arguments after return from—the extrinsic procedure. Scalar dummy

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arguments and scalar function results behave as if they are replicated on each processor. These mappings may, optionally, be explicit in the interface, but any other explicit mapping is not HPF conforming.

- 3. IN, OUT, and INOUT intent restrictions should be observed.
- 4. No HPF variable is modified unless it could be modified by an HPF procedure with the same explicit interface. Note that even though HPF\_LOCAL and HPF\_SERIAL routines are not permitted to access and modify HPF global data, other kinds of extrinsic routines may do so to the extent that an HPF procedure could.
- 5. When a procedure returns and the caller resumes execution, all objects accessible to the caller after the call are mapped exactly as they were before the call. In particular, the original distribution of arguments is restored, if necessary.
- 6. Exactly the same set of processors is visible to the HPF environment before and after the subprogram call.

Advice to implementors.

To ensure that all actions that logically precede the call are completed, multiple processors may need to be synchronized before the call is made.

If a variable accessible to the called routine has a replicated representation, then all copies may need to be updated prior to the call to contain the correct current value according to the sequential semantics of the source program.

Replicated variables, if updated in the procedure, must be updated consistently. More precisely, if a variable accessible to a procedure has a replicated representation and is updated by (one or more copies of) the procedure, then all copies of the replicated variable must have identical values when the last processor returns from the local procedure.

An implementation might check, before returning from the local subprogram, to make sure that replicated variables have been updated consistently by the subprogram. Note, however, that there is no requirement for an implementation to do so; it is merely an implementation tradeoff between speed and, for instance, debuggability.

Note that, as with a global HPF subprogram, actual arguments may be copied or remapped in any way, so long as the effect is undone on return from the subprogram.

To ensure that all actions of the procedure logically complete before execution in the caller is resumed, multiple processors may need to be synchronized after the call.

(End of advice to implementors.)

## 6.4 Examples of Extrinsic Procedures

Consider:

```
PROGRAM DUMPLING
INTERFACE
EXTRINSIC('HPF','LOCAL') SUBROUTINE GNOCCHI(P, L, X)
```

```
INTERFACE
1
                 SUBROUTINE P(Q)
2
3
                   REAL Q
                 END SUBROUTINE P
4
                 EXTRINSIC('COBOL', 'LOCAL') SUBROUTINE L(R)
5
                   REAL R(:,:)
6
                 END SUBROUTINE L
7
              END INTERFACE
8
              REAL X(:)
9
            END SUBROUTINE GNOCCHI
10
            EXTRINSIC('HPF', 'LOCAL') SUBROUTINE POTSTICKER(Q)
11
              REAL Q
12
            END SUBROUTINE POTSTICKER
13
            EXTRINSIC('COBOL','LOCAL') SUBROUTINE LEBERKNOEDEL(R)
14
              REAL R(:,:)
15
            END SUBROUTINE LEBERKNOEDEL
16
          END INTERFACE
17
18
          . . .
          CALL GNOCCHI(POTSTICKER, LEBERKNOEDEL, (/ 1.2, 3.4, 5.6 /) )
19
20
       END PROGRAM DUMPLING
21
22
          The main program, DUMPLING, when compiled by an HPF compiler, is implicitly of ex-
23
     trinsic kind HPF. Interfaces are declared to three external subroutines GNOCCHI, POTSTICKER,
24
     and LEBERKNOEDEL. The first two are of extrinsic kind HPF_LOCAL and the third is of an ex-
25
     trinsic kind specified by the language COBOL and the local model. Now, GNOCCHI accepts
26
     two dummy procedure arguments and so interfaces must be declared for those. Because
27
     no extrinsic-prefix is given for dummy argument P, its extrinsic kind is that of its host
28
     scoping unit, the declaration of subroutine GNOCCHI, which has extrinsic kind HPF_LOCAL.
29
     The declaration of the corresponding actual argument POTSTICKER needs to have an explicit
30
     extrinsic-prefix because its host scoping unit is program DUMPLING, of extrinsic kind HPF.
31
          Here are some more examples. In the first example, note that the declaration of the
32
     explicit size of BAGEL as 100 refers to its global size and not its local size:
33
34
       INTERFACE
35
          EXTRINSIC('HPF','LOCAL') FUNCTION BAGEL(X)
36
            REAL BAGEL(100)
37
            REAL X(:)
38
               !HPF$ DISTRIBUTE (CYCLIC) :: BAGEL, X
39
          END FUNCTION
40
       END INTERFACE
41
42
         In the next example, note that the ALIGN statement asserts that X, Y, and Z all have
43
     the same shape:
44
       INTERFACE OPERATOR (+)
45
          EXTRINSIC('C', 'LOCAL') FUNCTION LATKES(X, Y) RESULT(Z)
46
            REAL, DIMENSION(:,:), INTENT(IN) :: X
47
            REAL, DIMENSION(:,:), INTENT(IN) :: Y
48
```

REAL, DIMENSION(SIZE(X,1), SIZE(X,2)	) :: Z	1
!HPF\$ ALIGN WITH X :: Y, Z		2
!HPF\$ DISTRIBUTE (BLOCK, BLOCK) X		3
		4
END INTERFACE		5
In the interface block in this final example,	two optornal procedures one of them	6
extrinsic and one not, are associated with the same		7
a scalar of the same type as its array argument:	e generic procedure name, which returns	8
a scalar of the same type as its array argument.		9
INTERFACE KNISH		10 11
FUNCTION RKNISH(X)	!normal HPF interface	11
REAL X(:), RKNISH		13
END RKNISH		14
EXTRINSIC('SISAL') FUNCTION CKNISH(X)	!extrinsic interface	15
COMPLEX X(:), CKNISH		16
END CKNISH		17
END INTERFACE		18
		19
		20
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Section 7

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# **Intrinsic and Library Procedures**

HPF includes Fortran's intrinsic procedures. It also adds new intrinsic procedures in two categories: system inquiry intrinsic functions and computational intrinsic functions.

In addition to the new intrinsic functions, HPF defines a library module, HPF\_LIBRARY, that must be provided by vendors of any full HPF implementation.

This description of HPF intrinsic and library procedures follows the form and conventions of the Fortran standard. The material of Sections F95:13.1, F95:13.2, F95:13.3, F95:13.5.7, F95:13.8.1, F95:13.8.2, F95:13.9, and F95:13.10 is applicable to the HPF intrinsic and library procedures and to their descriptions in this section of the HPF document.

# 7.1 Notation

In the examples of this section, T and F are used to denote the logical values true and false.

## 7.2 System Inquiry Intrinsic Functions

In a multi-processor implementation, the processors may be arranged in an implementation-dependent multi-dimensional processor array. The system inquiry functions return values related to this underlying machine and processor configuration, including the size and shape of the underlying processor array. NUMBER\_OF\_PROCESSORS returns the total number of processors available to the program or the number of processors available to the program along a specified dimension of the processor array. PROCESSORS\_SHAPE returns the shape of the processor array.

The Fortran definition of restricted expression is extended to permit references to the HPF system inquiry intrinsic functions. In particular, at the end of the numbered list in Section 7.1.6.2 of the Fortran standard, add:

(13) A reference to one of the system inquiry functions NUMBER\_OF\_PROCESSORS or
 PROCESSORS\_SHAPE, where any argument is a restricted expression.

A variable that appears in a restricted expression in an HPF directive in the scoping unit of a module or main program must be an implied-DO variable or an argument in a reference to an array inquiry function, bit inquiry function, character inquiry function, kind inquiry function, or numeric inquiry function.

The values returned by the system inquiry intrinsic functions remain constant for the duration of one program execution. Thus, NUMBER\_OF\_PROCESSORS and PROCESSORS\_SHAPE have values that are restricted expressions and may be used wherever any other Fortran

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restricted expression may be used. In particular, NUMBER\_OF\_PROCESSORS may be used in a specification expression.

The values of system inquiry functions may not occur in initialization expressions, because they may not be assumed to be constants. In particular, HPF programs may be compiled to run on machines whose configurations are not known at compile time.

Note that the system inquiry functions query the physical machine, and have nothing to do with any **PROCESSORS** directive that may occur. If an HPF program is running on a physical partition of a larger machine, then it is the smaller partition that actually executes the HPF program whose parameters are returned by the system inquiry functions.

Some machines may not have a "natural" shape to return as the value of the function PROCESSORS\_SHAPE, for example, a machine with a tree topology. In these cases, the implementation must provide some reasonable, consistent description of the machine, such as an rank-one array of size NUMBER\_OF\_PROCESSORS(). The compiler will also have to arrange to map between this description and the underlying hardware processor identification mechanism.

Advice to users. SIZE(PROCESSORS\_SHAPE()) returns the rank of the processor array. References to system inquiry functions may occur in array declarations and in HPF directives, as in:

INTEGER, DIMENSION(SIZE(PROCESSORS\_SHAPE())) :: PSHAPE
!HPF\$ TEMPLATE T(100, 3\*NUMBER\_OF\_PROCESSORS())

(End of advice to users.)

## 7.3 Computational Intrinsic Functions

HPF adds one new computational intrinsic function, ILEN, which computes the number of bits needed to store an integer value.

## 7.4 Library Procedures

The mapping inquiry subroutines and computational functions described in this section are available in the HPF library module, HPF\_LIBRARY. Use of these procedures must be accompanied by an appropriate USE statement in each scoping unit in which they are used. They are not intrinsic.

# 7.4.1 Mapping Inquiry Subroutines

HPF provides data mapping directives that are advisory in nature. The mapping inquiry 43 subroutines allow the program to determine the actual mapping of an array at run time. It 44 may be especially important to know the exact mapping when an EXTRINSIC subprogram is 45 invoked. For these reasons, HPF includes mapping inquiry subroutines which describe how 46 an array is actually mapped onto a machine. To keep the number of routines small, the 47 inquiry procedures are structured as subroutines with optional INTENT (OUT) arguments. 48

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#### **Bit Manipulation Functions** 7.4.2

The HPF library includes three elemental bit-manipulation functions. LEADZ computes the number of leading zero bits in an integer's representation. POPCNT counts the number of one bits in an integer. POPPAR computes the parity of an integer.

#### 7.4.3**Array Reduction Functions**

8 HPF adds additional array reduction functions that operate in the same manner as the 9 Fortran SUM and ANY intrinsic functions. The new reduction functions are IALL, IANY, 10 **IPARITY**, and **PARITY**, which correspond to the commutative, associative binary operations  $1\,1$ IAND, IOR, IEOR, and .NEQV. respectively.

12In the specifications of these functions, the terms "XXX reduction" are used, where XXX 13 is one of the binary operators above. These are defined by means of an example. The IAND 14reduction of all the elements of **array** for which the corresponding element of **mask** is true 15is the scalar integer computed in **result** by

```
16
       result = IAND_IDENTITY_ELEMENT
18
       DO i_1 = LBOUND(array,1), UBOUND(array,1)
           DO i_n = LBOUND(array,n), UBOUND(array,n)
20
             IF ( mask(i_1,i_2,...,i_n) ) &
               result = IAND( result, array(i_1,i_2,...,i_n) )
           END DO
         . . .
^{25}
       END DO
```

Here, n is the rank of **array** and **IAND\_IDENTITY\_ELEMENT** is the integer which has all bits 28 equal to one. (The interpretation of an integer as a sequence of bits is given in F95:13.5.7.) 29 The other three reductions are similarly defined. The identity elements for IOR and IEOR 30 are zero. The identity element for .NEQV. is .FALSE. 31

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#### 7.4.4Array Combining Scatter Functions

The XXX\_SCATTER functions are generalized array reduction functions in which an arbi-35 trary subset of the elements of an array can be combined to produce an element of the 36 result; the subset corresponding to the result's elements are nonoverlapping. Each of the 37 eleven reduction operation in the language corresponds to one of the scatter functions, 38 while COPY\_SCATTER supports overwriting an existing value with any one of the values in 39 the corresponding subset. The way that elements of the source array are associated with 40 the elements of the result is described in this section; the method of combining their values 41 is described in the specifications of the individual functions in Section 7.7. 42 These functions have the general form

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XXX\_SCATTER(ARRAY, BASE, INDX1, ..., INDXn, MASK)

except in the special cases noted below. The allowed values of XXX are ALL, ANY, COPY, 46 COUNT, IALL, IANY, IPARITY, MAXVAL, MINVAL, PARITY, PRODUCT, and SUM. ARRAY, MASK, 47and all the INDX arrays are conformable. The INDX arrays are integer, and the number 48

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of INDX arguments must equal the rank of BASE. The argument MASK is logical, and it is optional. Except for COUNT\_SCATTER, ARRAY and BASE are arrays of the same type. For COUNT\_SCATTER, ARRAY is of type logical and BASE is of type integer. (For ALL\_SCATTER, ANY\_SCATTER, COUNT\_SCATTER, and PARITY\_SCATTER, the ARRAY argument must be logical. These functions do not have an optional MASK argument. To conform with the conventions of the Fortran standard, the required ARRAY argument to these functions is called MASK in their specifications in Section 7.7.) In all cases the result array is an array with the same type, kind type parameter, and shape as BASE.

For every element a in ARRAY there is a corresponding element in each of the INDX arrays, since they all have the same shape as ARRAY. For each j = 1, 2, ..., n, where n is the rank of BASE, let  $s_j$  be the value of the element of INDXj that corresponds to element a in ARRAY. These indices determine the element of the result that is affected by element a of ARRAY. For each of the indices  $s_j$ , let the corresponding index for BASE be given by  $b_j = s_j + LBOUND(BASE, j) - 1$ .

The integers  $b_j$ , j = 1, ..., n, form a subscript selecting an element of BASE: BASE $(b_1, b_2, ..., b_n)$ . Because BASE and the result are conformable, for each element of BASE there is a corresponding element of the result.

Thus the INDX arrays establish a mapping from all the elements of ARRAY onto selected elements of the result and BASE. Viewed in the other direction, this mapping associates with each element b of BASE a set S of elements from ARRAY.

If S is empty, then the element of the result corresponding to the element b of BASE has the same value as b.

If S is non-empty, then the elements of S will be combined with element b to produce an element of the result. The detailed specifications of the scatter functions describe the particular means of combining these values. As an example, for SUM\_SCATTER, if the elements of S are  $a_1, \ldots, a_m$ , then the element of the result corresponding to the element b of BASE is the result of evaluating SUM(( $a_1, a_2, \ldots, a_m, b/$ )).

Note that the elements of the INDX arrays must be non-negative, and that INDXj may not exceed SIZE(BASE, j). The result computed is not affected by the declared upper or lower bounds on indices of BASE; it depends only on the shape of BASE.

Note that, since a scalar is conformable with any array, a scalar may be used in place of an INDX array, in which case one hyperplane of the result is selected. See the example below.

If the optional, final MASK argument is present, then only the elements of ARRAY in positions for which MASK is true participate in the operation. All other elements of ARRAY and of the INDX arrays are ignored and cannot have any influence on any element of the result.

For example, if

A is the array
$$\begin{bmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \\ 7 & 8 & 9 \end{bmatrix}$$
;B is the array $\begin{bmatrix} -1 & -2 & -3 \\ -4 & -5 & -6 \\ -7 & -8 & -9 \end{bmatrix}$ ;I1 is the array $\begin{bmatrix} 1 & 1 & 1 \\ 2 & 1 & 1 \\ 3 & 2 & 1 \end{bmatrix}$ ;I2 is the array $\begin{bmatrix} 1 & 2 & 3 \\ 1 & 1 & 2 \\ 1 & 1 & 1 \end{bmatrix}$ ;

then

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```
SUM_SCATTER(A, B, I1, I2) is \begin{bmatrix} 14 & 6 & 0 \\ 8 & -5 & -6 \\ 0 & -8 & -9 \end{bmatrix};

SUM_SCATTER(A, B, 2, I2) is \begin{bmatrix} -1 & -2 & -3 \\ 30 & 3 & -3 \\ -7 & -8 & -9 \end{bmatrix};

SUM_SCATTER(A, B, I1, 2) is \begin{bmatrix} -1 & 24 & -3 \\ -4 & 7 & -6 \\ -7 & -1 & -9 \end{bmatrix};

SUM_SCATTER(A, B, 2, 2) is \begin{bmatrix} -1 & -2 & -3 \\ -4 & 40 & -6 \\ -7 & -8 & -9 \end{bmatrix}

If A is the array \begin{bmatrix} 10 & 20 & 30 & 40 & -10 \\ -7 & -8 & -9 \end{bmatrix}

If A is the array \begin{bmatrix} 10 & 20 & 30 & 40 & -10 \\ 3 & 2 & 2 & 1 & 1 \end{bmatrix},

then SUM_SCATTER(A, B, IND, MASK=(A .GT. 0)) is \begin{bmatrix} 41 & 52 & 13 & 4 \end{bmatrix}.
```

## 7.4.5 Array Prefix and Suffix Functions

In a scan of a vector, each element of the result is a function of the elements of the vector
 that precede it (for a prefix scan) or that follow it (for a suffix scan). These functions
 provide scan operations on arrays and subarrays. The functions have the general form

```
XXX_PREFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)
XXX_SUFFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)
```

except in the special cases noted below. The allowed values of XXX are ALL, ANY, COPY, COUNT, IALL, IANY, IPARITY, MAXVAL, MINVAL, PARITY, PRODUCT, and SUM.

When comments below apply to both prefix and suffix forms of the routines, we will refer to them as YYYFIX functions.

The arguments DIM, MASK, SEGMENT, and EXCLUSIVE are optional. The COPY\_YYYFIX functions do not have MASK or EXCLUSIVE arguments. The ALL\_YYYFIX, ANY\_YYYFIX, COUNT\_-YYYFIX, and PARITY\_YYYFIX functions do not have MASK arguments. Their ARRAY argument must be of type logical; it is denoted MASK in their specifications in Section 7.7.

The arguments MASK and SEGMENT must be of type logical. SEGMENT must have the same shape as ARRAY. MASK must be conformable with ARRAY. EXCLUSIVE is a logical scalar. DIM is a scalar integer between one and the rank of ARRAY.

**Result Value.** The result has the same shape as ARRAY, and, with the exception of COUNT\_YYYFIX, the same type and kind type parameter as ARRAY. (The result of COUNT\_YYYFIX is default integer.)

In every case, every element of the result is determined by the values of certain selected elements of ARRAY in a way that is specific to the particular function and is described in its specification. The optional arguments affect the selection of elements of ARRAY for each element of the result; the selected elements of ARRAY are said to contribute to the result element. This section describes fully which elements of ARRAY contribute to a given element of the result.

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If no elements of ARRAY are selected for a given element of the result, that result element is set to a default value that is specific to the particular function and is described in its specification.

For any given element r of the result, let a be the corresponding element of ARRAY. Every element of ARRAY contributes to r unless disqualified by one of the following rules.

- 1. If the function is  $XXX\_PREFIX$ , no element that follows a in the array element ordering of ARRAY contributes to r. If the function is  $XXX\_SUFFIX$ , no element that precedes a in the array element ordering of ARRAY contributes to r.
- 2. If the DIM argument is provided, an element z of ARRAY does not contribute to r unless all its indices, excepting only the index for dimension DIM, are the same as the corresponding indices of a. (It follows that if the DIM argument is omitted, then ARRAY, MASK, and SEGMENT are processed in array element order, as if temporarily regarded as rank-one arrays. If the DIM argument is present, then a family of completely independent scan operations are carried out along the selected dimension of ARRAY.)
- 3. If the MASK argument is provided, an element z of ARRAY contributes to r only if the element of MASK corresponding to z is true. (It follows that array elements corresponding to positions where the MASK is false do not contribute anywhere to the result. However, the result is nevertheless defined at all positions, even positions where the MASK is false.)
- 4. If the SEGMENT argument is provided, an element z of ARRAY does not contribute if there is some intermediate element w of ARRAY, possibly z itself, with all of the following properties:
  - (a) If the function is XXX\_PREFIX, w does not precede z but does precede a in the array element ordering; if the function is XXX\_SUFFIX, w does not follow z but does follow a in the array element ordering;
  - (b) If the DIM argument is present, all the indices of w, excepting only the index for dimension DIM, are the same as the corresponding indices of a; and
  - (c) The element of SEGMENT corresponding to w does not have the same value as the element of SEGMENT corresponding to a. (In other words, z can contribute only if there is an unbroken string of SEGMENT values, all alike, extending from z through a.)
- 5. If the EXCLUSIVE argument is provided and is true, then a itself does not contribute to r.

These general rules lead to the following important cases:

- Case (i): If ARRAY has rank one, element i of the result of XXX\_PREFIX(ARRAY) is determined by the first i elements of ARRAY; element SIZE(ARRAY) -i+1 of the result of XXX\_SUFFIX(ARRAY) is determined by the last i elements of ARRAY.
- Case (ii): If ARRAY has rank greater than one, then each element of the result of 46 XXX\_PREFIX(ARRAY) has a value determined by the corresponding element 47 a of the ARRAY and all elements of ARRAY that precede a in array element 48

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	order. For XXX_SUFFIX, $a$ is determined by the elements of ARRAY that correspond to or follow $a$ in array element order.
Case (iii):	Each element of the result of XXX_PREFIX(ARRAY,MASK=MASK) is deter- mined by selected elements of ARRAY, namely the corresponding element <i>a</i> of the ARRAY and all elements of ARRAY that precede <i>a</i> in array ele- ment order, but an element of ARRAY may contribute to the result only if the corresponding element of MASK is true. If this restriction results in selecting no array elements to contribute to some element of the result, then that element of the result is set to the default value for the given function.
Case (iv):	Each element of the result of XXX_PREFIX(ARRAY,DIM=DIM) is determined by selected elements of ARRAY, namely the corresponding element <i>a</i> of the ARRAY and all elements of ARRAY that precede <i>a</i> along dimension DIM; for example, in SUM_PREFIX(A(1:N,1:N), DIM=2), result element $(i_1, i_2)$ could be computed as SUM(A( $i_1, 1 : i_2$ )). More generally, in SUM_PREFIX(ARRAY, DIM), result element $i_1, i_2, \ldots, i_{DIM}, \ldots, i_n$ could be computed as SUM(ARRAY( $i_1, i_2, \ldots, :i_{DIM}, \ldots, i_n$ )). (Note the colon before $i_{DIM}$ in that last expression.)
Case $(v)$ :	If ARRAY has rank one, then element $i$ of the result of XXX_PREFIX (ARRAY, EXCLUSIVE=.TRUE.) is determined by the first $i - 1$ elements of ARRAY.
Case (vi):	The options may be used in any combination.
Advice to u true to false logical value	; thus a segment is indicated by a maximal contiguous subsequence of like
	F,T,F,F,F,T,F,F,T/) seven segments
(End of adv	ice to users.)
Rationale.	

One existing library delimits the segments by indicating the start of each segment. Another delimits the segments by indicating the *stop* of each segment. Each method has its advantages. There is also the question of whether this convention should change when performing a suffix rather than a prefix. HPF adopts the symmetric representation above. The main advantages of this representation are:

- (A) It is symmetrical, in that the same segment specifier may be meaningfully used for prefix and suffix without changing its interpretation (start versus stop).
- (B) The start-bit or stop-bit representation is easily converted to this form by using PARITY\_PREFIX or PARITY\_SUFFIX. These might be standard idioms for a compiler to recognize:

SUM\_PREFIX(FOO,SEGMENT=PARITY\_PREFIX(START\_BITS)) SUM\_PREFIX(FOO,SEGMENT=PARITY\_SUFFIX(STOP\_BITS)) SUM\_SUFFIX(FOO,SEGMENT=PARITY\_SUFFIX(START\_BITS)) SUM\_SUFFIX(FOO,SEGMENT=PARITY\_PREFIX(STOP\_BITS)) (End of rationale.) **Examples.** The examples below illustrate all possible combinations of optional arguments for SUM\_PREFIX. The default value for SUM\_YYYFIX is zero.  $1\,1$ Case (i): SUM\_PREFIX((/1,3,5,7/)) is  $\begin{bmatrix} 1 & 4 & 9 & 16 \end{bmatrix}$ . Case (ii): If B is the array  $\begin{vmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \\ 7 & 8 & 9 \end{vmatrix}$ , then SUM\_PREFIX(B) is the array  $\begin{bmatrix} 1 & 14 & 30 \\ 5 & 19 & 36 \\ 12 & 27 & 45 \end{bmatrix}$ Case (iii): If A is the array  $\begin{bmatrix} 3 & 5 & -2 & -1 & 7 & 4 & 8 \end{bmatrix}$ , then SUM\_PREFIX(A, MASK = A .LT. 6) is  $\begin{bmatrix} 3 & 8 & 6 & 5 & 5 & 9 & 9 \end{bmatrix}$ . Case (iv): If B is the array  $\begin{bmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \\ 7 & 8 & 9 \end{bmatrix}$ , then SUM\_PREFIX(B, DIM=1) is the array  $\begin{bmatrix} 1 & 2 & 3 \\ 5 & 7 & 9 \\ 12 & 15 & 18 \end{bmatrix}$  and SUM\_PREFIX(B, DIM=2) is the array  $\begin{bmatrix} 1 & 3 & 6 \\ 4 & 9 & 15 \\ 7 & 15 & 24 \end{bmatrix}$ .  $^{24}$  $^{25}$ Case (v): SUM\_PREFIX((/1,3,5,7/), EXCLUSIVE=.TRUE.) is  $\begin{bmatrix} 0 & 1 & 4 & 9 \end{bmatrix}$ .  $Case (vi): If B is the array \begin{bmatrix} 1 & 2 & 3 & 4 & 5 \\ 6 & 7 & 8 & 9 & 10 \\ 11 & 12 & 13 & 14 & 15 \end{bmatrix}, M is the array \begin{bmatrix} T & T & T & F & F & F \\ F & F & T & T & T \\ T & F & T & F & F \end{bmatrix}, and S is the array \begin{bmatrix} T & T & F & F & F \\ F & T & T & F & F \\ T & T & T & T & T \end{bmatrix},$ M is the array then: SUM\_PREFIX(B, DIM=2, MASK=M, SEGMENT=S, EXCLUSIVE=.TRUE.) is 0 11 11 24 SUM\_PREFIX(B, DIM=2, MASK=M, SEGMENT=S, EXCLUSIVE=.FALSE.) is 7 120 0 8 9 19 .  $11 \ 11 \ 24 \ 24 \ 24$  $0 \ 1 \ 3 \ 6 \ 10$ 0 0 0 8 17 SUM\_PREFIX(B, DIM=2, MASK=M, EXCLUSIVE=.TRUE.) is 

1 2 3	SUM_PREFIX(B, DIM=2, MASK=M, EXCLUSIVE=.FALSE.) is	$\left[\begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$
4 5 6		$\begin{bmatrix} 0 & 1 & 0 & 3 & 7 \\ 0 & 0 & 7 & 0 & 9 \\ 0 & 11 & 23 & 36 & 50 \end{bmatrix}.$
7 8 9 10	SUM_PREFIX(B, DIM=2, SEGMENT=S, EXCLUSIVE=.FALSE.) is	$\begin{bmatrix} 1 & 3 & 3 & 7 & 12 \\ 6 & 7 & 15 & 9 & 19 \\ 11 & 23 & 36 & 50 & 65 \end{bmatrix}.$
11 12 13 14	SUM_PREFIX(B, DIM=2, EXCLUSIVE=.TRUE.) is	$\begin{bmatrix} 11 & 23 & 36 & 50 & 65 \end{bmatrix}$ $\begin{bmatrix} 0 & 1 & 3 & 6 & 10 \\ 0 & 6 & 13 & 21 & 30 \\ 0 & 11 & 23 & 36 & 50 \end{bmatrix}.$
15 16 17 18	SUM_PREFIX(B, DIM=2, EXCLUSIVE=.FALSE.) is	$\begin{bmatrix} 1 & 3 & 6 & 10 & 15 \\ 6 & 13 & 21 & 30 & 40 \\ 11 & 23 & 36 & 50 & 65 \end{bmatrix}.$
19 20 21	SUM_PREFIX(B, MASK=M, SEGMENT=S, EXCLUSIVE=.TRUE.) is	$\left[\begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$
22 23 24 25	SUM_PREFIX(B, MASK=M, SEGMENT=S, EXCLUSIVE=.FALSE.) i	$\begin{bmatrix} 1 & 13 & 3 & 4 & 5 \\ 0 & 13 & 8 & 13 & 15 \\ 11 & 13 & 21 & 0 & 0 \end{bmatrix}.$
26 27 28 29	SUM_PREFIX(B, MASK=M, EXCLUSIVE=.TRUE.) is	$\begin{bmatrix} 0 & 12 & 14 & 38 & 51 \\ 1 & 14 & 17 & 42 & 56 \\ 1 & 14 & 25 & 51 & 66 \end{bmatrix}.$
30 31 32 33	SUM_PREFIX(B, MASK=M, EXCLUSIVE=.FALSE.) is	$\left[\begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$
34 35 36 37	SUM_PREFIX(B, SEGMENT=S, EXCLUSIVE=.TRUE.) is	$\left[\begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$
38 39 40	SUM_PREFIX(B, SEGMENT=S, EXCLUSIVE=.FALSE.) is	$\begin{bmatrix} 1 & 13 & 3 & 4 & 5 \\ 6 & 20 & 8 & 13 & 15 \\ 11 & 32 & 21 & 14 & 15 \end{bmatrix}.$
41 42 43 44	SUM_PREFIX(B, EXCLUSIVE=.TRUE.) is	$\left[\begin{array}{ccccccccccc} 0 & 18 & 39 & 63 & 90 \\ 1 & 20 & 42 & 67 & 95 \\ 7 & 27 & 50 & 76 & 105 \end{array}\right].$
45 46 47 48	SUM_PREFIX(B, EXCLUSIVE=.FALSE.) is	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$

#### 7.4.6**Array Sorting Functions** 1 2 HPF includes procedures for sorting multidimensional arrays. The SORT\_UP and SORT\_DOWN 3 functions return sorted arrays; the GRADE\_UP and GRADE\_DOWN functions return sorting per-4 mutations. An array can be sorted along a given axis, or the whole array may be viewed 5as a sequence in array element order. The grade functions use stable sorts, allowing for 6 convenient sorting of structures by major and minor keys. 7 8 9 10 7.5Generic Intrinsic and Library Procedures 111213 For all of the intrinsic and library procedures, the arguments shown are the names that 14must be used for keywords when using the keyword form for actual arguments. Many of the 15argument keywords have names that are indicative of their usage, as is the case in Fortran. 16See Section F95:13.11. 1718 19 20 7.5.1System Inquiry Intrinsic Functions 2122 NUMBER\_OF\_PROCESSORS(DIM) The number of executing processors 23 Optional DIM $^{24}$ PROCESSORS\_SHAPE() The shape of the executing processor array $^{25}$ 26 27 Mapping Inquiry Subroutines 7.5.228 29 30 HPF\_ALIGNMENT(ALIGNEE, LB, UB, STRIDE, AXIS\_MAP, IDENTITY\_MAP, & 31 NCOPIES) 32 Optional LB, UB, STRIDE, AXIS\_MAP, IDENTITY\_MAP, NCOPIES 33 HPF\_DISTRIBUTION(DISTRIBUTEE, AXIS\_TYPE, AXIS\_INFO, PROCESSORS\_RANK, & 34 PROCESSORS\_SHAPE) 35 Optional AXIS\_TYPE, AXIS\_INFO, PROCESSORS\_RANK, PROCESSORS\_SHAPE 36 HPF\_TEMPLATE (ALIGNEE, TEMPLATE\_RANK, LB, UB, AXIS\_TYPE, AXIS\_INFO, & 37 NUMBER\_ALIGNED) 38 Optional TEMPLATE\_RANK, LB, UB, AXIS\_TYPE, AXIS\_INFO, 39 NUMBER\_ALIGNED 40 4142 **Bit Manipulation Functions** 7.5.343 44 ILEN(I) Bit length (intrinsic) 45 LEADZ(I) Leading zeros 46 Number of one bits POPCNT(I) 47POPPAR(I) Parity 48

```
7.5.4 Array Reduction Functions
1
2
      IALL(ARRAY, DIM, MASK)
                                    Bitwise logical AND reduction
3
            Optional DIM, MASK
4
      IANY(ARRAY, DIM, MASK)
                                    Bitwise logical OR reduction
5
            Optional DIM, MASK
6
      IPARITY(ARRAY, DIM, MASK)
                                    Bitwise logical EOR reduction
7
            Optional DIM, MASK
8
      PARITY(MASK, DIM)
                                    Logical EOR reduction
9
            Optional DIM
10
1\,1
             Array Combining Scatter Functions
     7.5.5
12
      ALL_SCATTER(MASK, BASE, INDX1 ..., INDXn)
13
      ANY_SCATTER(MASK, BASE, INDX1, ..., INDXn)
14
      COPY_SCATTER(ARRAY, BASE, INDX1, ..., INDXn, MASK)
15
16
            Optional MASK
      COUNT_SCATTER(MASK, BASE, INDX1, ..., INDXn)
17
      IALL_SCATTER(ARRAY, BASE, INDX1, ..., INDXn, MASK)
18
19
            Optional MASK
      IANY_SCATTER(ARRAY, BASE, INDX1, ..., INDXn, MASK)
20
            Optional MASK
21
      IPARITY_SCATTER(ARRAY, BASE, INDX1, ..., INDXn, MASK)
22
            Optional MASK
23
      MAXVAL_SCATTER(ARRAY, BASE, INDX1, ..., INDXn, MASK)
^{24}
            Optional MASK
^{25}
      MINVAL_SCATTER(ARRAY, BASE, INDX1, ..., INDXn, MASK)
26
            Optional MASK
27
      PARITY_SCATTER(MASK, BASE, INDX1, ..., INDXn)
28
29
      PRODUCT_SCATTER(ARRAY, BASE, INDX1, ..., INDXn, MASK)
            Optional MASK
30
      SUM_SCATTER(ARRAY, BASE, INDX1, ..., INDXn, MASK)
31
            Optional MASK
32
33
            Array Prefix and Suffix Functions
34
     7.5.6
35
      ALL_PREFIX(MASK, DIM, SEGMENT, EXCLUSIVE)
36
            Optional DIM, SEGMENT, EXCLUSIVE
37
      ALL_SUFFIX(MASK, DIM, SEGMENT, EXCLUSIVE)
38
            Optional DIM, SEGMENT, EXCLUSIVE
39
      ANY_PREFIX(MASK, DIM, SEGMENT, EXCLUSIVE)
40
            Optional DIM, SEGMENT, EXCLUSIVE
41
      ANY_SUFFIX(MASK, DIM, SEGMENT, EXCLUSIVE)
42
            Optional DIM, SEGMENT, EXCLUSIVE
43
44
45
46
47
48
```

COPY_PREFIX(ARRAY, DIM, SEGMENT)	1
Optional DIM, SEGMENT	2
COPY_SUFFIX(ARRAY, DIM, SEGMENT)	3
Optional DIM, SEGMENT	4
COUNT_PREFIX(MASK, DIM, SEGMENT, EXCLUSIVE)	5
Optional DIM, SEGMENT, EXCLUSIVE	6
COUNT_SUFFIX(MASK, DIM, SEGMENT, EXCLUSIVE)	7
Optional DIM, SEGMENT, EXCLUSIVE	8
IALL_PREFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)	9
Optional DIM, MASK, SEGMENT, EXCLUSIVE	10
IALL_SUFFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)	11
Optional DIM, MASK, SEGMENT, EXCLUSIVE	12
IANY_PREFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)	13
Optional DIM, MASK, SEGMENT, EXCLUSIVE	14
IANY_SUFFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)	15
Optional DIM, MASK, SEGMENT, EXCLUSIVE	16
IPARITY_PREFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)	17
Optional DIM, MASK, SEGMENT, EXCLUSIVE	18
IPARITY_SUFFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)	19
Optional DIM, MASK, SEGMENT, EXCLUSIVE	20
MAXVAL_PREFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)	21
Optional DIM, MASK, SEGMENT, EXCLUSIVE	22
MAXVAL_SUFFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)	23
Optional DIM, MASK, SEGMENT, EXCLUSIVE	24
MINVAL_PREFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)	25
Optional DIM, MASK, SEGMENT, EXCLUSIVE	26
MINVAL_SUFFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)	27
Optional DIM, MASK, SEGMENT, EXCLUSIVE	28
PARITY_PREFIX(MASK, DIM, SEGMENT, EXCLUSIVE)	29
Optional DIM, SEGMENT, EXCLUSIVE	30
PARITY_SUFFIX(MASK, DIM, SEGMENT, EXCLUSIVE)	31
Optional DIM, SEGMENT, EXCLUSIVE	32
PRODUCT_PREFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)	33
Optional DIM, MASK, SEGMENT, EXCLUSIVE	34
PRODUCT_SUFFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)	35
Optional DIM, MASK, SEGMENT, EXCLUSIVE	36
SUM_PREFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)	37
Optional DIM, MASK, SEGMENT, EXCLUSIVE	38
SUM_SUFFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)	39
Optional DIM, MASK, SEGMENT, EXCLUSIVE	40
	41

7.5.7 Array Sort Fo	inctions
GRADE_DOWN(ARRAY,DIM	) Permutation that sorts into descending order
Optional DIM GRADE_UP(ARRAY,DIM) Optional DIM	Permutation that sorts into ascending order
SORT_DOWN (ARRAY, DIM) Optional DIM	Sort into descending order
SORT_UP(ARRAY,DIM) Optional DIM	Sort into ascending order
7.6 Specifications of	of Intrinsic Procedures
ILEN(I)	
<b>Description</b> . Re representation of a	turns one less than the length, in bits, of the two's-complement integer.
Class. Elemental	function.
Argument. I mu	ast be of type integer.
Result Type and	d Type Parameter. Same as I.
	If I is nonnegative, ILEN(I) has the value $\lceil \log_2(I+1) \rceil$ ; if I has the value $\lceil \log_2(-I) \rceil$ .
-	(4) = 3. ILEN(-4) = 2. 2**ILEN(N-1) rounds N up to a pow hereas 2**(ILEN(N)-1) rounds N down to a power of 2. Comp
of I, as the follow	l is one <i>less</i> than the length of the two's-complement representation ring explains. The shortest two's-complement representation o ring zero is the required sign bit. In 3-bit two's complement, 1
NUMBER_OF_PRO	CESSORS(DIM)
<b>Optional Argun</b>	nent. DIM
=	turns the total number of processors available to the program cessors available to the program along a specified dimension of t
Class. System inc	quiry function.
Arguments.	
$\mathtt{DIM}$ (optional)	must be scalar and of type integer with a value in trange $1 \leq \text{DIM} \leq n$ where <i>n</i> is the rank of the process array.

Result Type, Type Parameter, and Shape. Default integer scalar. 1 2 **Result Value.** The result has a value equal to the extent of dimension DIM of the 3 implementation-dependent hardware processor array or, if DIM is absent, the total 4 number of elements of the implementation-dependent hardware processor array. The 5 result is always greater than zero. 6 7 **Examples.** For a computer with 8192 processors arranged in a 128 by 64 rectangular 8 grid, the value of NUMBER\_OF\_PROCESSORS() is 8192; the value of NUMBER\_OF\_PROCES-9 SORS(DIM=1) is 128; and the value of NUMBER\_OF\_PROCESSORS(DIM=2) is 64. For a 10single-processor workstation, the value of NUMBER\_OF\_PROCESSORS() is 1; since the 11 rank of a scalar processor array is zero, no DIM argument may be used. 12 13 PROCESSORS\_SHAPE() 14 15**Description**. Returns the shape of the implementation-dependent processor array. 16 17**Class.** System inquiry function. 18 Arguments. None 19 20 **Result Type, Type Parameter, and Shape.** The result is a default integer  $^{21}$ array of rank one whose size is equal to the rank of the implementation-dependent 22 processor array. 23  $^{24}$ **Result Value.** The value of the result is the shape of the implementation-dependent  $^{25}$ processor array. 26 27 **Example.** In a computer with 2048 processors arranged in a hypercube, the value 28 of PROCESSORS\_SHAPE() is [2,2,2,2,2,2,2,2,2,2,2,2]. In a computer with 8192 proces-29 sors arranged in a 128 by 64 rectangular grid, the value of PROCESSORS\_SHAPE() is 30 [128,64]. For a single processor workstation, the value of PROCESSORS\_SHAPE() is [] 31 (the size-zero array of rank one). 32 33 7.7**Specifications of Library Procedures** 34 35 ALL\_PREFIX(MASK, DIM, SEGMENT, EXCLUSIVE) 36 37 **Optional Arguments.** DIM, SEGMENT, EXCLUSIVE 38 **Description.** Computes a segmented logical AND scan along dimension DIM of 39 MASK. 40 41**Class.** Transformational function. 42 43 Arguments. 44 45 must be of type logical. It must not be scalar. MASK 46 must be scalar and of type integer with a value in the **DIM** (optional) 47 range 1 < DIM < n, where n is the rank of MASK. 48

1 2		<b>SEGMENT</b> (optional)	must be of type logical and must have the same shape as MASK.	
3 4		EXCLUSIVE (optional)	must be of type logical and must be scalar.	
5		Result Type, Type Parameter, and Shape. Same as MASK.		
7 8 9 10		<b>Result Value.</b> Element r of the result has the value ALL((/ $a_1, \ldots, a_m$ /)) where $(a_1, \ldots, a_m)$ is the (possibly empty) set of elements of MASK selected to contribute to r by the rules stated in Section 7.4.5.		
10 11 12 13		Example. ALL_PREFIX( $\left[ T F F T T \right]$ .	(/T,F,T,T,T/), SEGMENT= (/F,F,F,T,T/) ) is	
14 15	ALL	_SCATTER(MASK,B	ASE,INDX1,, INDXn)	
16 17 18 19		index arrays INDX1,, II	lements of MASK to positions of the result indicated by NDXn. An element of the result is true if and only if the BASE and all elements of MASK scattered to that position	
20 21		Class. Transformational f	function.	
22 23		Arguments.		
24 25		MASK	must be of type logical. It must not be scalar.	
26 27		BASE	must be of type logical with the same kind type parameter as MASK. It must not be scalar.	
28 29 30 31		INDX1,,INDXn	must be of type integer and conformable with MASK. The number of INDX arguments must be equal to the rank of BASE.	
32 33		Result Type, Type Par	rameter, and Shape. Same as BASE.	
34 35 36 37			ent of the result corresponding to the element $b$ of BASE has $(a_m, b/)$ ), where $(a_1, \ldots, a_m)$ are the elements of MASK abed in Section 7.4.4.	
38 39 40		Example. ALL_SCATTER( $\begin{bmatrix} T & F & T \end{bmatrix}$ .	(/T, T, T, F/), (/T, T, T/), (/1, 1, 2, 2/) ) is	
41 42	ALL	_SUFFIX(MASK, DIN	A, SEGMENT, EXCLUSIVE)	
43 44		Optional Arguments.	DIM, SEGMENT, EXCLUSIVE	
45 46		<b>Description.</b> Computes DIM of MASK.	a reverse, segmented logical AND scan along dimension	
47 48		Class. Transformational	function.	

$\mathbf{Arguments.}$	
MASK	must be of type logical. It must not be scalar.
DIM (optional)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of MASK.
<b>SEGMENT</b> (optional)	must be of type logical and must have the same shape as MASK.
EXCLUSIVE (optional)	must be of type logical and must be scalar.
Result Type, Type P	arameter, and Shape. Same as MASK.
	r of the result has the value ALL((/ $a_1, \ldots, a_m$ /)) where bly empty) set of elements of MASK selected to contribute to Section 7.4.5.
Example. ALL_SUFFIX( $\left[ F F T T T \right]$ .	(/T,F,T,T,T/), SEGMENT= (/F,F,F,T,T/) ) is
NY_PREFIX(MASK, D	DIM, SEGMENT, EXCLUSIVE)
<b>Optional Arguments.</b>	DIM, SEGMENT, EXCLUSIVE
<b>Description.</b> Computes	s a segmented logical OR scan along dimension DIM of MASK.
Class. Transformational	l function.
Arguments.	
MASK	must be of type logical. It must not be scalar.
DIM (optional)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of MASK.
<b>SEGMENT</b> (optional)	must be of type logical and must have the same shape as MASK.
EXCLUSIVE (optional)	must be of type logical and must be scalar.
Result Type, Type P	arameter, and Shape. Same as MASK.
	r of the result has the value ANY((/ $a_1, \ldots, a_m$ /)) where bly empty) set of elements of MASK selected to contribute to Section 7.4.5.
	(/F,T,F,F,F/), SEGMENT= (/F,F,F,T,T/) ) is

Example. ANY\_PREFIX( (/F,T,F,F,F/), SEGMENT= (/F,F,F,T,T/) ) is [F T T F F].

47 48

ANY\_SCATTER(MASK,BASE,INDX1, ..., INDXn) 1 2 **Description.** Scatters elements of MASK to positions of the result indicated by 3 index arrays INDX1, ..., INDXn. An element of the result is true if and only if the 4 corresponding element of BASE or any element of MASK scattered to that position is 5 true. 6 Class. Transformational function. 7 8 Arguments. 9 MASK must be of type logical. It must not be scalar. 10 11BASE must be of type logical with the same kind type parameter 12as MASK. It must not be scalar. 13 INDX1,..., INDXn must be of type integer and conformable with MASK. The 14 number of INDX arguments must be equal to the rank of 15BASE. 1617**Result Type, Type Parameter, and Shape.** Same as BASE. 18 **Result Value.** The element of the result corresponding to the element b of **BASE** has 19 the value ANY(  $(/a_1, a_2, ..., a_m, b/)$  ), where  $(a_1, ..., a_m)$  are the elements of MASK 20 associated with b as described in Section 7.4.4. 21 22 Example. ANY\_SCATTER( (/T, F, F, F/), (/F, F, T/), (/1, 1, 2, 2/) ) is 23 TFT. 24  $^{25}$ ANY\_SUFFIX(MASK, DIM, SEGMENT, EXCLUSIVE) 26 **Optional Arguments.** DIM, SEGMENT, EXCLUSIVE 27 28 **Description.** Computes a reverse, segmented logical OR scan along dimension DIM 29 of MASK. 30 Class. Transformational function. 31 32 Arguments. 33 MASK must be of type logical. It must not be scalar. 34 35 must be scalar and of type integer with a value in the DIM (optional) 36 range  $1 \leq \text{DIM} \leq n$ , where n is the rank of MASK. 37 SEGMENT (optional) must be of type logical and must have the same shape as 38 MASK. 39 EXCLUSIVE (optional) must be of type logical and must be scalar. 40 41 **Result Type, Type Parameter, and Shape.** Same as MASK. 42 **Result Value.** Element r of the result has the value  $ANY((/a_1, \ldots, a_m/))$  where 43  $(a_1,\ldots,a_m)$  is the (possibly empty) set of elements of MASK selected to contribute to 44 r by the rules stated in Section 7.4.5. 4546 Example. ANY\_SUFFIX( (/F,T,F,F,F/), SEGMENT= (/F,F,F,T,T/) ) is 47 T T F F F. 48

COF	PY_PREFIX(ARRAY,	DIM, SEGMENT)	1	
	Optional Arguments. DIM, SEGMENT			
	Description. Computes a segmented copy scan along dimension DIM of ARRAY.			
	Class. Transformational function.			
	Arguments.		8 9	
	ARRAY	may be of any type. It must not be scalar.	10	
	DIM (optional)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of ARRAY.	11 12	
	<b>SEGMENT</b> (optional)	must be of type logical and must have the same shape as ARRAY.	13 14 15	
	Result Type, Type Pa	rameter, and Shape. Same as ARRAY.	16 17	
	<b>Result Value.</b> Element r of the result has the value $a_1$ where $(a_1, \ldots, a_m)$ is the set, in array element order, of elements of <b>ARRAY</b> selected to contribute to r by the rules stated in Section 7.4.5. Note that this set is never empty.			
	Example. COPY_PREFIX( (/1,2,3,4,5/), SEGMENT= (/F,F,F,T,T/) ) is [ 1 1 1 4 4 ].			
COF	<b>OPY_SCATTER</b> ( <b>ARRAY</b> , <b>BASE</b> , <b>INDX1</b> ,, <b>INDX</b> n, <b>MASK</b> )			
	Optional Argument. MASK			
	indicated by index arrays	ements of ARRAY selected by MASK to positions of the result s INDX1,, INDXn. Each element of the result is equal to RRAY scattered to that position or, if there is none, to the BASE.	28 29 30 31 32 33	
	Class. Transformational	function.	34	
	Arguments.		35 36	
	ARRAY	may be of any type. It must not be scalar.	37	
	BASE	must be of the same type and kind type parameter as ARRAY.	38 39 40	
	INDX1,,INDXn	must be of type integer and conformable with ARRAY. The number of INDX arguments must be equal to the rank of BASE.	41 42 43 44	
	MASK (optional)	must be of type logical and must be conformable with ARRAY.	44 45 46	
	Result Type Type Pa	rameter and Shape Same as BASE	47	

Result Type, Type Parameter, and Shape. Same as BASE.

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**Result Value.** Let S be the set of elements of ARRAY associated with element b of 1 **BASE** as described in Sector 7.4.4. 2 3 If S is empty, then the element of the result corresponding to the element b of BASE 4 has the same value as b. 5 If S is non-empty, then the element of the result corresponding to the element b of 6 **BASE** is the result of choosing one element from S. HPF does not specify how the 7 choice is to be made; the mechanism is implementation dependent. 8 9 Example. COPY\_SCATTER((/1, 2, 3, 4/), (/7, 8, 9/), (/1, 1, 2, 2/)) is 10 [x, y, 9], where x is a member of the set  $\{1, 2\}$  and y is a member of the set 11  $\{3,4\}.$ 1213 COPY\_SUFFIX(ARRAY, DIM, SEGMENT) 14 15**Optional Arguments.** DIM, SEGMENT 16**Description.** Computes a reverse, segmented copy scan along dimension DIM of 1718 ARRAY. 19 Class. Transformational function. 20 21 Arguments. 22 23 ARRAY may be of any type. It must not be scalar.  $^{24}$ DIM (optional) must be scalar and of type integer with a value in the 25range  $1 \leq \text{DIM} \leq n$ , where n is the rank of ARRAY. 26 27 SEGMENT (optional) must be of type logical and must have the same shape as 28 ARRAY. 29 30 **Result Type, Type Parameter, and Shape.** Same as ARRAY. 31 **Result Value.** Element r of the result has the value  $a_m$  where  $(a_1, \ldots, a_m)$  is the 32 set, in array element order, of elements of ARRAY selected to contribute to r by the 33 rules stated in Section 7.4.5. Note that this set is never empty. 34 35 Example. COPY\_SUFFIX( (/1,2,3,4,5/), SEGMENT= (/F,F,F,T,T/) ) is 36 3 3 3 5 5 . 37 38 39 COUNT\_PREFIX(MASK, DIM, SEGMENT, EXCLUSIVE) 40 **Optional Arguments.** DIM, SEGMENT, EXCLUSIVE 4142 **Description.** Computes a segmented COUNT scan along dimension DIM of MASK. 43 44 Class. Transformational function. 45 46 Arguments. 47

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DIM (optional)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of MASK.	1 2
SEGMENT (optional)	must be of type logical and must have the same shape as MASK.	3 4 5
EXCLUSIVE (optional)	must be of type logical and must be scalar.	6

**Result Type, Type Parameter, and Shape.** The result is of type default integer and of the same shape as MASK.

**Result Value.** Element r of the result has the value COUNT((/  $a_1, \ldots, a_m$  /)) where  $(a_1, \ldots, a_m)$  is the (possibly empty) set of elements of MASK selected to contribute to r by the rules stated in Section 7.4.5.

```
Example. COUNT_PREFIX( (/F,T,T,T,T/), SEGMENT= (/F,F,F,T,T/) ) is
\begin{bmatrix} 0 & 1 & 2 & 1 & 2 \end{bmatrix}.
```

#### COUNT\_SCATTER(MASK, BASE, INDX1, ..., INDXn)

**Description.** Scatters elements of MASK to positions of the result indicated by index arrays INDX1, ..., INDXn. Each element of the result is the sum of the corresponding element of BASE and the number of true elements of MASK scattered to that position.

Class. Transformational function.

#### Arguments.

MASK	must be of type logical. It must not be scalar.	27
BASE	must be of type integer. It must not be scalar.	28 29
INDX1,,INDXn	must be of type integer and conformable with MASK. The number of INDX arguments must be equal to the rank of	30 31
	BASE.	32 33

#### **Result Type, Type Parameter, and Shape.** Same as BASE.

**Result Value.** The element of the result corresponding to the element b of BASE has the value  $b + COUNT((a_1, a_2, ..., a_m))$  ), where  $(a_1, ..., a_m)$  are the elements of MASK associated with b as described in Section 7.4.4.

Example. COUNT\_SCATTER((/T, T, T, F/),(/1, -1, 0/),(/1, 1, 2, 2/)) is 300.

#### COUNT\_SUFFIX(MASK, DIM, SEGMENT, EXCLUSIVE)

**Optional Arguments.** DIM, SEGMENT, EXCLUSIVE

**Description.** Computes a reverse, segmented COUNT scan along dimension DIM of MASK.

1	Class. Transformational function.			
2 3	Argumer	its.		
4 5	MASK		must be of type logical. It must not be scalar.	
6 7	DIM (optio	nal)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of MASK.	
8	SEGMENT (	optional)	must be of type logical and must have the same shape as MASK.	
10 11	EXCLUSIVE	E (optional)	must be of type logical and must be scalar.	
12 13 14	Result Type, Type Parameter, and Shape. The result is of type default integer and of the same shape as MASK.			
15 16 17 18	where $(a_1, a_2)$	$(\ldots, a_m)$ is the	t r of the result has the value COUNT((/ $a_1, \ldots, a_m$ /)) (possibly empty) set of elements of MASK selected to constated in Section 7.4.5.	
19 20 21	$ \begin{array}{c} \mathbf{Example} \\ \begin{bmatrix} 2 & 1 & 1 \end{bmatrix} \end{array} $	-	X( (/T,F,T,T,T/), SEGMENT= (/F,F,F,T,T/) ) is	
$\mathbf{GR}^{22}_{23}$	ADE_DOV	VN(ARRAY	,DIM)	
24	Optional	Optional Argument. DIM		
25 26 27	<b>Description.</b> Produces a permutation of the indices of an array, expressed as one- based coordinates, and sorted by descending array element values.			
28 29	Class. Transformational function.			
30 31	Argumer	nts.		
32 33	ARRAY		must be of type integer, real, or character. It must not be scalar.	
34 35 36 37 38	DIM (optio	nal)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where <i>n</i> is the rank of <b>ARRAY</b> . The corresponding actual argument must not be an optional dummy argument.	
39 40 41 42	If DIM is p	resent, the resu	cameter, and Shape. The result is of type default integer. It has the same shape as ARRAY. If DIM is absent, the result E(ARRAY)), SIZE(ARRAY) /).	
43	Result Value.			
44 45 46	Case (i):	The result of S = GRADE_DO	WN(ARRAY)	
47 48		+ SPRE	CAD (LBOUND(ARRAY), DIM=2, NCOPIES=SIZE(ARRAY))-1 one computes the rank-one array B of size SIZE(ARRAY) by	

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FORALL  $(K=1:SIZE(B)) B(K)=ARRAY(S(1,K),S(2,K),\ldots,S(N,K))$ 1 where N has the value SIZE(SHAPE(ARRAY)), then B is sorted in descending order; moreover, all of the columns of S are distinct, that is, if 3  $j \neq m$  then ALL(S(:,j) .EQ. S(:,m)) will be false. The sort is stable; 4 if  $j \leq m$  and B(j) = B(m), then ARRAY(S(1,j),S(2,j),...,S(n,j)) 5precedes ARRAY (S(1, m), S(2, m), ..., S(n, m)) in the array element or-6 dering of ARRAY. The collating sequence for an array of type CHARACTER 7is that used by the Fortran intrinsic functions, namely ASCII.

# Case (ii): The result of R = GRADE\_DOWN(ARRAY, DIM=K) + LBOUND(ARRAY, DIM=K) -1 has the property that if one computes the array $B(i_1, i_2, ..., i_k, ..., i_n) =$ ARRAY $(i_1, i_2, \ldots, R(i_1, i_2, \ldots, i_k, \ldots, i_n), \ldots, i_n)$ then for all $i_1, i_2, ..., (\text{omit } i_k), ..., i_n$ , the vector $B(i_1, i_2, ..., :, ..., i_n)$ is sorted in descending order; moreover, $R(i_1, i_2, \ldots, :, \ldots, i_n)$ is a permutation of all the integers between 1 and SIZE(ARRAY, DIM=K), inclusive. The sort is stable; that is, if j < m and $B(i_1, i_2, ..., j, ..., i_n) = B(i_1, i_2, ..., m, ..., i_n)$ , then $R(i_1, i_2, \ldots, j, \ldots, i_n) \leq R(i_1, i_2, \ldots, m, \ldots, i_n)$ . The collating sequence for an array of type CHARACTER is that used by the Fortran intrinsic functions, namely ASCII.

#### Examples.

 $^{25}$ Case (i): GRADE\_DOWN( (/30, 20, 30, 40, -10/) ) is a rank two array of shape  $\begin{bmatrix} 1 & 5 \end{bmatrix}$  with the value  $\begin{bmatrix} 4 & 1 & 3 & 2 & 5 \end{bmatrix}$ . (To produce a rank-one result, the optional DIM = 1 argument must be used.) 26 27 28 If **A** is the array  $\begin{bmatrix} 1 & 9 & 2 \\ 4 & 5 & 2 \\ 1 & 2 & 4 \end{bmatrix}$ , 29 30 31 32 then GRADE\_DOWN(A) has the value 33 34 Case (ii): If A is the array  $\begin{bmatrix} 1 & 9 & 2 \\ 4 & 5 & 2 \\ 1 & 2 & 4 \end{bmatrix}$ 35 36 37 38 then GRADE\_DOWN(A, DIM = 1) has the value  $\begin{bmatrix} 2 & 1 & 3 \\ 1 & 2 & 1 \\ 3 & 3 & 2 \end{bmatrix}$ , 39 40 41  $\left[\begin{array}{rrrrr} 2 & 3 & 1 \\ 2 & 1 & 3 \\ 3 & 2 & 1 \end{array}\right].$ 42 and  $GRADE_DOWN(A, DIM = 2)$  has the value 43 44 45 46 GRADE\_UP(ARRAY,DIM)

**Optional Argument.** DIM

**Description.** Produces a permutation of the indices of an array, expressed as one-1 based coordinates, and sorted by ascending array element values. 2 3 Class. Transformational function. 4 5 6 Arguments. 7 must be of type integer, real, or character. It must not ARRAY 8 be scalar. 9 10 DIM (optional) must be scalar and of type integer with a value in the 11range 1 < DIM < n, where n is the rank of ARRAY. The 12corresponding actual argument must not be an optional 13 dummy argument. 14 15**Result Type, Type Parameter, and Shape.** The result is of type default integer. 16If DIM is present, the result has the same shape as ARRAY. If DIM is absent, the result 17has shape (/ SIZE(SHAPE(ARRAY)), SIZE(ARRAY) /). 18 19 Result Value. 20 21 Case (i): The result of 22  $S = GRADE_UP(ARRAY)$ 23 + SPREAD(LBOUND(ARRAY), DIM=2, NCOPIES=SIZE(ARRAY))-1  $^{24}$ is such that if one computes the rank-one array B of size SIZE(ARRAY) by 25FORALL  $(K=1:SIZE(B)) B(K)=ARRAY(S(1,K),S(2,K),\ldots,S(N,K))$ 26 where N has the value SIZE(SHAPE(ARRAY)), then B is sorted in ascending 27 order; moreover, all of the columns of S are distinct, that is, if  $j \neq m$  then 28 ALL(S(:, j) .EQ. S(:, m)) will be false. The sort is stable; if j < m29 and B(j) = B(m), then ARRAY(S(1,j),S(2,j),...,S(n,j)) precedes 30 ARRAY  $(S(1, m), S(2, m), \ldots, S(n, m))$  in the array element ordering of 31 ARRAY. The collating sequence for an array of type CHARACTER is that used 32 by the Fortran intrinsic functions, namely ASCII. 33 *Case (ii):* The result of 34  $R = GRADE_UP(ARRAY, DIM=K) + LBOUND(ARRAY, DIM=K) - 1$ 35 has the property that if one computes the array 36  $B(i_1, i_2, ..., i_k, ..., i_n) =$ 37 ARRAY  $(i_1, i_2, \ldots, R(i_1, i_2, \ldots, i_k, \ldots, i_n), \ldots, i_n)$ 38 then for all  $i_1, i_2, ..., (\text{omit } i_k), ..., i_n$ , the vector  $B(i_1, i_2, ..., :, ..., i_n)$ 39 is sorted in ascending order; moreover,  $R(i_1, i_2, \ldots, :, \ldots, i_n)$  is a permu-40 tation of all the integers between 1 and SIZE(ARRAY, DIM=K), inclusive. 41The sort is stable; that is, if j < m and 42  $B(i_1, i_2, ..., j, ..., i_n) = B(i_1, i_2, ..., m, ..., i_n)$ , then 43  $R(i_1, i_2, \ldots, j, \ldots, i_n) \leq R(i_1, i_2, \ldots, m, \ldots, i_n)$ . The collating sequence 44 for an array of type CHARACTER is that used by the Fortran intrinsic func-45tions, namely ASCII. 46 47Examples. 48

Case (i): GRADE\_UP( (/30, 20, 30, 40, -10/) ) is a rank two array of shape 1  $\begin{bmatrix} 1 & 5 \end{bmatrix}$  with the value  $\begin{bmatrix} 5 & 2 & 1 & 3 & 4 \end{bmatrix}$ . (To produce a rank-one 2 3 result, the optional DIM = 1 argument must be used.) 4 If **A** is the array  $\begin{bmatrix} 1 & 9 & 2 \\ 4 & 5 & 2 \\ 1 & 2 & 4 \end{bmatrix}$ , 56 then GRADE\_UP(A) has the value 10 Case (ii): If **A** is the array  $\begin{vmatrix} 1 & 9 & 2 \\ 4 & 5 & 2 \\ 1 & 2 & 4 \end{vmatrix}$ , 11 1213 then GRADE\_UP(A, DIM = 1) has the value  $\begin{bmatrix} 1 & 3 & 1 \\ 3 & 2 & 2 \\ 2 & 1 & 3 \end{bmatrix}$ , 141516  $\left[\begin{array}{rrrrr} 1 & 3 & 2 \\ 3 & 1 & 2 \\ 1 & 2 & 3 \end{array}\right].$ 17and  $GRADE_UP(A, DIM = 2)$  has the value 18 19 20 21HPF\_ALIGNMENT(ALIGNEE, LB, UB, STRIDE, AXIS\_MAP, IDEN-22 TITY\_MAP, NCOPIES) 23 Optional Arguments. LB, UB, STRIDE, AXIS\_MAP, IDENTITY\_MAP, NCOPIES  $^{24}$  $^{25}$ **Description.** Returns information regarding the correspondence of a variable and 26 the *align-target* (array or template) to which it is ultimately aligned. 27 28 Class. Mapping inquiry subroutine. 29 30 Arguments. 31 32 ALIGNEE may be of any type. It may be scalar or array valued. It 33 must not be an assumed-size array. If it is a member of an 34 aggregate variable group, then it must be an aggregate 35 cover of the group. (See Section 3.8 for the definitions 36 of "aggregate variable group" and "aggregate cover.") It 37 must not be a pointer that is disassociated or an allocat-38 able array that is not allocated. It is an INTENT (IN) 39 argument. 40 If ALIGNEE is a pointer, information about the align-41 ment of its target is returned. The target must not 42 be an assumed-size dummy argument or a section of an 43 assumed-size dummy argument. 44 LB (optional) must be of type default integer and of rank one. Its 45size must be at least equal to the rank of ALIGNEE. It 46 is an INTENT (OUT) argument. The first element of the 47 $i^{th}$  axis of ALIGNEE is ultimately aligned to the LB(i)<sup>th</sup> 48

1 2 3 4		<i>align-target</i> element along the axis of the <i>align-target</i> associated with the $i^{th}$ axis of ALIGNEE. If the $i^{th}$ axis of ALIGNEE is a collapsed axis, LB(i) is implementation dependent.
5 6 7 8 9 10 11 12 13	UB (optional)	must be of type default integer and of rank one. Its size must be at least equal to the rank of ALIGNEE. It is an INTENT (OUT) argument. The last element of the $i^{th}$ axis of ALIGNEE is ultimately aligned to the UB(i) <sup>th</sup> align-target element along the axis of the align-target associated with the $i^{th}$ axis of ALIGNEE. If the $i^{th}$ axis of ALIGNEE is a collapsed axis, UB(i) is implementation dependent.
14 15 16 17 18 19	STRIDE (optional)	must be of type default integer and of rank one. Its size must be at least equal to the rank of ALIGNEE. It is an INTENT (OUT) argument. The i <sup>th</sup> element of STRIDE is set to the stride used in aligning the elements of ALIGNEE along its i <sup>th</sup> axis. If the i <sup>th</sup> axis of ALIGNEE is a collapsed axis, STRIDE(i) is zero.
20 21 22 23 24 25	AXIS_MAP (optional)	must be of type default integer and of rank one. Its size must be at least equal to the rank of ALIGNEE. It is an INTENT (OUT) argument. The i <sup>th</sup> element of AXIS_MAP is set to the <i>align-target</i> axis associated with the i <sup>th</sup> axis of ALIGNEE. If the i <sup>th</sup> axis of ALIGNEE is a collapsed axis, AXIS_MAP(i) is 0.
26 27 28 29 30 31 32 33 34 35 36	IDENTITY_MAP (optional)	must be scalar and of type default logical. It is an INTENT (OUT) argument. It is set to true if the ultimate <i>align-</i> <i>target</i> associated with ALIGNEE has a shape identical to ALIGNEE, the axes are mapped using the identity per- mutation, and the strides are all positive (and therefore equal to 1, because of the shape constraint); otherwise it is set to false. If a variable has not appeared as an <i>alignee</i> in an ALIGN or REALIGN directive, and does not have the INHERIT attribute, then IDENTITY_MAP must be true; it can be true in other circumstances as well.
37 38 39 40	NCOPIES (optional)	must be scalar and of type default integer. It is an INTENT (OUT) argument. It is set to the number of copies of ALIGNEE that are ultimately aligned to <i>align-target</i> . For a non-replicated variable, it is set to one.
41 42 43	<b>Examples.</b> If <b>ALIGNEE</b> is are set.	s scalar, then no elements of LB, UB, STRIDE, or AXIS_MAP $% \label{eq:mass_scalar}$
44 45	Given the declarations	
46	REAL PI = 3.1415927	
47		3(20,30),C(20,40,10),D(40)
48	!HPF\$ TEMPLATE T(40,20)	

!HPF\$ ALIGN A(I,:) WITH T(1+3\*I,2:20:2) !HPF\$ ALIGN C(I,\*,J) WITH T(J,21-I) !HPF\$ ALIGN D(I) WITH T(I,4) !HPF\$ PROCESSORS PROCS(4,2), SCALARPROC !HPF\$ DISTRIBUTE T(BLOCK,BLOCK) ONTO PROCS !HPF\$ DISTRIBUTE B(CYCLIC,BLOCK) ONTO PROCS !HPF\$ DISTRIBUTE ONTO SCALARPROC :: PI

assuming that the actual mappings are as the directives specify, the results of calling HPF\_ALIGNMENT are:

	А	В	С	D
LB	[4, 2]	[1, 1]	[20, N/A, 1]	[1]
UB	[31, 20]	[20,30]	[1, N/A, 10]	[40]
STRIDE	[3, 2]	[1, 1]	[-1, 0, 1]	[1]
AXIS_MAP	[1, 2]	[1, 2]	[2,  0,  1]	[1]
IDENTITY_MAP	false	true	false	false
NCOPIES	1	1	1	1

where "N/A" denotes a implementation-dependent result. To illustrate the use of NCOPIES, consider:

LOGICAL BOZO(20,20)	,RONALD_MCDONALD(20)	21		
!HPF\$ TEMPLATE EMMETT_KEL	!HPF\$ TEMPLATE EMMETT_KELLY(100,100)			
!HPF\$ ALIGN RONALD_MCDONA	!HPF\$ ALIGN RONALD_MCDONALD(I) WITH BOZO(I,*)			
!HPF\$ ALIGN BOZO(J,K) WIT		24		
		25		
CALL HPF_ALIGNMENT(RONALD_MC	DONALD, NCOPIES = NC) sets NC to 20. Now consider:	26		
LOGICAL BOZO(20,20)		27		
HPF\$ TEMPLATE WILLIE_WHI:	•	28		
HPF\$ ALIGN RONALD_MCDONA		29		
HPF\$ ALIGN BOZO(J,*) WIT		30		
INTE ALIGN DOLD(3,*) WIT	I WILLIE_WILDIEL(0+3)	31		
CALL HPF_ALIGNMENT(RONALD_MCDONALD, NCOPIES = NC) sets NC to one.				
HPF_DISTRIBUTION(DISTRIBUTEE, AXIS_TYPE,				
AXIS_INFO, PROCESSORS_RANK, PROCESSORS_SHAPE)				
<b>Optional Arguments.</b> AXIS_TYPE, AXIS_INFO, PROCESSORS_RANK,				
PROCESSORS_SHAPE		38		
-	STRIBUTION subroutine returns information regarding the	40		
distribution of the ultimat	te align-target associated with a variable.	41		
<b>Class.</b> Mapping inquiry s	ubrouting	42		
Class. Mapping inquiry s		43		
Arguments.		44		
		45		
DISTRIBUTEE	may be of any type. It may be scalar or array valued. It	46		
	must not be an assumed-size array. If it is a member of an	47		
	aggregate variable group, then it must be an aggregate	48		

1 2 3 4 5 6 7		<ul><li>cover of the group. (See Section 3.8 for the definitions of "aggregate variable group" and "aggregate cover.") It must not be a pointer that is disassociated or an allocatable array that is not allocated. It is an INTENT (IN) argument.</li><li>If DISTRIBUTEE is a pointer, information about the distribution of its target is returned. The target must not</li></ul>
8 9 10		be an assumed-size dummy argument or a section of an assumed-size dummy argument.
11 12 13 14 15 16 17 18 19 20 21	AXIS_TYPE (optional)	must be a rank one array of type default character. It may be of any length, although it must be of length at least 9 in order to contain the complete value. Its elements are set to the values below as if by a char- acter intrinsic assignment statement. Its size must be at least equal to the rank of the <i>align-target</i> to which DISTRIBUTEE is ultimately aligned; this is the value re- turned by HPF_TEMPLATE in TEMPLATE_RANK). It is an INTENT (OUT) argument. Its i <sup>th</sup> element contains infor- mation on the distribution of the i <sup>th</sup> axis of that <i>align- target</i> . The following values are defined by HPF (imple- mentations may define other values):
22 23 24 25 26 27 28 29 30 31		<ul> <li>'BLOCK' The axis is distributed BLOCK. The corresponding element of AXIS_INFO contains the block size.</li> <li>'COLLAPSED' The axis is collapsed (distributed with the "*" specification). The value of the corresponding element of AXIS_INFO is implementation dependent.</li> <li>'CYCLIC' The axis is distributed CYCLIC. The corresponding element of AXIS_INFO contains the block size.</li> </ul>
32 33 34 35 36 37 38 39 40	AXIS_INFO (optional)	must be a rank one array of type default integer, and size at least equal to the rank of the <i>align-target</i> to which DISTRIBUTEE is ultimately aligned (which is returned by HPF_TEMPLATE in TEMPLATE_RANK). It is an INTENT (OUT) argument. The i <sup>th</sup> element of AXIS_INFO contains the block size in the block or cyclic distribution of the i <sup>th</sup> axis of the ultimate <i>align-target</i> of DISTRIBUTEE; if that axis is a collapsed axis, then the value is implementation de- pendent.
41 42 43 44 45	PROCESSORS_RANK (option:	al) must be scalar and of type default integer. It is set to the rank of the processor arrangement onto which DISTRIBUTEE is distributed. It is an INTENT (OUT) ar- gument.
46 47 48	PROCESSORS_SHAPE (option	nal) must be a rank one array of type default integer and of size at least equal to the value, $m$ , returned in PROCES- SORS_RANK. It is an INTENT (OUT) argument. Its first $m$

elements are set to the shape of the processor arrangement onto which DISTRIBUTEE is mapped. (It may be necessary to call HPF\_DISTRIBUTION twice, the first time to obtain the value of PROCESSORS\_RANK in order to allocate PROCESSORS\_SHAPE.)

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**Example.** Given the declarations in the example illustrating HPF\_ALIGNMENT, and assuming that the actual mappings are as the directives specify, the results of HPF\_DISTRIBUTION are:

	А	В	ΡI
AXIS_TYPE	['BLOCK', 'BLOCK']	['CYCLIC', 'BLOCK']	[]
AXIS_INFO	[10, 10]	[1, 15]	[ ]
PROCESSORS_SHAPE	[4, 2]	[2, 2]	[ ]
PROCESSORS_RANK	2	2	0

# HPF\_TEMPLATE(ALIGNEE, TEMPLATE\_RANK, LB, UB, AXIS\_TYPE, AXIS\_INFO, NUMBER\_ALIGNED)

**Optional Arguments.** LB, UB, AXIS\_TYPE, AXIS\_INFO, NUMBER\_ALIGNED, TEMPLATE\_RANK

**Description.** The HPF\_TEMPLATE subroutine returns information regarding the ultimate *align-target* associated with a variable; HPF\_TEMPLATE returns information concerning the variable from the point of view of its ultimate *align-target*, while HPF\_ALIGNMENT returns information from the variable's point of view.

Class. Mapping inquiry subroutine.

ALIGNEE may be of any type. It may be scalar or array valued. It 32 must not be an assumed-size array. If it is a member of an 33 aggregate variable group, then it must be an aggregate 34 cover of the group. (See Section 3.8 for the definitions 35 of "aggregate variable group" and "aggregate cover.") It 36 must not be a pointer that is disassociated or an allocat-37 able array that is not allocated. It is an INTENT (IN) 38 argument. 39 40 If ALIGNEE is a pointer, information about the align-41ment of its target is returned. The target must not 42 be an assumed-size dummy argument or a section of an 43 assumed-size dummy argument. 44 TEMPLATE\_RANK (optional) must be scalar and of type default integer. It is an INTENT 45(OUT) argument. It is set to the rank of the ultimate 46 align-target. This can be different from the rank of the 47

ALIGNEE, due to collapsing and replicating.

1 2 3 4 5 6	LB (optional)	must be of type default integer and of rank one. Its size must be at least equal to the rank of the <i>align-target</i> to which ALIGNEE is ultimately aligned; this is the value returned in TEMPLATE_RANK. It is an INTENT (OUT) argu- ment. The i <sup>th</sup> element of LB contains the declared <i>align-</i> <i>target</i> lower bound for the i <sup>th</sup> template axis.
7 8 9 10 11 12 13	UB (optional)	must be of type default integer and of rank one. Its size must be at least equal to the rank of the <i>align-target</i> to which ALIGNEE is ultimately aligned; this is the value returned in TEMPLATE_RANK. It is an INTENT (OUT) argu- ment. The i <sup>th</sup> element of UB contains the declared <i>align-</i> <i>target</i> upper bound for the i <sup>th</sup> template axis.
14 15 16 17 18 19 20 21 22 23 24	AXIS_TYPE (optional)	must be a rank one array of type default character. It may be of any length, although it must be of length at least 10 in order to contain the complete value. Its elements are set to the values below as if by a char- acter intrinsic assignment statement. Its size must be at least equal to the rank of the <i>align-target</i> to which ALIGNEE is ultimately aligned; this is the value returned in TEMPLATE_RANK. It is an INTENT (OUT) argument. The i <sup>th</sup> element of AXIS_TYPE contains information about the i <sup>th</sup> axis of the <i>align-target</i> . The following values are de- fined by HPF (implementations may define other values):
25 26 27 28 29		'NORMAL' The <i>align-target</i> axis has an axis of ALIGNEE aligned to it. For elements of AXIS_TYPE assigned this value, the corresponding element of AXIS_INFO is set to the number of the axis of ALIGNEE aligned to this <i>align-target</i> axis.
30 31 32 33 34 35		'REPLICATED' ALIGNEE is replicated along this <i>align-tar-</i> <i>get</i> axis. For elements of AXIS_TYPE assigned this value, the corresponding element of AXIS_INFO is set to the number of copies of ALIGNEE along this <i>align-</i> <i>target</i> axis.
36 37 38 39 40		'SINGLE' ALIGNEE is aligned with one coordinate of the <i>align-target</i> axis. For elements of AXIS_TYPE assigned this value, the corresponding element of AXIS_INFO is set to the <i>align-target</i> coordinate to which ALIGNEE is aligned.
41 42 43 44 45 46	AXIS_INFO (optional)	must be of type default integer and of rank one. Its size must be at least equal to the rank of the <i>align-target</i> to which ALIGNEE is ultimately aligned; this is the value returned in TEMPLATE_RANK. It is an INTENT (OUT) argu- ment. See the description of AXIS_TYPE above.
46 47 48	NUMBER_ALIGNED (optional)	must be scalar and of type default integer. It is an INTENT (OUT) argument. It is set to the total number

of variables aligned to the ultimate *align-target*. This is the number of variables that are moved if the *align-target* is redistributed.

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**Example.** Given the declarations in the example illustrating HPF\_ALIGNMENT, and assuming that the actual mappings are as the directives specify, the results of HPF\_TEMPLATE are:

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		А	С	D	9
	LB	[1, 1]	[1, 1]	[1, 1]	10
	UB	[40, 20]	[40, 20]	[40,  20]	11
	AXIS_TYPE	['NORMAL',	['NORMAL',	['NORMAL',	12
		'NORMAL']	'NORMAL']	'SINGLE']	13
	AXIS_INFO	[1, 2]	[3, 1]	[1, 4]	14
	NUMBER_ALIGNED	3	3	3	15
	TEMPLATE_RANK	2	2	2	16
					17
IALL(ARRA	AY, DIM, MAS	SK)			18
Ϋ́,		,			19
Option	al Arguments. D	IM, MASK			20
					21
-	otion. Computes	a bitwise logic	al AND reduc	ction along dimension <b>DIM</b> of	22
ARRAY.					23
					24
Class.	Transformational f	unction.			25
•					26
Argum	ents.				27
ARRAY		must be of typ	pe integer. It	must not be scalar.	28 29
DTM (	• 1)	. 1 1	1 6 4	• . • .1 1 • .1	30
$DIM\ (\mathrm{opt}$	tional)			e integer with a value in the	31
				<i>n</i> is the rank of <b>ARRAY</b> . The	32
			_	nent must not be an optional	33
		dummy argun	nent.		34
MASK (oj	ptional)	must be of ty	vpe logical an	d must be conformable with	35
		ARRAY.			36
					37
$\mathbf{Result}$	Type, Type Par	ameter, and	Shape. The	result is of type integer with	38
the sam	e kind type param	eter as ARRAY.	It is scalar if	DIM is absent or if ARRAY has	39
rank one	e; otherwise, the re	esult is an arra	y of rank $n$ –	1 and shape	40
$(d_1, d_2, .$	$\ldots, d_{DIM-1}, d_{DIM}$	$+1,\ldots,d_n$ ) whe	re $(d_1, d_2,, d_n)$	$d_n$ ) is the shape of ARRAY.	41
					42
$\mathbf{Result}$	Value.				43
~ …					44
Case~(i)				luction of all the elements of	45
	ARRAY. If ARRA	Y has size zerc	), the result is	s equal to a implementation-	46

Case (i): The result of IALL(ARRAY) is the IAND reduction of all the elements of 45
 ARRAY. If ARRAY has size zero, the result is equal to a implementation-46
 dependent integer value x with the property that IAND(I, x) = I for all 47
 integers I of the same kind type parameter as ARRAY. See Section 7.4.3. 48

1 2 3 4	ements of ARRA tains no true el integer value a	ALL (ARRAY, MASK=MASK) is the IAND reduction of all the el- AY corresponding to the true elements of MASK; if MASK con- lements, the result is equal to a implementation-dependent r (of the same kind type parameter as ARRAY) with the HAND(I = R) = I for all integers I
5 6 7 8 9 10	Case (iii): If ARRAY has r that of IALL(A $(s_1, s_2, \dots, s_{DI})$ equal to IALL(	IAND(I, x) = I for all integers I. ank one, IALL(ARRAY, DIM [,MASK]) has a value equal to RRAY [,MASK]). Otherwise, the value of element $M_{-1}, s_{DIM+1}, \ldots, s_n$ of IALL(ARRAY, DIM [,MASK]) is ARRAY( $s_1, s_2, \ldots, s_{DIM-1}, :, s_{DIM+1}, \ldots, s_n$ ) K( $s_1, s_2, \ldots, s_{DIM-1}, :, s_{DIM+1}, \ldots, s_n$ )])
11 12	Examples.	
13	Case (i): The value of I	ALL((/7, 6, 3, 2/)) is 2.
1415 16	Case (ii): The value of I odd elements of	ALL(C, MASK = BTEST(C,O)) is the IAND reduction of the of C.
17 18		$y \begin{bmatrix} 2 & 3 & 5 \\ 3 & 7 & 7 \end{bmatrix}, \text{ then IALL(B, DIM = 1) is } \begin{bmatrix} 2 & 3 & 5 \end{bmatrix}$
19 20	and IALL(B, I	DIM = 2) is $\begin{bmatrix} 0 & 3 \end{bmatrix}$ .
21 22 IA	ALL_PREFIX(ARRAY, I	DIM, MASK, SEGMENT, EXCLUSIVE)
23 24	Optional Arguments. I	DIM, MASK, SEGMENT, EXCLUSIVE
25 26	<b>Description.</b> Computes a of ARRAY.	a segmented bitwise logical AND scan along dimension $\mathtt{DIM}$
27 28	<b>Class.</b> Transformational	function.
29 30	Arguments.	
31	ARRAY	must be of type integer. It must not be scalar.
32 33	DIM (optional)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of ARRAY.
34 35 36	MASK (optional)	must be of type logical and must be conformable with ARRAY.
37 38	SEGMENT (optional)	must be of type logical and must have the same shape as ARRAY.
39	EXCLUSIVE (optional)	must be of type logical and must be scalar.
40 41	Result Type, Type Par	rameter, and Shape. Same as ARRAY.
42 43 44 45		of the result has the value IALL((/ $a_1, \ldots, a_m$ /)) where ly empty) set of elements of ARRAY selected to contribute Section 7.4.5.
46 47 48	Example. IALL_PREFIX( $\begin{bmatrix} 1 & 1 & 0 & 4 & 4 \end{bmatrix}$ .	(/1,3,2,4,5/), SEGMENT= (/F,F,F,T,T/) ) is

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# IALL\_SCATTER(ARRAY, BASE, INDX1, ..., INDXn, MASK)

### **Optional Argument. MASK**

Description. Scatters elements of ARRAY selected by MASK to positions of the result indicated by index arrays INDX1, ..., INDXn. The j<sup>th</sup> bit of an element of the result is 1 if and only if the j<sup>th</sup> bits of the corresponding element of **BASE** and of the elements of ARRAY scattered to that position are all equal to 1.

Class. Transformational function.

#### Arguments.

ARRAY	must be of type integer. It must not be scalar.	13
BASE	must be of type integer with the same kind type param- eter as <b>ARRAY</b> . It must not be scalar.	14 15 16
INDX1,,INDXn	must be of type integer and conformable with ARRAY. The number of INDX arguments must be equal to the rank of BASE.	17 18 19
MASK (optional)	must be of type logical and must be conformable with ARRAY.	20 21 22

## **Result Type, Type Parameter, and Shape.** Same as BASE.

**Result Value.** The element of the result corresponding to the element b of BASE has the value IALL(  $(/a_1, a_2, ..., a_m, b/)$ ), where  $(a_1, ..., a_m)$  are the elements of ARRAY associated with b as described in Section 7.4.4.

Example. IALL\_SCATTER((/1, 2, 3, 6/), (/1, 3, 7/), (/1, 1, 2, 2/)) is 027.

# IALL\_SUFFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)

#### **Optional Arguments.** DIM, MASK, SEGMENT, EXCLUSIVE

Description. Computes a reverse, segmented bitwise logical AND scan along dimension DIM of ARRAY.

Class. Transformational function.

# Arguments.

ARRAY	must be of type integer. It must not be scalar.	43
DIM (optional)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of ARRAY.	44 45 46
MASK (optional)	must be of type logical and must be conformable with ARRAY.	47 48

SEGMENT (optional) must be of type logical and must have the same shape as 1 ARRAY. 2 3 **EXCLUSIVE** (optional) must be of type logical and must be scalar. 4 **Result Type, Type Parameter, and Shape.** Same as ARRAY. 5 6 **Result Value.** Element r of the result has the value IALL( $(/ a_1, \ldots, a_m /)$ ) where 7  $(a_1,\ldots,a_m)$  is the (possibly empty) set of elements of ARRAY selected to contribute 8 to r by the rules stated in Section 7.4.5. 9 10 Example. IALL\_SUFFIX( (/1,3,2,4,5/), SEGMENT= (/F,F,F,T,T/) ) is 11  $0 \ 2 \ 2 \ 4 \ 5$  . 1213 IANY(ARRAY, DIM, MASK) 14 15**Optional Arguments.** DIM, MASK 16**Description.** Computes a bitwise logical OR reduction along dimension DIM of 17ARRAY. 18 19 Class. Transformational function. 20 21 Arguments. 22 ARRAY must be of type integer. It must not be scalar. 23 **DIM** (optional) must be scalar and of type integer with a value in the  $^{24}$ range 1 < DIM < n, where n is the rank of ARRAY. The 25corresponding actual argument must not be an optional 26 dummy argument. 27 28 MASK (optional) must be of type logical and must be conformable with 29 ARRAY. 30 **Result Type, Type Parameter, and Shape.** The result is of type integer with 31 the same kind type parameter as ARRAY. It is scalar if DIM is absent or if ARRAY has 32 rank one; otherwise, the result is an array of rank n-1 and shape 33  $(d_1, d_2, \ldots, d_{DIM-1}, d_{DIM+1}, \ldots, d_n)$  where  $(d_1, d_2, \ldots, d_n)$  is the shape of ARRAY. 34 35 Result Value. 36 37 *Case (i):* The result of IANY(ARRAY) is the IOR reduction of all the elements of 38 ARRAY. If ARRAY has size zero, the result has the value zero. See Sec-39 tion 7.4.3. 40 Case (ii): The result of IANY(ARRAY, MASK=MASK) is the IOR reduction of all the 41elements of ARRAY corresponding to the true elements of MASK; if MASK 42 contains no true elements, the result is zero. 43 Case (iii): If ARRAY has rank one, IANY(ARRAY, DIM [,MASK]) has a value equal to 44 that of IANY(ARRAY [,MASK]). Otherwise, the value of element 45  $(s_1, s_2, \ldots, s_{DIM-1}, s_{DIM+1}, \ldots, s_n)$  of IANY(ARRAY, DIM [,MASK]) is 46 equal to IANY(ARRAY( $s_1, s_2, \ldots, s_{DIM-1}, \ldots, s_{DIM+1}, \ldots, s_n$ ) 47  $[, MASK = MASK(s_1, s_2, \dots, s_{DIM-1}, :, s_{DIM+1}, \dots, s_n)])$ 48

	Examples.		1
	<i>Case (i):</i> The value of I	ANY((/9, 8, 3, 2/)) is 11.	2 3
		ANY(C, MASK = BTEST(C,O)) is the IOR reduction of the	4
	odd elements o		5
			6
	Case (iii): If B is the arra	$\operatorname{ay}\left[\begin{array}{cc}2&3&5\\0&4&2\end{array}\right],  \text{then IANY(B, DIM = 1) is } \left[\begin{array}{cc}2&7&7\end{array}\right]$	7
		$DIM = 2) \text{ is } \begin{bmatrix} 7 & 6 \end{bmatrix}.$	8 9
	.,		10
IAN	Y_PREFIX(ARRAY,	DIM, MASK, SEGMENT, EXCLUSIVE)	11
	X · · ·		12
	Optional Arguments.	DIM, MASK, SEGMENT, EXCLUSIVE	13
	<b>Description</b> . Computes	a segmented bitwise logical OR scan along dimension DIM	14 15
	of ARRAY.		16
	Class. Transformational	function	17
	Class. Italisiofiliational		18
	Arguments.		19
			20 21
	ARRAY	must be of type integer. It must not be scalar.	22
	DIM (optional)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of ARRAY.	23 24
	MASK (optional)	must be of type logical and must be conformable with ARRAY.	25 26
	SEGMENT (optional)	must be of type logical and must have the same shape as	27
		ARRAY.	28
	EXCLUSIVE (optional)	must be of type logical and must be scalar.	29 30
		must be of type logical and must be scalar.	31
	Result Type, Type Pa	rameter, and Shape. Same as ARRAY.	32
			33
		of the result has the value IANY((/ $a_1, \ldots, a_m$ /)) where by empty) set of elements of ARRAY selected to contribute	34
	$(a_1, \ldots, a_m)$ is the (possion) to r by the rules stated in		35 36
			37
		(/1,2,3,2,5/), SEGMENT= (/F,F,F,T,T/) ) is	38
	$\begin{bmatrix} 1 & 3 & 3 & 2 & 7 \end{bmatrix}$ .		39
			40
IAN	Y_SCATTER(ARRAY	$(\mathbf{BASE}, \mathbf{INDX1},, \mathbf{INDXn}, \mathbf{MASK})$	41 42
	Optional Argument. M	ASK	43
	<b></b>		44
		ements of ARRAY selected by MASK to positions of the result	45
	is 1 if and only if the i <sup>th</sup>	INDX1,, INDXn. The $j^{th}$ bit of an element of the result bit of the corresponding element of BASE or of any of the	46
		ed to that position is equal to 1.	47 48
		1 1	-0

1	Class. Transformationa	l function.
2 3	Arguments.	
4	ARRAY	must be of type integer. It must not be scalar.
5 6 7	BASE	must be of type integer with the same kind type param- eter as <b>ARRAY</b> . It must not be scalar.
8 9 10	INDX1,,INDXn	must be of type integer and conformable with ARRAY. The number of INDX arguments must be equal to the rank of BASE.
11 12	MASK (optional)	must be of type logical and must be conformable with ARRAY.
13 14	Result Type, Type P	arameter, and Shape. Same as BASE.
15 16 17 18	has the value IANY( (/	ement of the result corresponding to the element $b$ of BASE $a_1, a_2,, a_m, b/$ ) ), where $(a_1,, a_m)$ are the elements of as described in Section 7.4.4.
19 20 21	Example. IANY_SCATT $\begin{bmatrix} 3 & 7 & 2 \end{bmatrix}$ .	TER((/1, 2, 3, 6/), (/1, 3, 7/), (/1, 1, 2, 2/)) is
	ANY_SUFFIX(ARRAY.	DIM, MASK, SEGMENT, EXCLUSIVE)
23 24	· · · ·	. DIM, MASK, SEGMENT, EXCLUSIVE
25 26 27		s a reverse, segmented bitwise logical OR scan along dimen-
28	Class. Transformationa	l function.
29 30	Arguments.	
31	ARRAY	must be of type integer. It must not be scalar.
32 33 34	DIM (optional)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of ARRAY.
35 36	MASK (optional)	must be of type logical and must be conformable with ARRAY.
37 38	SEGMENT (optional)	must be of type logical and must have the same shape as ARRAY.
39 40	EXCLUSIVE (optional)	must be of type logical and must be scalar.
41	Result Type, Type P	arameter, and Shape. Same as ARRAY.
42 43 44 45		a r of the result has the value IANY((/ $a_1, \ldots, a_m$ /)) where bibly empty) set of elements of ARRAY selected to contribute in Section 7.4.5.
46 47 48	Example. IANY_SUFFIX $\begin{bmatrix} 7 & 3 & 3 & 7 & 5 \end{bmatrix}$ .	K( (/4,2,3,2,5/), SEGMENT= (/F,F,F,T,T/) ) is

	Arguments.	DIM, MASK
Descripti DIM of ARR	-	a bitwise logical exclusive OR reduction along dimension
Class. Tra	ansformational	function.
Argumen	ts.	
ARRAY		must be of type integer. It must not be scalar.
DIM (optio	nal)	must be scalar and of type integer with a value in th range $1 \leq \text{DIM} \leq n$ , where n is the rank of ARRAY. Th corresponding actual argument must not be an optional dummy argument.
MASK (opti	onal)	must be of type logical and must be conformable wit ARRAY.
the same k rank one; d	kind type param otherwise, the r $, d_{DIM-1}, d_{DIM}$	<b>arameter, and Shape.</b> The result is of type integer with neter as ARRAY. It is scalar if DIM is absent or if ARRAY has result is an array of rank $n - 1$ and shape $d_{l+1}, \ldots, d_n$ where $(d_1, d_2, \ldots, d_n)$ is the shape of ARRAY.
Case (i):	The result of IPARITY(ARRAY) is the IEOR reduction of all the elements of ARRAY. If ARRAY has size zero, the result has the value zero. See Section 7.4.3.	
	The result of IPARITY(ARRAY, MASK=MASK) is the IEOR reduction of all the elements of ARRAY corresponding to the true elements of MASK; if MASK contains no true elements, the result is zero.	
Case (ii):	the elements o	f ARRAY corresponding to the true elements of $MASK$ ; if $MAS$
	the elements of contains no tr If ARRAY has no to that of IPA $(s_1, s_2, \ldots, s_D)$ equal to IPAR.	of ARRAY corresponding to the true elements of MASK; if MAS ue elements, the result is zero. rank one, IPARITY(ARRAY, DIM [,MASK]) has a value equa RITY(ARRAY [,MASK]). Otherwise, the value of element
	the elements of contains no tr If ARRAY has no to that of IPA $(s_1, s_2, \dots, s_D)$ equal to IPAR: [,MASK = MAS	of ARRAY corresponding to the true elements of MASK; if MAS ue elements, the result is zero. rank one, IPARITY(ARRAY, DIM [,MASK]) has a value equa RITY(ARRAY [,MASK]). Otherwise, the value of element $IM_{-1}, s_{DIM+1}, \ldots, s_n$ ) of IPARITY(ARRAY, DIM [,MASK]) is ITY(ARRAY( $s_1, s_2, \ldots, s_{DIM-1}, \vdots, s_{DIM+1}, \ldots, s_n$ )
Case (iii):	the elements of contains no tr If ARRAY has no to that of IPA $(s_1, s_2, \ldots, s_D)$ equal to IPAR: [,MASK = MAS 5.	of ARRAY corresponding to the true elements of MASK; if MAS ue elements, the result is zero. rank one, IPARITY(ARRAY, DIM [,MASK]) has a value equa RITY(ARRAY [,MASK]). Otherwise, the value of element $IM_{-1}, s_{DIM+1}, \ldots, s_n$ ) of IPARITY(ARRAY, DIM [,MASK]) is ITY(ARRAY( $s_1, s_2, \ldots, s_{DIM-1}, :, s_{DIM+1}, \ldots, s_n$ )
Case (iii): Examples Case (i): Case (ii):	the elements of contains no tr If ARRAY has no to that of IPA $(s_1, s_2, \dots, s_D)$ equal to IPAR [,MASK = MAS 5. The value of I the odd eleme	of ARRAY corresponding to the true elements of MASK; if MASK ue elements, the result is zero. rank one, IPARITY(ARRAY, DIM [,MASK]) has a value equal RITY(ARRAY [,MASK]). Otherwise, the value of element $IM_{-1}, s_{DIM+1}, \ldots, s_n$ ) of IPARITY(ARRAY, DIM [,MASK]) is ITY(ARRAY( $s_1, s_2, \ldots, s_{DIM-1}, :, s_{DIM+1}, \ldots, s_n$ ) $K(s_1, s_2, \ldots, s_{DIM-1}, :, s_{DIM+1}, \ldots, s_n)$ ]) IPARITY((/13, 8, 3, 2/)) is 4. IPARITY(C, MASK = BTEST(C,O)) is the IEOR reduction of onts of C.
Case (iii): Examples Case (i): Case (ii):	the elements of contains no tr If ARRAY has no to that of IPA $(s_1, s_2, \ldots, s_D)$ equal to IPAR [,MASK = MAS s. The value of I the value of I the odd element If B is the array	of ARRAY corresponding to the true elements of MASK; if MASK ue elements, the result is zero. rank one, IPARITY(ARRAY, DIM [,MASK]) has a value equal RITY(ARRAY [,MASK]). Otherwise, the value of element $IM_{-1}, s_{DIM+1}, \ldots, s_n$ ) of IPARITY(ARRAY, DIM [,MASK]) is ITY(ARRAY( $s_1, s_2, \ldots, s_{DIM-1}, \vdots, s_{DIM+1}, \ldots, s_n$ ) $K(s_1, s_2, \ldots, s_{DIM-1}, \vdots, s_{DIM+1}, \ldots, s_n)$ ]) IPARITY((/13, 8, 3, 2/)) is 4.

IPARITY\_PREFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE) 1 2 **Optional Arguments.** DIM, MASK, SEGMENT, EXCLUSIVE з 4 **Description**. Computes a segmented bitwise logical exclusive OR scan along di-5 mension DIM of ARRAY. 6 7 Class. Transformational function. 8 9 Arguments. 10 ARRAY must be of type integer. It must not be scalar. 11 12DIM (optional) must be scalar and of type integer with a value in the 13 range 1 < DIM < n, where n is the rank of ARRAY. 14 MASK (optional) must be of type logical and must be conformable with 15ARRAY. 16SEGMENT (optional) must be of type logical and must have the same shape as 1718 ARRAY. 19 **EXCLUSIVE** (optional) must be of type logical and must be scalar. 20 21 **Result Type, Type Parameter, and Shape.** Same as ARRAY. 22 23 **Result Value.** Element r of the result has the value IPARITY((/  $a_1, \ldots, a_m$  /)) 24 where  $(a_1, \ldots, a_m)$  is the (possibly empty) set of elements of ARRAY selected to con-25tribute to r by the rules stated in Section 7.4.5. 26 Example. IPARITY\_PREFIX( (/1,2,3,4,5/), SEGMENT= (/F,F,F,T,T/) ) is 27  $1 \ 3 \ 0 \ 4 \ 1$ . 28 29 30 IPARITY\_SCATTER(ARRAY, BASE, INDX1, ..., INDXn, MASK) 31 **Optional Argument.** MASK 32 33 Description. Scatters elements of ARRAY selected by MASK to positions of the re-34 sult indicated by index arrays INDX1, ..., INDXn. The j<sup>th</sup> bit of an element of the 35 result is 1 if and only if there are an odd number of ones among the j<sup>th</sup> bits of the 36 corresponding element of BASE and the elements of ARRAY scattered to that position. 37 38 Class. Transformational function. 39 40 Arguments. 4142 must be of type integer. It must not be scalar. ARRAY 43 BASE must be of type integer with the same kind type param-44 eter as ARRAY. It must not be scalar. 45 INDX1,..., INDXn must be of type integer and conformable with ARRAY. The 46 number of INDX arguments must be equal to the rank of 47 BASE. 48

	MASK (optional)	must be of type logical and must be conformable with ARRAY.	1 2
	Result Type, Type Par	rameter, and Shape. Same as BASE.	3 4
	<b>Result Value.</b> The element of the result corresponding to the element $b$ of BASE has the value IPARITY( $(/a_1, a_2,, a_m, b/)$ ), where $(a_1,, a_m)$ are the elements of ARRAY associated with $b$ as described in Section 7.4.4.		
	Example. IPARITY_SCAT $\begin{bmatrix} 2 & 6 & 7 \end{bmatrix}$ .	TER((/1,2,3,6/), (/1,3,7/), (/1,1,2,2/)) is	9 10 11
IPAI	RITY_SUFFIX(ARRA	Y, DIM, MASK, SEGMENT, EXCLUSIVE)	12 13
	X	DIM, MASK, SEGMENT, EXCLUSIVE	14 15
	<b>Description.</b> Computes along dimension DIM of AF	a reverse, segmented bitwise logical exclusive OR scan RRAY.	16 17 18
	Class. Transformational	function.	19
	Arguments.		20 21
	ARRAY	must be of type integer. It must not be scalar.	22 23
	DIM (optional)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of ARRAY.	24 25
	MASK (optional)	must be of type logical and must be conformable with ARRAY.	26 27 28
	$\texttt{SEGMENT} \ (\texttt{optional})$	must be of type logical and must have the same shape as ARRAY.	29 30
	EXCLUSIVE (optional)	must be of type logical and must be scalar.	31 32
	Result Type, Type Pa	rameter, and Shape. Same as ARRAY.	33
	Degult Value Element	r of the pecult has the value TDADITY(() $r$ $r$ ())	34 35
		r of the result has the value IPARITY((/ $a_1, \ldots, a_m$ /)) (possibly empty) set of elements of ARRAY selected to con-	36
	tribute to $r$ by the rules s		37
	Example. IPARITY_SUFF $\begin{bmatrix} 0 & 1 & 3 & 1 & 5 \end{bmatrix}$ .	IX( (/1,2,3,4,5/), SEGMENT= (/F,F,F,T,T/) ) is	38 39 40
TTA	- D7(I)		41 42
LEA	DZ(I)		43
	<b>Description</b> . Return the	number of leading zeros in an integer.	44 45
	Class. Elemental function	1.	46
	Argument. I must be of	type integer	47
	<b>1115 DE DE DE DE</b>	opponnoceor.	48

1	Result Type and T	ype Parameter. Same as I.
2 3 4 5 6 7 8	I. The model for the i F95:13.5.7 LEADZ(0) is	result is a count of the number of leading 0-bits in the integer interpretation of an integer as a sequence of bits is in Section SBIT_SIZE(I). For nonzero I, if the leftmost one bit of I occurs re the rightmost bit is bit 0) then LEADZ(I) is BIT_SIZE(I) -
9 10 11 12 13 14 15	MINVAL( (/ (J, J=0 J=BIT_SIZE(I)-1, 0, results from LEADZ(I)	has the value BIT_SIZE(3) - 2. For scalar I, LEADZ(I) .EQ. , BIT_SIZE(I) ) /), MASK=M ) where M =(/ (BTEST(I,J), -1), .TRUE. /). A given integer I may produce different , depending on the number of bits in the representation of the . That is because LEADZ counts bits from the most significant EN.
16 17	MAXVAL_PREFIX(AB	RAY, DIM, MASK, SEGMENT, EXCLUSIVE)
18 19 20	Ň	ts. DIM, MASK, SEGMENT, EXCLUSIVE
21 22	<b>Description.</b> Compu	tes a segmented MAXVAL scan along dimension DIM of ARRAY.
23 24	Class. Transformatio	nal function.
25 26	Arguments.	
27 28	ARRAY	must be of type integer or real. It must not be scalar.
29 30 31	DIM (optional)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of ARRAY.
32 33	MASK (optional)	must be of type logical and must be conformable with ARRAY.
34 35 36	<b>SEGMENT</b> (optional)	must be of type logical and must have the same shape as ARRAY.
37 38	EXCLUSIVE (optional)	must be of type logical and must be scalar.
39 40 41	Result Type, Type	Parameter, and Shape. Same as ARRAY.
42 43 44 45	where $(a_1,\ldots,a_m)$ is t	ent $r$ of the result has the value MAXVAL((/ $a_1, \ldots, a_m$ /)) the (possibly empty) set of elements of ARRAY selected to con- es stated in Section 7.4.5.
46 47 48	Example. MAXVAL_PR $\left[\begin{array}{cccc} 3 & 4 & 4 & 2 & 5 \end{array}\right]$ .	EFIX( (/3,4,-5,2,5/), SEGMENT= (/F,F,F,T,T/) ) is

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# MAXVAL\_SCATTER(ARRAY, BASE, INDX1, ..., INDXn, MASK)

#### **Optional Argument. MASK**

**Description.** Scatters elements of ARRAY selected by MASK to positions of the result indicated by index arrays INDX1, ..., INDXn. Each element of the result is assigned the maximum value of the corresponding element of BASE and the elements of ARRAY scattered to that position.

Class. Transformational function.

#### Arguments.

ARRAY	must be of type integer or real. It must not be scalar.	13
BASE	must be of the same type and kind type parameter as ARRAY. It must not be scalar.	14
DADE		15
		16
INDX1,,INDXn	must be of type integer and conformable with ARRAY. The	17
	number of INDX arguments must be equal to the rank of	18
	BASE.	19
	must be of type logical and must be conformable with	20
MASK (optional)		21
	ARRAY.	22

#### **Result Type, Type Parameter, and Shape.** Same as BASE.

**Result Value.** The element of the result corresponding to the element b of BASE has the value MAXVAL (  $(a_1, a_2, ..., a_m, b/)$  ), where  $(a_1, ..., a_m)$  are the elements of ARRAY associated with b as described in Section 7.4.4.

Example. MAXVAL\_SCATTER((/1, 2, 3, 1/), (/4, -5, 7/), (/1, 1, 2, 2/)) is **4 3 7**.

# MAXVAL\_SUFFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)

#### **Optional Arguments.** DIM, MASK, SEGMENT, EXCLUSIVE

Description. Computes a reverse, segmented MAXVAL scan along dimension DIM of ARRAY.

Class. Transformational function.

#### Arguments.

ARRAY	must be of type integer or real. It must not be scalar.	43
DIM (optional)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of ARRAY.	44 45 46
MASK (optional)	must be of type logical and must be conformable with ARRAY.	47 48

1 2		$\texttt{SEGMENT} \ (\texttt{optional})$	must be of type logical and must have the same shape as ARRAY.	
3 4		EXCLUSIVE (optional)	must be of type logical and must be scalar.	
5		Result Type, Type Par	cameter, and Shape. Same as ARRAY.	
6 7 8 9		<b>Result Value.</b> Element r of the result has the value MAXVAL((/ $a_1, \ldots, a_m$ /) where $(a_1, \ldots, a_m)$ is the (possibly empty) set of elements of ARRAY selected to contribute to r by the rules stated in Section 7.4.5.		
10 11 12 13		<b>Example.</b> MAXVAL_SUFFI: $\begin{bmatrix} 4 & 4 & -5 & 5 \end{bmatrix}$ .	X( (/3,4,-5,2,5/), SEGMENT= (/F,F,F,T,T/) ) is	
14	MIN	VAL_PREFIX(ARRA	Y, DIM, MASK, SEGMENT, EXCLUSIVE)	
15 16		Optional Arguments.	DIM, MASK, SEGMENT, EXCLUSIVE	
17		<b>Description.</b> Computes a segmented MINVAL scan along dimension DIM of ARRAY.		
18 19		Class. Transformational function.		
20 21		Arguments.		
22		ARRAY	must be of type integer or real. It must not be scalar.	
23 24 25		DIM (optional)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of ARRAY.	
26 27		MASK (optional)	must be of type logical and must be conformable with ARRAY.	
28 29 30		${\tt SEGMENT}~({\rm optional})$	must be of type logical and must have the same shape as ARRAY.	
31		EXCLUSIVE (optional)	must be of type logical and must be scalar.	
32 33		Result Type, Type Par	cameter, and Shape. Same as ARRAY.	
34 35 36 37		<b>Result Value.</b> Element r of the result has the value MINVAL((/ $a_1, \ldots, a_m$ /)) where $(a_1, \ldots, a_m)$ is the (possibly empty) set of elements of ARRAY selected to contribute to r by the rules stated in Section 7.4.5.		
38 39 40		Example. MINVAL_PREFINE $\begin{bmatrix} 1 & 1 & -3 & 4 \end{bmatrix}$ .	X( (/1,2,-3,4,5/), SEGMENT= (/F,F,F,T,T/) ) is	
41 42	MIN	VAL_SCATTER(ARR	AY,BASE,INDX1,, INDXn, MASK)	
43		Optional Argument. MASK		
44 45 46 47 48		indicated by index arrays	ments of ARRAY selected by MASK to positions of the result INDX1,, INDXn. Each element of the result is assigned corresponding element of BASE and the elements of ARRAY	

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Class. Transformational function. Arguments. must be of type integer or real. It must not be scalar. ARRAY BASE must be of the same type and kind type parameter as ARRAY. It must not be scalar. must be of type integer and conformable with **ARRAY**. The INDX1,...,INDXn number of INDX arguments must be equal to the rank of BASE. MASK (optional) must be of type logical and must be conformable with ARRAY. Result Type, Type Parameter, and Shape. Same as BASE. **Result Value.** The element of the result corresponding to the element b of BASE has the value MINVAL(  $(/a_1, a_2, ..., a_m, b/)$  ), where  $(a_1, ..., a_m)$  are the elements of ARRAY associated with b as described in Section 7.4.4. Example. MINVAL\_SCATTER((/ 1,-2,-3,6 /), (/ 4,3,7 /), (/ 1,1,2,2 /)) is -2 -3 7 . MINVAL\_SUFFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE) **Optional Arguments.** DIM, MASK, SEGMENT, EXCLUSIVE Description. Computes a reverse, segmented MINVAL scan along dimension DIM of ARRAY. Class. Transformational function. Arguments. ARRAY must be of type integer or real. It must not be scalar. **DIM** (optional) must be scalar and of type integer with a value in the range 1 < DIM < n, where n is the rank of ARRAY. must be of type logical and must be conformable with MASK (optional) ARRAY.

ARRAY.

must be of type logical and must have the same shape as

**EXCLUSIVE** (optional) must be of type logical and must be scalar.

Result Type, Type Parameter, and Shape. Same as ARRAY.

SEGMENT (optional)

**Result Value.** Element r of the result has the value MINVAL((/ $a_1, \ldots, a_m$ /)) where  $(a_1, \ldots, a_m)$  is the (possibly empty) set of elements of ARRAY selected to contribute to r by the rules stated in Section 7.4.5.

Example. MINVAL\_SUFFIX( (/1,2,-3,4,5/), SEGMENT= (/F,F,F,T,T/) ) is [-3 -3 -3 4 5].

Optional	Argument. DIM
<b>Descript</b> i dimension	on. Determine whether an odd number of values are true in MASK along DIM.
Class. Tr	ansformational function.
Argumer	nts.
MASK	must be of type logical. It must not be scalar.
DIM (optio	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of MASK. The corresponding actual argument must not be an optional dummy argument.
the same rank one;	<b>ype, Type Parameter, and Shape.</b> The result is of type logical with kind type parameter as MASK. It is scalar if DIM is absent or if MASK has otherwise, the result is an array of rank $n - 1$ and shape $(d_{DIM-1}, d_{DIM+1}, \ldots, d_n)$ where $(d_1, d_2, \ldots, d_n)$ is the shape of MASK.
${f Result}$ V	alue.
<i>Case</i> ( <i>i</i> ):	The result of PARITY(MASK) is the .NEQV. reduction of all the elements of MASK. If MASK has size zero, the result has the value false. See Section 7.4.3.
Case (ii):	If MASK has rank one, PARITY(MASK, DIM) has a value equal to that of PARITY(MASK). Otherwise, the value of element $(s_1, s_2, \ldots, s_{DIM-1}, s_{DIM+1}, \ldots, s_n)$ of PARITY(MASK, DIM) is equal to PARITY(MASK $(s_1, s_2, \ldots, s_{DIM-1}, \vdots, s_{DIM+1}, \ldots, s_n)$ )
Example	s.
Case (i):	The value of PARITY((/T, T, T, F/)) is true.
Case (ii):	If B is the array $\begin{bmatrix} T & T & F \\ T & T & T \end{bmatrix}$ , then PARITY(B, DIM = 1) is $\begin{bmatrix} F & F & T \end{bmatrix}$
	and PARITY(B, DIM = 2) is $\begin{bmatrix} F & T \end{bmatrix}$ .
PARITY_PRI	EFIX(MASK, DIM, SEGMENT, EXCLUSIVE)
Optional	Arguments. DIM, SEGMENT, EXCLUSIVE
Descripti DIM of MAS	on. Computes a segmented logical exclusive OR scan along dimension SK.
Class. Tr	ansformational function.
Argumen	nts.
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MASK	must be of type logical. It must not be scalar.	1
DIM (optional)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of MASK.	2 3 4
SEGMENT (optional)	must be of type logical and must have the same shape as MASK.	5
EXCLUSIVE (optional)	must be of type logical and must be scalar.	7 8
		9

**Result Type, Type Parameter, and Shape.** Same as MASK.

**Result Value.** Element r of the result has the value PARITY((/  $a_1, \ldots, a_m$  /)) where  $(a_1, \ldots, a_m)$  is the (possibly empty) set of elements of MASK selected to contribute to r by the rules stated in Section 7.4.5.

Example. PARITY\_PREFIX( (/T,F,T,T,T/), SEGMENT= (/F,F,F,T,T/) ) is  $\begin{bmatrix} T & T & F & T & F \end{bmatrix}$ .

# PARITY\_SCATTER(MASK, BASE, INDX1, ..., INDXn)

**Description.** Scatters elements of MASK to positions of the result indicated by index arrays INDX1, ..., INDXn. An element of the result is true if and only if the number of true values among the corresponding element of BASE and the elements of MASK scattered to that position is odd.

Class. Transformational function.

Arguments.

MASK	must be of type logical. It must not be scalar.	31
IIADIA		32
BASE	must be of type logical with the same kind type parameter	33
	as MASK. It must not be scalar.	34
		35
INDX1,,INDXn	must be of type integer and conformable with MASK. The	36
	number of INDX arguments must be equal to the rank of BASE.	37
		38
		39

#### **Result Type, Type Parameter, and Shape.** Same as BASE.

**Result Value.** The element of the result corresponding to the element b of BASE has the value PARITY (  $(a_1, a_2, ..., a_m, b/)$  ), where  $(a_1, ..., a_m)$  are the elements of MASK associated with b as described in Section 7.4.4.

Example. PARITY\_SCATTER((/ T,T,T,T /), (/ T,F,F /), (/ 1,1,1,2 /)) is [FTF].

PARITY\_SUFFIX(MASK, DIM, SEGMENT, EXCLUSIVE) 1 2 **Optional Arguments.** DIM, SEGMENT, EXCLUSIVE 3 4 **Description**. Computes a reverse, segmented logical exclusive OR scan along di-5 mension DIM of MASK. 6 7 8 Class. Transformational function. 9 10 Arguments.  $1\,1$ 12MASK must be of type logical. It must not be scalar. 13 **DIM** (optional) 14 must be scalar and of type integer with a value in the range 1 < DIM < n, where n is the rank of MASK. 1516SEGMENT (optional) must be of type logical and must have the same shape as 17MASK. 18 19 **EXCLUSIVE** (optional) must be of type logical and must be scalar. 20 21 Result Type, Type Parameter, and Shape. Same as MASK. 22 23 **Result Value.** Element r of the result has the value PARITY( $(/a_1, \ldots, a_m/)$ ) 24 where  $(a_1, \ldots, a_m)$  is the (possibly empty) set of elements of MASK selected to con- $^{25}$ tribute to r by the rules stated in Section 7.4.5. 26 27 28 Example. PARITY\_SUFFIX( (/T,F,T,T,T/), SEGMENT= (/F,F,F,T,T/) ) is 29 FTTFT. 30 31 32 POPCNT(I)33 **Description**. Return the number of one bits in an integer. 34 35 Class. Elemental function. 36 37 38 Argument. I must be of type integer. 39 40 **Result Type and Type Parameter.** Same as I. 4142 **Result Value.** POPCNT(I) is the number of one bits in the binary representation of 43 the integer I. The model for the interpretation of an integer as a sequence of bits is 44 in Section F95:13.5.7 45 46 Example. POPCNT(I) = COUNT((/ (BTEST(I,J), J=0, BIT\_SIZE(I)-1) /)), for 47scalar I. 48

POP	PAR(I)		1
	Description. Return th	e parity of an integer.	2 3
	Class. Elemental function	on.	4 5
	Argument. I must be of type integer.		
	Result Type and Type Parameter. Same as I.		
	<b>Result Value.</b> POPPAR(I) is 1 if there are an odd number of one bits in I a if there are an even number. The model for the interpretation of an integ sequence of bits is in Section F95:13.5.7		
	Example. For scalar I,	<pre>POPPAR(I) = MERGE(1,0,BTEST(POPCNT(I),0)).</pre>	14 15 16
PRC	DUCT_PREFIX(AR	RAY, DIM, MASK, SEGMENT, EXCLUSIVE)	17 18
	Optional Arguments.	DIM, MASK, SEGMENT, EXCLUSIVE	19 20
	<b>Description</b> . Computes	s a segmented PRODUCT scan along dimension DIM of ARRAY.	20 21 22
	Class. Transformational	Class. Transformational function.	
	Arguments.	Arguments.	
	ARRAY	must be of type integer, real, or complex. It must not be scalar.	27 28 29
	DIM (optional)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of ARRAY.	30 31
	MASK (optional)	must be of type logical and must be conformable with ARRAY.	32 33 34
	<b>SEGMENT</b> (optional)	must be of type logical and must have the same shape as ARRAY.	35 36 37
	EXCLUSIVE (optional)	must be of type logical and must be scalar.	38
	Result Type, Type Pa	arameter, and Shape. Same as ARRAY.	39 40 41
	<b>Result Value.</b> Element r of the result has the value PRODUCT((/ $a_1, \ldots, a_m$ ) where $(a_1, \ldots, a_m)$ is the (possibly empty) set of elements of ARRAY selected to tribute to r by the rules stated in Section 7.4.5.		
	<b>Example.</b> PRODUCT_PRE $\begin{bmatrix} 1 & 2 & 6 & 4 & 20 \end{bmatrix}$ .	FIX( (/1,2,3,4,5/), SEGMENT= (/F,F,F,T,T/) ) is	45 46 47 48

1	PRODUCT_SCATTER(ARRAY, BASE, INDX1,, INDXn, MASK)						
2 3	Optional Argument. MASK	Optional Argument. MASK					
4 5 6 7 8	<b>Description.</b> Scatters elements of ARRAY selected by MASK to positions of the result indicated by index arrays INDX1,, INDXn. Each element of the result is equal to the product of the corresponding element of BASE and the elements of ARRAY scattered to that position.						
9	Class. Transformational function.						
10 11	Arguments.						
12 13	ARRAY must be of type integer, real scalar.	, or complex. It must not be					
14 15 16	BASE must be of the same type a ARRAY. It must not be scalar						
17 18 19	INDX1,,INDXn must be of type integer and c number of INDX arguments r BASE.						
20 21 22	MASK (optional) must be of type logical and ARRAY.	must be conformable with					
23 24	Result Type, Type Parameter, and Shape. Same	Result Type, Type Parameter, and Shape. Same as BASE.					
25 26 27 28	has the value PRODUCT( (/ $a_1, a_2,, a_m, b$ /) ), where (	<b>Result Value.</b> The element of the result corresponding to the element $b$ of BASE has the value PRODUCT ( $(a_1, a_2,, a_m, b/)$ ), where $(a_1,, a_m)$ are the elements of ARRAY associated with $b$ as described in Section 7.4.4.					
29 30 31	Example. PRODUCT_SCATTER((/ 1,2,3,1 /), (/ 4,- is [ 8 -15 7 ].	5,7 /), (/ 1,1,2,2 /))					
32 33	PRODUCT_SUFFIX(ARRAY, DIM, MASK, SEG	MENT, EXCLUSIVE)					
34	<b>Optional Arguments.</b> DIM, MASK, SEGMENT, EXCLUSIV	Έ					
35 36 37	<b>Description.</b> Computes a reverse, segmented <b>PRODUCT</b> ARRAY.	scan along dimension DIM of					
38 39	Class. Transformational function.						
40 41	Arguments.						
42 43	ARRAY must be of type integer, real scalar.	, or complex. It must not be					
44 45 46	$\texttt{DIM}$ (optional) must be scalar and of type range $1 \leq \texttt{DIM} \leq n$ , where $n$	0					
47 48	MASK (optional) must be of type logical and ARRAY.	must be conformable with					

SI	EGMENT (o	optional)	must be of type logical and must have the same shape as ARRAY.	1 2
E	XCLUSIVE	$\Sigma$ (optional)	must be of type logical and must be scalar.	3 4
R	lesult T	уре, Туре Раг	cameter, and Shape. Same as ARRAY.	5 6
W	<b>Result Value.</b> Element r of the result has the value PRODUCT((/ $a_1, \ldots, a_m$ /)) where $(a_1, \ldots, a_m)$ is the (possibly empty) set of elements of ARRAY selected to contribute to r by the rules stated in Section 7.4.5.			7 8 9 10
Г	xample. 6 6 3	7	IX( (/1,2,3,4,5/), SEGMENT= (/F,F,F,T,T/) ) is	11 12 13
SORT	DOWN	(ARRAY,DI	M)	14 15
			,	16
0	ptional	Argument. DI	M	17
п	escrinti	<b>on.</b> Sort by des	conding value	18
	CSCI PU	on. Bort by des	cending value.	19
С	lass. Tra	ansformational f	unction.	20 21
А	rgumen	.ts.		22 23
Al	ARRAY		must be of type integer, real, or character. It must not be scalar.	24 25
D	IM (optio	nal)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where <i>n</i> is the rank of <b>ARRAY</b> . The corresponding actual argument must not be an optional dummy argument.	26 27 28 29 30
		<b>ype, Type Pa</b> type parameter	rameter, and Shape. The result has the same shape, as ARRAY.	31 32 33
R	lesult Va	alue.		34 35
C	vector of the sa the same numb The collating s		SORT_DOWN(ARRAY), when ARRAY is one-dimensional, is a ame shape as ARRAY, containing the same elements (with per of instances) but sorted in descending element order. Sequence for an array of type CHARACTER is that used by rinsic functions, namely ASCII.	36 37 38 39 40 41
C	Case (ii):		SORT_DOWN(ARRAY) for a multi-dimensional ARRAY is the uld be obtained by reshaping ARRAY to a rank-one array	42 43

Case (1): The result of SURT\_DOWN(ARRAY) for a multi-dimensional ARRAY is the result that would be obtained by reshaping ARRAY to a rank-one array 43
 V using array element order, sorting that rank-one array in descending 44
 order, as in Case(i), and finally restoring the result to the original shape. 45
 That is, it gives the same result as RESHAPE( SORT\_DOWN(V), SHAPE = 46
 SHAPE(ARRAY) ), where V = RESHAPE( ARRAY, SHAPE = (/ M /) and M 47
 = SIZE(ARRAY). 48

Case (iii): The result of SORT\_DOWN(ARRAY, DIM=k) contains the same elements as A, 1 but each one-dimensional array section of the form ARRAY  $(i_1, i_2, \ldots, i_{k-1})$ : 2  $(i_{k+1},\ldots,i_n)$ , where n is the rank of ARRAY, has been sorted in descending 3 element order, as in Case(i) above. 4 5 Examples. 6 7 SORT\_DOWN( (/30, 20, 30, 40, -10/) ) Case (i): 8 has the value  $\begin{bmatrix} 40 & 30 & 30 & 20 & -10 \end{bmatrix}$ . 9 Case (ii): If **A** is the array  $\begin{bmatrix} 1 & 9 & 2 \\ 4 & 5 & 2 \\ 1 & 2 & 4 \end{bmatrix}$ , 10 1112then SORT\_DOWN(A) has the value  $\begin{bmatrix} 9 & 4 & 2 \\ 5 & 2 & 1 \\ 4 & 2 & 1 \end{bmatrix}$ 13 14 1516Case (iii): If **A** is the array  $\begin{bmatrix} 1 & 9 & 2 \\ 4 & 5 & 2 \\ 1 & 2 & 4 \end{bmatrix}$ , 1718 19  $\left[\begin{array}{rrrr} 4 & 9 & 4 \\ 1 & 5 & 2 \\ 1 & 2 & 2 \end{array}\right].$ 20 then SORT\_DOWN(A, DIM = 1) has the value 21 22 23 24 SORT\_UP(ARRAY,DIM) 25**Optional Argument.** DIM 26 27 **Description.** Sort by ascending value. 28 29 Class. Transformational function. 30 Arguments. 31 32 ARRAY must be of type integer, real, or character. It must not 33 be scalar. 34 DIM (optional) must be scalar and of type integer with a value in the 35 range 1 < DIM < n, where n is the rank of ARRAY. The 36 corresponding actual argument must not be an optional 37 dummy argument. 38 39 **Result Type, Type Parameter, and Shape.** The result has the same shape, 40 type, and type parameter as ARRAY. 4142 Result Value. 43 Case (i): The result of SORT\_UP(ARRAY), when ARRAY is one-dimensional, is a vector 44 of the same shape as ARRAY, containing the same elements (with the same 45 number of instances) but sorted in ascending element order. The collating 46 sequence for an array of type CHARACTER is that used by the Fortran 47

intrinsic functions, namely ASCII.

 $\left|\begin{array}{cccc} 1 & 5 & 2 \\ 4 & 9 & 4 \end{array}\right|.$ 

- Case (ii): The result of SORT\_UP(ARRAY) for a multi-dimensional ARRAY is the result that would be obtained by reshaping ARRAY to a rank-one array V using array element order, sorting that rank-one array in ascending order, as in Case(i), and finally restoring the result to the original shape. That is, it gives the same result as RESHAPE( SORT\_UP(V), SHAPE = SHAPE(ARRAY) ), where V = RESHAPE( ARRAY, SHAPE = (/ M /) and M = SIZE(ARRAY).
- Case (iii): The result of SORT\_UP(ARRAY, DIM=k) contains the same elements as A, but each one-dimensional array section of the form ARRAY( $i_1, i_2, \ldots, i_{k-1}$ ,: , $i_{k+1}, \ldots, i_n$ ), where n is the rank of ARRAY, has been sorted in ascending element order, as in Case(i) above.

### Examples.

Case (i):	SORT_UP( (/30, 20, 30, 40, $-10$ /) ) has the value $\begin{bmatrix} -10 & 20 & 30 & 30 & 40 \end{bmatrix}$ .	
Case (ii):	If <b>A</b> is the array $\begin{bmatrix} 1 & 9 & 2 \\ 4 & 5 & 2 \\ 1 & 2 & 4 \end{bmatrix}$ ,	
	then SORT_UP(A) has the value $\begin{bmatrix} 1 & 2 & 4 \\ 1 & 2 & 5 \\ 2 & 4 & 9 \end{bmatrix}$ ,	
Case (iii):	: If <b>A</b> is the array $\begin{bmatrix} 1 & 9 & 2 \\ 4 & 5 & 2 \\ 1 & 2 & 4 \end{bmatrix}$ ,	
		$\begin{bmatrix} 1 & 2 & 2 \end{bmatrix}$

then SORT\_UP(A, DIM = 1) has the value f(A) = 1

## SUM\_PREFIX(ARRAY, DIM, MASK, SEGMENT, EXCLUSIVE)

## **Optional Arguments.** DIM, MASK, SEGMENT, EXCLUSIVE

Description. Computes a segmented SUM scan along dimension DIM of ARRAY.

Class. Transformational function.

### Arguments.

		39
ARRAY	must be of type integer, real, or complex. It must not be	40
	scalar.	41
DIM (optional)	must be scalar and of type integer with a value in the	42
( - ),	range $1 \leq \text{DIM} \leq n$ , where n is the rank of ARRAY.	43
MACK (aptional)	must be of type lowicel and must be conformable with	44
MASK (optional)	must be of type logical and must be conformable with	45
	ARRAY.	46
<b>SEGMENT</b> (optional)	must be of type logical and must have the same shape as	47
	ARRAY.	48

1	EXCLUSIVE (optional)	must be of type logical and must be scalar.		
- 3 4	Result Type, Type Parameter, and Shape. Same as ARRAY.			
5 6 7 8	<b>Result Value.</b> Element r of the result has the value $SUM((/a_1,, a_m /))$ where $(a_1,, a_m)$ is the (possibly empty) set of elements of ARRAY selected to contribute to r by the rules stated in Section 7.4.5.			
9 10 11 12	Example. SUM_PREFIX( $\begin{bmatrix} 1 & 3 & 6 & 4 & 9 \end{bmatrix}$ .	(/1,2,3,4,5/), SEGMENT= (/F,F,F,T,T/) ) is		
13 14 <b>SUN</b>	1_SCATTER(ARRAY	,BASE,INDX1,, INDXn, MASK)		
15 16	Optional Argument. M	ASK		
17 18 19 20 21 22	indicated by index arrays	ements of ARRAY selected by MASK to positions of the result INDX1,, INDXn. Each element of the result is equal to ding element of BASE and the elements of ARRAY scattered		
22 23 24	Class. Transformational	function.		
24 25 26	Arguments.			
27 28 29	ARRAY	must be of type integer, real, or complex. It must not be scalar.		
30 31	BASE	must be of the same type and kind type parameter as ARRAY. It must not be scalar.		
32 33 34 35	INDX1,,INDXn	must be of type integer and conformable with <b>ARRAY</b> . The number of <b>INDX</b> arguments must be equal to the rank of <b>BASE</b> .		
36 37 38	MASK (optional)	must be of type logical and must be conformable with ARRAY.		
39 40	Result Type, Type Pa	rameter, and Shape. Same as BASE.		
41 42 43 44 45		ent of the result corresponding to the element $b$ of BASE has $, a_m, b/$ ) ), where $(a_1,, a_m)$ are the elements of ARRAY ibed in Section 7.4.4.		
46 47 48	Example. SUM_SCATTER $\begin{bmatrix} 7 & -1 & 7 \end{bmatrix}$ .	a((/1, 2, 3, 1/), (/4, -5, 7/), (/1, 1, 2, 2/)) is		

SUM_SUFFIX(ARRAY	, DIM, MASK, SEGMENT, EXCLUSIVE)	1		
<b>Optional Argument</b>	<b>Optional Arguments.</b> DIM, MASK, SEGMENT, EXCLUSIVE			
<b>Description.</b> Comp ARRAY.	utes a reverse, segmented SUM scan along dimension DIM of	4 5		
Class. Transformatio	nal function.	7 8		
Arguments.		EXCLUSIVE ed SUM scan along dimension DIM of eger, real, or complex. It must not be 1 of type integer with a value in the where $n$ is the rank of ARRAY. igical and must be conformable with ical and must be scalar. be. Same as ARRAY. e value SUM((/ $a_1,, a_m$ /)) where nents of ARRAY selected to contribute GMENT= (/F,F,F,T,T/) ) is ARRAY and the state of the stat		
ARRAY	must be of type integer, real, or complex. It must not be scalar.	11		
DIM (optional)	must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of ARRAY.	14		
MASK (optional)	must be of type logical and must be conformable with <b>ARRAY</b> .	16		
SEGMENT (optional)	must be of type logical and must have the same shape as ARRAY.			
EXCLUSIVE (optional)	must be of type logical and must be scalar.			
Result Type, Type	Parameter, and Shape. Same as ARRAY.			
	ent $r$ of the result has the value SUM((/ $a_1, \ldots, a_m$ /)) where ossibly empty) set of elements of ARRAY selected to contribute ed in Section 7.4.5.	24 25 26		
Example. SUM_SUFFI $\left[\begin{array}{cccc} 6 & 5 & 3 & 9 & 5 \end{array}\right]$ .	X( (/1,2,3,4,5/), SEGMENT= (/F,F,F,T,T/) ) is	28 29 30 31 32		
		35		
		38		
		40		
		42 43 44		

## Part III

# **HPF** Approved Extensions

This major section describes the syntax and semantics of features of approved extensions to High Performance Fortran. In most cases, these features build on concepts found in HPF itself; it may therefore be necessary to refer back to Parts I and II for background information.

## Section 8

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# **Approved Extensions for Data Mapping**

This section describes a set of data mapping features that extend the capabilities provided by the base set as described in Section 3. These extensions can be divided into two categories.

20 The first set of extensions provides the user greater control over the mapping of the data. 21 These include directives for dynamic remapping of data, which allow the user to redistribute 22 and realign at run time data that has been declared DYNAMIC. The ONTO clause used in 23 the **DISTRIBUTE** directive is extended to allow direct distribution to subsets of processors. 24 Explicit mapping of pointers and components of derived types are also introduced. Two  $^{25}$ new distributions are included: the GEN\_BLOCK distribution, which generalizes the block 26 distribution, and the INDIRECT distribution, which allows the mapping of individual array  $^{27}$ elements to be specified through a mapping array.

The programmer can use the second set of extensions to provide the compiler with 29 information useful for generating efficient code. This category includes the **RANGE** directive, 30 which allows the user to specify the range of distributions that a dynamically distributed 31 array, a pointer, or a dummy argument may have. The SHADOW directive allows the user to 32 specify the amount of additional space required on a processor to accommodate non-local 33 elements in a nearest-neighbor computation. 34

35 Since this section deals with extensions, we repeat some of the sections of Sections 3 36 and 4, providing new rules and extending old ones where necessary. In particular, subsections 8.13, 8.14 and 8.15 extend the corresponding subsections in Section 3 based on the 38 approved extensions described here.

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#### **Extended Model** 8.1

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The fundamental model for allocation of data to abstract processors still remains a two-level 46 mapping as described in Section 3. However, it is extended to allow the dynamic remapping 47of the data objects as illustrated by the following diagram: 48

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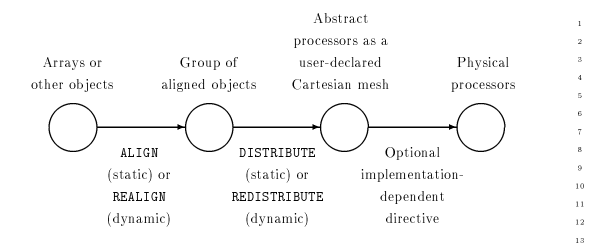
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Thus, objects can be remapped at execution time using the executable directives **REALIGN** and **REDISTRIBUTE**. Any object that is the root of an alignment tree (i.e., is not explicitly aligned to another object) can be explicitly redistributed. Redistributing such an object causes all objects ultimately aligned with it also to be redistributed so as to maintain the alignment relationships.

Any object that is not a root of an alignment tree can be explicitly realigned but not explicitly redistributed. Such a realignment does not change the mapping of any other object. Note that such remapping of data may require communication among the processors.

By analogy with the Fortran ALLOCATABLE attribute, HPF includes the DYNAMIC attribute. It is not permitted to REALIGN an array that has not been declared DYNAMIC. Similarly, it is not permitted to REDISTRIBUTE an array or template that has not been declared DYNAMIC.

Saved local variables, variables in common, and variables accessed by use association must not be implicitly remapped (e.g., by having variable distribution formats or being aligned with entities having variable distribution formats). Of these three categories of variables, only variables accessed by use association may have the DYNAMIC attribute.

As in Section 3.1, an object is considered to be *explicitly mapped* if it appears in an HPF mapping directive within the scoping unit in which it is declared; otherwise it is *implicitly mapped*. The definition of a mapping directive in Section 3.1 is extended as follows: A mapping directive is an ALIGN, DISTRIBUTE, INHERIT, DYNAMIC, RANGE, or SHADOW directive, or any directive that confers an alignment, a distribution, or the INHERIT, DYNAMIC, RANGE, or SHADOW attributes.

## 8.2 Syntax of Attributed Forms of Extended Data Mapping Directives

Like other mapping directives, the executable directives REALIGN and REDISTRIBUTE also come in two forms (statement form and attribute form) but may not be combined with other attributes in a single directive. The RANGE and SHADOW attributes may be combined with other attributes in a single directive.

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1	H801	$combined\-attribute\-extended$	is	ALIGN align-attribute-stuff
2			or	DISTRIBUTE dist-attribute-stuff
3			or	INHERIT
4			or	TEMPLATE
5			$\mathbf{or}$	PROCESSORS
6			or	DIMENSION ( <i>explicit-shape-spec-list</i> )
7			$\mathbf{or}$	DYNAMIC
8			$\mathbf{or}$	RANGE range-attr-stuff
9			or	SHADOW shadow-attr-stuff
10			or	SUBSET
11				

Constraint: The SUBSET attribute may be applied only to a processors arrangement.

The SUBSET attribute is discussed in Section 9; the rest are discussed below.

8.3 The REDISTRIBUTE Directive

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The REDISTRIBUTE directive is similar to the DISTRIBUTE directive but is considered executable. An object or template may be redistributed at any time, provided it has been declared DYNAMIC (see Section 8.5). Any other objects currently ultimately aligned with an array (or template) when it is redistributed are also remapped to reflect the new distribution, in such a way as to preserve alignment relationships (see Section 3.4). (This can require a lot of computational and communication effort at run time; the programmer must take care when using this feature.)

The DISTRIBUTE directive may appear only in the *specification-part* of a scoping unit. 27 The **REDISTRIBUTE** directive may appear only in the *execution-part* of a scoping unit. The 28 principal difference between DISTRIBUTE and REDISTRIBUTE is that DISTRIBUTE must con-29 tain only a *specification-expr* as the argument to a distribution format such as **BLOCK** or 30 CYCLIC, whereas in **REDISTRIBUTE** such an argument may be any integer expression. An-31 other difference is that DISTRIBUTE is an attribute, and so can be combined with other 32 attributes as part of a *combined-directive*, whereas **REDISTRIBUTE** is not an attribute (al-33 though a **REDISTRIBUTE** statement may be written in the style of attributed syntax, using 34 "::" punctuation). 35

The syntax of the **REDISTRIBUTE** directive is:

37 38 39 40	H802 redis	tribute- $directive$		REDISTRIBUTE distributee dist-directive-stuff REDISTRIBUTE dist-attribute-stuff :: distributee-list
41 42 43	Constraint:	A <i>distributee</i> that appears attribute (see Section 8.5		A REDISTRIBUTE directive must have the DYNAMIC
44 45 46	Constraint:	A <i>distributee</i> in a REDIST ALIGN or REALIGN directi		JTE directive may not appear as an <i>alignee</i> in an
47 48	Constraint:	Neither the <i>dist-format-c</i> may begin with "*".	laus	e nor the $dist$ -target in a REDISTRIBUTE directive

Note that, although an object may not have both the INHERIT attribute and the DISTRIBUTE attribute, any object—whether or not it has the INHERIT attribute—may appear as a *distributee* in a REDISTRIBUTE directive, provided that it has the DYNAMIC attribute and that it does not appear as an *alignee* in a ALIGN or REALIGN directive.

If a range directive (see Section 8.11) has been used to restrict the set of distribution formats allowed for a *distributee*, then the new mapping must match one of the formats specified in the range directive.

The statement form of a **REDISTRIBUTE** directive may be considered an abbreviation for an attributed form that happens to mention only one *distributee*; for example,

!HPF\$ REDISTRIBUTE distributee ( dist-format-list ) ONTO dist-target

is equivalent to

!HPF\$ REDISTRIBUTE ( dist-format-list ) ONTO dist-target :: distributee

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## 8.4 The REALIGN Directive

The REALIGN directive is similar to the ALIGN directive but is considered executable. An array (or template) may be realigned at any time, provided it has been declared DYNAMIC (see Section 8.5). Unlike redistribution (also in Section 8.5), realigning a data object does not cause any other object to be remapped. (However, realignment of even a single object, if it is large, can require a lot of computational and communication effort at run time; the programmer must take care when using this feature.)

The ALIGN directive may appear only in the *specification-part* of a scoping unit. The REALIGN directive is similar but may appear only in the *execution-part* of a scoping unit. The principal difference between ALIGN and REALIGN is that ALIGN must contain only a *specification-expr* as a *subscript* or in a *subscript-triplet*, whereas in REALIGN an expression as a *subscript* or in a *subscript-triplet* need not be a *specification-expr*. Another difference is that ALIGN is an attribute, and so can be combined with other attributes as part of a *combined-directive*, whereas REALIGN is not an attribute (although a REALIGN statement may be written in the style of attributed syntax, using "::" punctuation).

The syntax of **REALIGN** is as follows:

H803 realig	gn-directive is R	EALIGN alignee align-directive-stuff	34
	or R	EALIGN align-attribute-stuff :: alignee-list	35
			36
Constraint:	0 0 11	REALIGN directive must have the DYNAMIC at-	37
	tribute (see Section 8.5).		38
Constraint			39
Constraint: If the <i>align-target</i> specified in the <i>align-with-clause</i> has the DYNAMIC attribute, then each <i>alignee</i> must also have the DYNAMIC attribute.			40
	then each <i>ungrice</i> must also have		41
Constraint:	An <i>alignee</i> in a <b>REALIGN</b> directi	ve may not appear as a <i>distributee</i> in a	42
DISTRIBUTE or REDISTRIBUTE directive.		directive.	43
			44
Note th	at, although an object may not h	nave both the INHERIT attribute and the ALIGN	45

Note that, although an object may not have both the INHERIT attribute and the ALIGN 45 attribute, any object—whether or not it has the INHERIT attribute—may appear as an 46 alignee in a REALIGN directive, provided it has the DYNAMIC attribute and that it does not 47 appear as a distributee in a DISTRIBUTE or REDISTRIBUTE directive. 48 If a range directive (see Section 8.11) has been used to restrict the set of distribution formats allowed for an *alignee*, then the new mapping must match one of the formats specified in the range directive.

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## 8.5 The DYNAMIC Directive

The **DYNAMIC** attribute specifies that an object may be dynamically realigned or redistributed.

10	H804	$dynamic\mathchar`-directive$	is	${\tt DYNAMIC} \ a lignee-or-distributee-list$
11	H805	alignee-or-distributee	is	aliqnee
12	11000	anymee or aborriouree		distributee
13			01	
14				

- Constraint: An object in COMMON may not be declared DYNAMIC and may not be aligned to an object (or template) that is DYNAMIC. (To get this kind of effect, modules must be used instead of COMMON blocks.)
- <sup>18</sup> Constraint: A component of a derived type may have the DYNAMIC attribute only if it also
   <sup>19</sup> has the POINTER attribute. (See Section 8.9 for further discussion.)
- Constraint: An object with the SAVE attribute may not be declared DYNAMIC and may not be aligned to an object (or template) that is DYNAMIC.
- A REALIGN directive may not be applied to an *alignee* that does not have the DYNAMIC attribute. A REDISTRIBUTE directive may not be applied to a *distributee* that does not have the DYNAMIC attribute.

A DYNAMIC directive may be combined with other directives, with the attributes stated in any order, consistent with the Fortran attribute syntax.

- <sup>29</sup> Examples:
- 31 !HPF\$ DYNAMIC A,B,C,D,E
- <sup>32</sup> !HPF\$ DYNAMIC:: A,B,C,D,E
- 33 !HPF\$ DYNAMIC, ALIGN WITH SNEEZY:: X,Y,Z
- <sup>34</sup> !HPF\$ ALIGN WITH SNEEZY, DYNAMIC:: X,Y,Z
- 35 !HPF\$ DYNAMIC, DISTRIBUTE(BLOCK, BLOCK) :: X,Y
- <sup>36</sup> !HPF\$ DISTRIBUTE(BLOCK, BLOCK), DYNAMIC :: X,Y
- The first two examples mean exactly the same thing. The next two examples mean exactly the same second thing. The last two examples mean exactly the same third thing. The three directives

```
    <sup>41</sup> !HPF$ TEMPLATE A(64,64),B(64,64),C(64,64),D(64,64)
    <sup>42</sup> !HPF$ DISTRIBUTE(BLOCK, BLOCK) ONTO P:: A,B,C,D
    <sup>43</sup> UDEA DYNAMIC A D C D
```

HPF\$ DYNAMIC A,B,C,D

```
    may be combined into a single directive as follows:
```

```
47 !HPF$ TEMPLATE, DISTRIBUTE(BLOCK, BLOCK) ONTO P, &
48 !HPF$ DIMENSION(64,64), DYNAMIC :: A,B,C,D
```

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An ALLOCATABLE object may also be given the DYNAMIC attribute. If an ALLOCATE state-1 ment is immediately followed by REDISTRIBUTE and/or REALIGN directives, the meaning in 2 principle is that the array is first created with the statically declared mapping, if any, then 3 immediately remapped. In practice there is an obvious optimization: create the array in 4 the processors to which it is about to be remapped, in a single step. HPF implementors are 5strongly encouraged to implement this optimization and HPF programmers are encouraged to rely upon it. Here is an example:

	REAL,ALLOCATABLE(:,:) :: TINKER, EVERS	9
!HPF\$	DYNAMIC :: TINKER, EVERS	10
	REAL, ALLOCATABLE :: CHANCE(:)	11
!HPF\$	DISTRIBUTE(BLOCK),DYNAMIC :: CHANCE	12
	•••	13
	READ 6, M, N	14
	ALLOCATE(TINKER(N*M,N*M))	15
!HPF\$	REDISTRIBUTE TINKER(CYCLIC, BLOCK)	16
	ALLOCATE(EVERS(N,N))	17
!HPF\$	REALIGN EVERS(:,:) WITH TINKER(M::M,1::M)	18
	ALLOCATE(CHANCE(10000))	19
!HPF\$	REDISTRIBUTE CHANCE(CYCLIC)	20

While CHANCE is by default always allocated with a BLOCK distribution, it should be possible for a compiler to notice that it will immediately be remapped to a CYCLIC distribution. Similar remarks apply to TINKER and EVERS. (Note that EVERS is mapped in a thinlyspread-out manner onto TINKER; adjacent elements of EVERS are mapped to elements of TINKER separated by a stride M. This thinly-spread-out mapping is put in the lower left corner of TINKER, because EVERS(1,1) is mapped to TINKER(M,1).)

In Section 5.1, a list is given of operations that, if performed in a do loop, cause the iterations of the loop to interfere with each other, and thereby prevent the loop from being characterized as INDEPENDENT. To that list must be added:

• Any REALIGN or REDISTRIBUTE directive performed in the loop interferes with any access to or any other remapping of the same data.

*Rationale.* REALIGN and REDISTRIBUTE may change the processor storing a particular array element, which interferes with any assignment or use of that element. Similarly, multiple remapping operations may cause the same element to be stored in multiple locations. (End of rationale.)

#### 8.6 Remapping and Subprogram Interfaces

If the dummy argument of any subprogram has the DYNAMIC attribute, then an explicit interface is required for the subprogram (see subsection 8.14). The rules on the interaction of the REALIGN and REDISTRIBUTE directives with a subprogram argument interface are:

1. A dummy argument may be declared DYNAMIC. However, it is subject to the general restrictions concerning the use of the name of an array to stand for its associated template.

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- The effect of any redistribution of the dummy after the procedure returns to the caller is dependent on the attribute of the actual argument. If the actual argument associated with the dummy has also been declared DYNAMIC, then any explicit remapping of the dummy is visible in the caller after the procedure returns. If a range directive (see Section 8.11) has been used to restrict the set of distribution formats allowed for the actual argument, then the new mapping must match one of the formats specified in the range directive.
  - A dummy argument whose associated actual argument has the DYNAMIC attribute may be used in REALIGN and REDISTRIBUTE as an *alignee* or *distributee* if and only if the associated actual argument is a whole array, not an array section.
    - If the actual argument associated with the dummy has not been declared DYNAMIC then the original mapping of the actual has to be restored on return. When the subprogram returns and the caller resumes execution, all objects accessible to the caller after the call that are not declared DYNAMIC are mapped exactly as they were before the call.
  - 2. If an array or any section thereof is accessible by two or more paths, it is not HPFconforming to remap it through any of those paths. For example, if an array is passed as an actual argument, it is forbidden to realign that array, or to redistribute an array or template to which it was aligned at the time of the call, until the subprogram has returned from the call. This prevents nasty aliasing problems. An example:

```
MODULE FOO
23
                     REAL A(10,10)
^{24}
              !HPF$ DYNAMIC :: A
25
                     END
26
27
                     PROGRAM MAIN
28
                     USE FOO
29
                     CALL SUB(A(1:5,3:9))
30
                     END
31
32
                     SUBROUTINE SUB(B)
33
                     USE FOO
34
                     REAL B(:,:)
35
              !HPF$ DYNAMIC :: B
36
                     . . .
37
              !HPF$ REDISTRIBUTE A
                                                     !Nonconforming
38
                     . . .
39
                     END
40
```

Situations such as this are forbidden, for the same reasons that an assignment to A at the statement marked "Nonconforming" would also be forbidden. In general, in *any* situation where assignment to a variable would be nonconforming by reason of aliasing, remapping of that variable by an explicit REALIGN or REDISTRIBUTE directive is also forbidden.

<sup>47</sup> Note that it is permitted to remap a host-associated or use-associated variable in a <sup>48</sup> subprogram if it has been declared **DYNAMIC** and is accessible only through a single

path. Such remappings stay in effect even after the subprogram has returned to its caller.

## 8.7 Mapping to Processor Subsets

This extension allows objects to be directly distributed to processor subsets by allowing a processor subset to be specified where a processor could be named, e.g., in a **DISTRIBUTE** directive. The specified subset must be a proper subset of the named processor arrangement.

The syntax of the extended *dist-target* is as follows:

		10			
H806 exten	ded-dist-target is processors-name [ ( section-subscript-list ) ]	11			
	$\mathbf{or}$ * processors-name [ ( section-subscript-list ) ]	12			
	or *	13			
Constraint	The section-subscripts in the section-subscript-list may not be vector-subscripts	14			
Constraint.	and are restricted to be either <i>subscripts</i> or <i>subscript-triplets</i> .	15			
	and are restricted to be either subscripts of subscript-tripters.	16			
Constraint:	In the <i>section-subscript-list</i> , the number of <i>section-subscripts</i> must equal the	17			
	rank of the <i>processor-name</i> .	18			
a		19			
Constraint:	Within a DISTRIBUTE directive, each <i>section-subscript</i> must be a <i>specification-</i>				
	expr.	21			
Constraint:	Within a DISTRIBUTE or a REDISTRIBUTE directive, if both a <i>dist-format-list</i>	22 23			
	and a <i>dist-target</i> appear, the number of elements of the <i>dist-format-list</i> that				
	are not "*" must equal the number of <i>subscript-triplets</i> in the named processor arrangement.				
a					
Constraint:	Within a DISTRIBUTE or a REDISTRIBUTE directive, if a <i>dist-target</i> appears but	27 28			
	not a <i>dist-format-list</i> , the rank of each <i>distributee</i> must equal the number of	29			
	subscript-triplets in the named processor arrangement.				
!Example	1	31			
-	OCESSORS P(10)	32			
	AL A(100)	33			
	STRIBUTE A(BLOCK) ONTO P(2:5)	34			
		35			
		36			
!Example	2	37			
_	OCESSORS Q(10,10)	38			
	AL A(100,100)	39			
	STRIBUTE B(BLOCK, BLOCK) ONTO Q(5:10,5:10)	40			
		41			
In Evar	nple 1, the array A is distributed by block across the processors $P(2)$ to $P(5)$	42			

In Example 1, the array A is distributed by block across the processors P(2) to P(5) while in the second example, the array B is distributed across the lower right quadrant of the processor array Q.

Advice to users. This extension is most useful in conjunction with the tasking construct, see Section 9.4, which allows multiple independent phases of a computation to execute simultaneously on different subsets of processors. A similar situation arises

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when the code uses multiple data structures which can be computed in parallel where the computation on each individual object also exhibits parallelism, e.g., the multiple blocks in a multi-block grid used in some fluid dynamics calculation. Here, the individual blocks have to be distributed over subsets of processors to exploit both levels of parallelism. (*End of advice to users.*)

## 8.8 Pointers

## 8.8.1 Mapped Pointers

As an approved extension to HPF, pointers and targets can be explicitly mapped. Formally, this implies that the constraints that a *distributee* and an *alignee* may not have the POINTER or TARGET attribute as stated in Sections 3.3 and 3.4 respectively, have to be removed.

As in the case of an allocatable object, the mapping specification for a pointer does not take effect immediately but plays a role when the pointer becomes pointer associated with a target either through allocation or through pointer assignment.

When a pointer with an explicit mapping is used in an ALLOCATE statement, the data is allocated with the specified mapping.

```
For example:
```

```
REAL, POINTER, DIMENSION(:) :: A, B

!HPF$ ALIGN B(I) WITH A(I)

!HPF$ DISTRIBUTE A(BLOCK)

...

ALLOCATE(A(100))

ALLOCATE(B(50))

...

ALLOCATE(B(200)) ! Nonconforming
```

Pointer A is declared to have a BLOCK distribution while pointer B is declared to be
identically aligned with A. When A is allocated, it is created with a block distribution. When
B is allocated, it is aligned with the first 50 elements of A. Note that the allocation statements
may not occur in the opposite order, since an object may be aligned to another only if it
has already been created or allocated. Also, the second allocation for B is nonconforming,
since a larger object, B here, cannot be aligned with a smaller object, A in this case.

A pointer **P** with an explicit mapping can be pointer associated with a target **T** through a pointer assignment statement under the following conditions:

- 1. The mapping of T is a specialization of the mapping of P (in particular, T must be a whole array); and
  - 2. If **P** is explicitly aligned, its ultimate align target has a fully-specified non-transcriptive distribution; and
  - 3. P and T are either both DYNAMIC or neither is.
- Here are some examples:

```
47 REAL, POINTER, DIMENSION(:,:) :: P
```

<sup>48</sup> !HPF\$ DISTRIBUTE P(BLOCK, BLOCK)

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```
REAL, TARGET, DIMENSION (100, 100) :: B, C, D
!HPF$ DISTRIBUTE B(BLOCK, BLOCK)
!HPF$ DISTRIBUTE C(BLOCK, CYCLIC)
....
P => B ! Conforming
P => B(1:50, 1:50) ! Nonconforming: target must be a whole array.
P => C ! Nonconforming: the distribution in the
! second dimension does not match
P => D ! Nonconforming: D is not explicitly mapped
...
```

The intuitive reason that the pointer assignment  $P \Rightarrow B(1:50, 1:50)$  above is nonconforming is similar to the reason that the example on page 53 (illustrating the difference between INHERIT A and DISTRIBUTE A \* ONTO \*) is nonconforming: Suppose for instance that the array B is distributed over a  $2 \times 2$  processor arrangement. Then the section B(1:50, 1:50) would live entirely on processor (1, 1). This mapping is not correctly described by a (BLOCK, BLOCK) distribution for P.

The following pointer assignment is valid even though no processor arrangement is specified for the pointer; in this case, the mapping of B is a specialization of the mapping of P:

```
^{21}
      REAL, POINTER, DIMENSION(:) :: P
                                                                                     22
      REAL, TARGET, DIMENSION(100) :: B
                                                                                     23
!HPF$ PROCESSORS PROC(NUMBER_OF_PROCESSORS())
                                                                                     ^{24}
!HPF$ DISTRIBUTE P(BLOCK)
                                                                                     ^{25}
!HPF$ DISTRIBUTE (BLOCK) ONTO PROC :: B
                                                                                     26
      . . .
                                                                                     27
                            ! Conforming
      P => B
                                                                                     28
      . . .
                                                                                     29
                                                                                     30
      REAL, POINTER, DIMENSION(:) :: P
                                                                                     31
!HPF$ DISTRIBUTE * :: P
                                                                                     32
      REAL, TARGET, DIMENSION(100) :: B, C
                                                                                     33
!HPF$ DISTRIBUTE B(BLOCK), C(CYCLIC)
                                                                                     34
      . . .
                                                                                     35
      P => B
                            ! Conforming
                                                                                     36
      P => C
                            ! Conforming
                                                                                     37
                            ! Nonconforming: target must be a whole array
      P => C(1:50)
                                                                                     38
      . . .
                                                                                     39
```

Here, the \* is used to indicate a transcriptive distribution for the pointer P and thus it can be pointer associated with both targets B and C distributed by BLOCK and CYCLIC respectively. However, it still cannot be used to point to an array section such as C(1:50). To do that, the pointer must have the INHERIT attribute:

```
REAL, POINTER, DIMENSION(:) :: P45!HPF$ INHERIT :: P46REAL, TARGET, DIMENSION(100) :: B, C47!HPF$ DISTRIBUTE B(BLOCK), C(CYCLIC)48
```

1	••	•		
2	Р	=> B	! Conforming	
3	Р	=> C	! Conforming	
4	Р	=> C(1:50)	! Conforming	
5		•	C C	
6				
7	To allo	w pointers to ha	ve transcriptive distributions, we have to change the cons	straint
8	for dist-form	<i>nat-clause</i> as sp	ecified in Section 3.3, to read as follows:	介
9				
10	Constraint:	If either the $a$	<i>list-format-clause</i> or the <i>dist-target</i> in a DISTRIBUTE dim	rective
11			" then every distributee must be a dummy argument, ex	
12			has the POINTER attribute.	1 5
13				
14	The co	nstraint for <i>alia</i>	n-spec as specified in Section 3.4, should be changed to r	ead as↑
15	follows:	listianit ioi <i>ung</i>	<i>n-spec</i> as specified in Section 5.4, should be changed to r	cau as II
16	10110 w.5.			
17	<b>a</b>			,
18	Constraint:	0 1	c in an ALIGN directive begins with "*" then every aligned	e must
19		be a dummy a	rgument, except if the alignee has the POINTER attribute.	
20				
20		nstraint for <i>inhe</i>	eritee as specified in Section 4.4.2, should be changed to r	ead as $\uparrow$
	follows:			
22 23				
	Constraint:	An <i>inheritee</i> m	sust be a dummy argument, $except$ if the alignee has the PC	DINTER
24		attribute.		
25				
26	When 1	pointers with su	ch transcriptive mappings are used in an ALLOCATE state	ement,
27	-		y arbitrary mapping for the allocated data. A range decla	
28	-		sed to restrict the set of distribution formats.	
29	· ·	/	YNAMIC attribute, then any target associated with the p	pointer
30	-		have the DYNAMIC attribute) may be remapped using a RE	
31	· ·		t under the following restriction:	
32			l in REALIGN and REDISTRIBUTE as an <i>alignee</i> , <i>align-tar</i>	raet or
33	-	-	is currently associated with a whole array, not an array s	
34		-	ect is remapped, the new mapping is visible through any p	
35		a associated with		Jointer
36	that may be	associated with	i the object.	
37				
38	8.8.2 Po	inters and Su	ıbprograms	
39		1		. 1
40	-		nt is not explicitly mapped, then the actual argument mu	st also
41	-	citly mapped.		
42			gument has an explicit mapping, then the actual argumen	
43		-	assignment as stated above, with one exception: If the	
44			attribute, it is not necessary that the corresponding d	ummy
45	argument h	ave the DYNAMIC	C attribute. That is, item 3 on page 151 is weakened to	
46				
47			argument has the $\texttt{DYNAMIC}$ attribute, then the correspondence of the the correspondence of the transformation of tra	onding
48	actual	argument must	also have the DYNAMIC attribute.	

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A range declaration (see Section 8.11) can be used to restrict the set of distribution formats of the actual.

A pointer dummy argument may have the DYNAMIC attribute. In this case, the actual argument must also have the DYNAMIC attribute. The target associated with the dummy argument may be redistributed under the restrictions stated in the last subsection. Following Fortran rules, if the actual is also visible (through host- or use-association), the target may be redistributed only through the dummy argument. If the dummy argument is redistributed, then the actual argument has the new mapping on return from the procedure. In such a case, the new mapping must match the range restrictions (if any) of the actual.

#### 8.8.3 **Restrictions on Pointers and Targets**

If, on invocation of a procedure P: (a) a dummy argument has the TARGET attribute, and (b) the corresponding actual argument has the **TARGET** attribute and is not an array section with a vector subscript (and therefore is an object A or a section of an array A), then the program is HPF-conforming only if:

- 1. No remapping of the actual argument occurs during the call; or
- 2. the remainder of program execution would be unaffected if
  - (a) each pointer associated with any portion of A before the call were to acquire undefined pointer association status on entry to P and, if not reassigned during execution of P, were to be restored on exit to the pointer association status it had before entry.
  - (b) each pointer associated with any portion of the dummy argument or with any portion of A during execution of P were to acquire undefined pointer association status on exit from P; and

Advice to users. One way of ensuring that no remapping occurs is to give the dummy argument the INHERIT attribute. (End of advice to users.)

*Rationale.* These restrictions are made in order to support the following part of the Fortran standard (F95:12.4.1.1) in the face of implicit remapping across the subprogram interface:

If the dummy argument does not have the TARGET or POINTER attribute, any pointers associated with the actual argument do not become associated with the corresponding dummy argument on invocation of the procedure.

If the dummy argument has the TARGET attribute and the corresponding actual argument has the TARGET attribute but is not an array section with a vector subscript:

- 1. Any pointers associated with the actual argument become associated with the corresponding dummy argument on invocation of the procedure.
- 2. When execution of the procedure completes, any pointers associated 47 with the dummy argument remain associated with the actual argument.

```
If the dummy argument has the TARGET attribute and the corresponding
1
                actual argument does not have the TARGET attribute or is an array section
2
3
                with a vector subscript, any pointers associated with the dummy argument
                become undefined when execution of the procedure completes.
4
5
           (End of rationale.)
6
7
          Here is an example that illustrates the restrictions of this section:
8
9
               INTEGER, TARGET, DIMENSION (10) :: ACT
10
               INTEGER, POINTER, DIMENSON (:) :: POINTS_TO_ACT, POINTS_TO_DUM
11
        !HPF$ DISTRIBUTE ACT(BLOCK)
12
13
               POINTS_TO_ACT => ACT
14
               CALL F(ACT)
15
               POINTS_TO_DUM(1) = 1
                                                     ! ILLEGAL
16
17
18
               CONTAINS
19
                 SUBROUTINE F(DUM)
                   INTEGER, TARGET, DIMENSION(10) :: DUM
20
                 !HPF$ DISTRIBUTE DUM(CYCLIC)
21
22
                   POINTS_TO_DUM => DUM
23
                   POINTS_TO_ACT(1) = 1
                                                     ! ILLEGAL
24
                 END SUBROUTINE
25
               END
26
27
28
          The assignment to POINTS_TO_DUM(1) is illegal because it violates item 2b; the assign-
29
     ment to POINTS_TO_ACT(1) is illegal because it violates item 2a.
30
31
            Mapping of Derived Type Components
     8.9
32
33
     An ALIGN, DISTRIBUTE, or DYNAMIC directive may appear within a derived-type-def wherever
34
     a component-def-stmt may appear. Every alignee or distributee within such a directive must
35
     be the name of a component defined within that derived-type-def. To allow mapping of the
36
     structure components, the rules have to be extended as follows:
37
38
39
     H807 distributee-extended
                                            is
                                                object-name
40
                                            or template-name
41
                                            or component-name
42
                                            or structure-component
43
44
          A derived type is said to be an explicitly mapped type if any of its components is
45
     explicitly mapped or if any of its components is of an explicitly mapped type.
46
     Constraint: A component of a derived type may be explicitly distributed only if the type
47
                  of the component is not an explicitly mapped type.
48
```

Constraint:	An object of a derived ty type is not an explicitly n	-	nay be explicitly distributed only if the derived bed type.	1 2
Constraint:	A distributee in a DISTRI	BUTH	E directive may not be a <i>structure-component</i> .	3 4
Constraint:			directive which occurs in a <i>derived-type-def</i> must component of the derived type.	5 6
Constraint:	A component-name may curing within the derived		r as a <i>distributee</i> in a <b>DISTRIBUTE</b> directive oc- e definition only.	7 8 9
Constraint:	directive and every part-r	<i>∙ef</i> e	<i>re-component</i> may occur only in a <b>REDISTRIBUTE</b> xcept the rightmost must be scalar (rank zero). he <i>structure-component</i> must have the <b>DYNAMIC</b>	10 11 12 13 14
H808 align	ee-extended	is or or	object-name component-name structure-component	14 15 16 17
Constraint:	A component of a derived the component is not an e	•	pe may be explicitly aligned only if the type of citly mapped type.	18 19 20
Constraint:	An object of a derived ty is not an explicitly mapped	1	nay be explicitly aligned only if the derived type vpe.	21 22
Constraint:	An <i>alignee</i> in an ALIGN d	irect	ive may not be a <i>structure-component</i> .	23 24
Constraint:	An <i>alignee</i> in an <b>ALIGN</b> di <i>component-name</i> of a con		we that occurs in a <i>derived-type-def</i> must be the ent of the derived type.	25 26 27
Constraint:	A <i>component-name</i> may o within the derived type d		as an <i>alignee</i> only in an ALIGN directive occuring tion.	28 29
Constraint:	rective and every part-re	f ex	<i>re-component</i> may occur only in a <b>REALIGN</b> dicept the rightmost must be scalar (rank zero). he <i>structure-component</i> must have the <b>DYNAMIC</b>	30 31 32 33 34
H809 align	target-extended	is or or or	object-name template-name component-name structure-component	35 36 37 38
Constraint:			ear as an align target only in an ALIGN directive performance of the p	39 40 41
Constraint:	In an <i>align-target</i> that is rightmost must be scalar		<i>structure-component</i> , every <i>part-ref</i> except the k zero).	42 43 44
the derived components		that	nponents of derived type can be mapped within when any objects of that type are created the ied mapping.	44 45 46 47 48

Consider the following example:

1	TYPE DT
2	REAL C(100)
3	!HPF\$ DISTRIBUTE C(BLOCK) ONTO P
4	END TYPE DT
5	
6	TYPE (DT) :: S1
7	TYPE (DT) :: S2(100)
8	
9	a derived type with one component, array C, which is specified to be distributed block.
10	Therefore the scalar variable S1 of derived type DT has a structure component S1%C that is
11	distributed block onto the processor arrangement P. Similarly, the component C of each of
12	the elements of the array S2 will also be distributed block onto the processor arrangement
13	P.
14	An align directive inside a derived type definition may align a component of the derived
15	type with another component of the same derived type or with another object. A structure
16	component can be used as a target to align other objects including components of derived
17	types.
18	Example:
19	!HPF\$ TEMPLATE T(100)
20	!HPF\$ DISTRIBUTE T(CYLIC)
21	
22	TYPE DT
23	REAL, DIMENSION(100) :: A, B, C
24	!HPF\$ ALIGN WITH A :: B
25	!HPF\$ DISTRIBUTE (BLOCK) :: A
26	HPF\$ ALIGN WITH T :: C
27	END TYPE DT
28	
29	Here variables of derived type $DT$ will be created such the component $B$ is aligned with $A$ ,
30	which is itself distributed block, and such that the component $C$ is aligned with a template
31	T that is external to the derived type definition.
32	Note that if a derived type component is given a partial mapping, it is up to the
33	compiler to choose the rest of the mapping of that component. However, it is expected
34	that the compiler will choose the same mapping for this component of all variables of
35	such a derived type. For example, consider a modification of the above code in which the
36	distribution of the component $A$ is omitted. $B$ and $A$ are specified to be aligned but no
37	distribution is given for A. In such a situation, it is expected that all variables of the derived
38	type $DT$ will be created such that the component $A$ (and in turn the component $B$ ) have the
39	same distribution.

The constraints for the mapping of derived type components allow the mapping of structure variables at only one level. Consider for example the following code in which a derived type contains a components that is itself a derived type:

44	TYPE SIMPLE
45	REAL S(100)
46	!HPF\$ DISTRIBUTE S(BLOCK)
47	END TYPE SIMPLE
48	

```
!HPF$ TEMPLATE, DISTRIBUTE(BLOCK, *) :: HAIRY_TEMPLATE(47,73)
                                                                                      1
                                                                                      2
        TYPE COMPLICATED
                                                                                      3
          INTEGER SIZE
                                                                                      4
          REAL RV(100,100), KV(100,100), QV(47,73)
                                                                                      5
  ! Arrays RV, KV, and QV may be mapped
                                                                                      6
  !HPF$
          DISTRIBUTE (BLOCK, BLOCK) :: RV, KV
  !HPF$
          ALIGN WITH HAIRY_TEMPLATE :: QV
                                                                                      8
          TYPE(SIMPLE) SV(100)
                                                                                      9
  ! The following directive is not valid because SIMPLE
                                                                                      10
  ! is an explicitly mapped type.
                                                                                      11
  !HPF$
          DISTRIBUTE SV(BLOCK)
                                                                                      12
        END TYPE COMPLICATED
                                                                                      13
                                                                                      14
        TYPE(COMPLICATED) LOTSOF(20)
                                                                                      15
                                                                                      16
  ! The following directive is not valid because COMPLICATED
                                                                                      17
  ! is an explicitly mapped type.
                                                                                      18
  !HPF$ DISTRIBUTE LOTSOF(BLOCK)
                                                                                      19
                                                                                      20
Here, a component of the derived type SIMPLE has been mapped; thus objects of this
                                                                                      21
type, e.g., SV in type COMPLICATED, cannot be distributed. The array LOTSOF cannot be
                                                                                      22
distributed for the same reason.
                                                                                      23
    Structure components having the POINTER attribute can be remapped using the
                                                                                      ^{24}
REALIGN or REDISTRIBUTE directive if they have been declared DYNAMIC. For example, the
                                                                                      ^{25}
following code fragment can be used to allocate and map multiple blocks (called SUBGRID
                                                                                      26
here) of a multi-block grid:
                                                                                      27
                                                                                      28
  !HPF$ PROCESSORS P( number_of_processors() )
                                                                                      29
                                                                                      30
        TYPE SUBGRID
                                                                                      31
          INTEGER SIZE
                                                                                      32
          INTEGER LO, HI
                                    ! target subset of processors
                                                                                      33
          REAL, POINTER BL(:)
                                                                                      34
  !HPF$
          DYNAMIC BL
                                                                                      35
        END TYPE SUBGRID
                                                                                      36
                                                                                      37
        TYPE (SUBGRID), ALLOCATABLE :: GRID(:)
                                                                                      38
                                                                                      39
        READ (*,*) SUBGRID_COUNT
                                                                                      40
        ALLOCATE GRID(SUBGRID_COUNT)
                                                                                      41
        DO I = 1, SUBGRID_COUNT
                                                                                      42
          READ(*,*) GRID(I)%SIZE
                                                                                      43
        END DO
                                                                                      44
                                                                                      45
  ! Compute processor subsets for each subgrid, setting
                                                                                      46
  ! the LO and HI values
                                                                                      47
        CALL FIGURE_THE_PROCS ( GRID, number_of_processors())
                                                                                      48
```

1	! Allocate each subgrid and distribute to the computed processors subset
2	DO I = 1, SUBGRID_COUNT
3	ALLOCATE( GRID(I)%BL( GRID(I)%SIZE ) )
4	!HPF\$       REDISTRIBUTE GRID(I)%BL(BLOCK) ONTO P( GRID(I)%LO : GRID(I)%HI )
5	END DO
	END DO
6	Rationale. Components of derived types can be remapped only if they have the
7	POINTER attribute in addition to the DYNAMIC attribute. This restriction has been
8	placed to disallow mappings which cannot be directly specified using HPF directives.
9	Consider, for instance, the following code fragment:
10	Consider, for instance, the following code fragment.
11	!HPF\$ PROCESSORS P(4)
12	
13	TYPE DT
14	REAL C(100)
15	HPF\$ DISTRIBUTE C(BLOCK) ONTO P
16	
17	!HPF\$ DYNAMIC C ! Nonconforming END TYPE DT
18	
19	TYPE $(PT) \dots C(10)$
20	TYPE (DT) :: S(10)
21	···· I – 2
22	J = 3
23	 !HPF\$ REDISTRIBUTE S(J)%C(CYCLIC) ONTO P
24	HPP\$ REDISTRIBULE S(J)%C(CYCLIC) UNIO P
25	$C(\cdot, )$ $U(C(0))$
26	S(:)%C(2)
27	Here the component C of derived type DT has been deelened DYNAMIC. Thus, the error
28	Here the component C of derived type DT has been declared DYNAMIC. Thus, the array variable S consists of 10 elements each of which is a structure with a component C
29	initially distributed block. The <b>REDISTRIBUTE</b> directive remaps the structure com-
30	ponent $C$ of the Jth element of $S$ so that it is distributed cyclic. Consider now the
31	mapping of the data object referred to by the expression $S(:) \& C(2)$ which picks out
32	the second element from each of the ten structures that make up the array variable
33	<b>S</b> . After the redistribution of one of the elements of $S$ (element 3 in this case), each
34	element of the object will reside on processor $P(1)$ except for the third element, which
35	will reside on processor $P(2)$ . Such a distribution cannot be specified directly using
36	HPF directives.
37	
38	The Fortran standard disallows such expressions for components with the POINTER
39	attribute. In particular, if a <i>part-name</i> in a data reference has the <b>POINTER</b> attribute
40	then each <i>part-ref</i> to its left must be scalar (F95:6.1.2). Thus, we avoid the above
41	situation by
42	• disallowing the remapping of components that do not have the <b>POINTER</b> attribute,
43	• disanowing the remapping of components that do not have the POINTER attribute, and
44	
45	• relying on the Fortran standard to disallow expressions such as the above for
46	components with the POINTER attribute.
47	(End of rationale.)
48	

## 8.10 New Distribution Formats

This section describes two new distribution formats. The syntax is extended as follows:

- H810 extended-dist-format is BLOCK [ ( int-expr ) ] or CYCLIC [ ( int-expr ) ] or GEN\_BLOCK ( int-array ) or INDIRECT ( int-array ) or \*
- Constraint: An *int-array* appearing in a *extended-dist-format* of a DISTRIBUTE directive or REDISTRIBUTE directive must be an integer array of rank 1.
- Constraint: An *int-array* appearing in a *extended-dist-format* of a **DISTRIBUTE** directive must be a *restricted-expr*.
- Constraint: The size of any *int-array* appearing with a GEN\_BLOCK distribution must be equal to the extent of the corresponding dimension of the target processor arrangement.
- Constraint: The size of any *int-array* appearing with an INDIRECT distribution must be equal to the extent of the corresponding dimension of the *distributee* to which the distribution is to be applied.

The "generalized" block distribution, GEN\_BLOCK, allows contiguous segments of an array, of possibly unequal sizes, to be mapped onto processors. The sizes of the segments are specified by values of a user-defined integer mapping array, one value per target processor of the mapping. That is, the *ith* element of the mapping array specifies the size of the block to be stored on the *ith* processor of the target processor arrangement. Thus, the values of the mapping arrays are restricted to be non-negative numbers and their sum must be greater than or equal to the extent of the corresponding dimension the array being distributed.

The mapping array has to be a restricted expression when used in the DISTRIBUTE <sup>30</sup> directive, but can be an array variable in a REDISTRIBUTE directive. In the latter case, <sup>31</sup> changing the value of the map array after the directive has been executed will not change <sup>32</sup> the mapping of the distributed array. <sup>33</sup>

Let l and u be the lower and upper bounds of the dimension of the *distributee*, MAP be the mapping array and let BS(i):BE(i) be the resultant elements mapped to the *ith* processor in the corresponding dimension of the target processor arrangements. Then,

$$BS(1) = l,$$
  

$$BE(i) = \min(BS(i) + MAP(i) - 1, u),$$

$$BS(i) = BE(i-1) + 1.$$

Example:

 $^{21}$ 

!HPF\$ DYNAMIC B

```
2
                 . . .
3
               new = ...
        !HPF$ REDISTRIBUTE ( B( GEN_BLOCK(new) )
4
5
      Given the above specification, array elements A(1:2) are mapped on P(1), A(3:27) are
6
      mapped on P(2), A(28:47) are mapped on P(3), no elements are mapped on P(4), A(48:55)
7
      are mapped on P(5), and A(56:100) are mapped on P(6). The array B is distributed based
8
      on the array new whose values are computed at runtime.
9
10
           Advice to implementors. Accessing elements of an array distributed using the gen-
11
           eralized block distribution may require accessing the values of the mapping array at
12
           runtime. However, since the size of such an array is equal to that of the processor
13
           arrangement, it can in most cases be replicated over all processors.
14
           For dynamic arrays, an independent copy of the mapping array will have to be main-
15
           tained internally so that a change in the values of the mapping array does not affect
16
           the access of the distributed array. (End of advice to implementors.)
17
18
          There are many scientific applications in which the structure of the underlying domain is
19
      such that it does not map directly onto Fortran data structures. For example, in many CFD
20
      applications an unstructured mesh (consisting of triangles in 2D or tetrahedra in 3D) is used
21
      to represent the underlying domain. The nodes of such a mesh are generally represented by
22
      a one-dimensional array while another is used to represent their interconnections. Mapping
23
      such arrays using the structured distribution mechanisms, BLOCK and CYCLIC, results in
24
      mappings in which unrelated elements are mapped onto the same processor. This in turn
25
      leads to massive amounts of unnecessary communication. What is required is a mechanism
26
      to map a related set of arbitrary array elements onto the same processor. The INDIRECT
27
      distribution provides such a mechanism.
28
          The INDIRECT distribution allows a many-to-one mapping of elements of a dimension
29
      of a data array to a dimension of the target processor arrangement. An integer array is
30
      used to specify the target processor of each individual element of the array dimension being
31
      distributed. That is, the ith element of the mapping array provides the processor number
32
      onto which the ith array element is to be mapped. Since the mapping array maps array
33
      elements onto processor elements, the extent of the mapping array must match the extent of
34
      the dimension of the array it is distributing. Also, the values of the mapping array must lie
35
      between the lower and upper bound of the target dimension of the processor arrangement.
36
          The mapping array has to be a restricted expression when used in the DISTRIBUTE
37
     directive, but can be an array variable in a REDISTRIBUTE directive. In the latter case,
38
      changing the value of the mapping array after the directive has been executed will not
39
      change the mapping of the distributed array.
40
          Example:
41
42
        !HPF$ PROCESSORS P(4)
43
                 REAL A(100), B(50)
44
                 INTEGER map1(100), map2(50)
45
                 PARAMETER (map1 = /1,3,4,3,3,2,1,4, ..../)
46
        !HPF$ DYNAMIC B
47
        !HPF$ DISTRIBUTE A( INDIRECT(map1) ) ONTO P
48
```

!HPF\$ DISTRIBUTE map2(BLOCK) C	NTO P	1
		2
map2 =		3
!HPF\$ DISTRIBUTE B( INDIRECT(m	nap2) ) ONTO P	4
••••		5
Here, the array <b>A</b> is distributed st	atically using the constant array map1. Thus:	6
A(1) is mapped onto P(1	),	7
A(2) is mapped onto $P(3)$		8 9
A(3) is mapped onto P(4		9 10
A(4) is mapped onto P(3		10
A(5) is mapped onto $P(3)$		12
A(6) is mapped onto P(2		13
A(7) is mapped onto $P(1)$		14
A(5) is mapped onto P(4 The array B is declared dynamic	and is redistributed using the mapping array map2.	15
		16
	neral, the INDIRECT distribution is going to be used	17
	with an array variable as the map array. Also, since	18
	nust be the same as the array being distributed, it	19
	cely using the <b>BLOCK</b> distribution. This raises several	20
	his distribution, the runtime system should maintain	21
, , , , , , , , , , , , , , , , , , , ,	ing array so that if the program modifies the mapping change. Using an array variable as a mapping array	22
	element of the array will not be known until runtime.	23
-	required to figure out the location of a specific array	24
element. (End of advice to imple		25
		26 27
8.11 The RANGE Directive		21
		29
The RANGE attribute is used to restric	ct the possible distribution formats for an object or	30
template that has the $\texttt{DYNAMIC}$ attrib	ute or a transcriptive distribution format (including	31
pointers).		32
H811 range-directive	is RANGE ranger range-attr-stuff	33
5	5 5 66	34
H812 ranger	is object-name	35
	or <i>template-name</i>	36
H813 range-attr-stuff	${f is}$ range-distribution-list	37
H814 range-distribution	is ( range-attr-list )	38
IIII Tunge and no and no		39

39
40
41
42
43 44
45
46
47

## 8.11. THE RANGE DIRECTIVE

1	Constraint: At least one of the following must be true:
2	• The <i>ranger</i> has the DYNAMIC attribute.
3 4	• The <i>ranger</i> has the INHERIT attribute.
5	• The ranger is specified with a dist-format-clause of * in a DISTRIBUTE or
6	combined directive.
7 8	Constraint: The length of each range-attr-list must be equal to the rank of the ranger.
9 10	Constraint: The <i>ranger</i> must not appear as an alignee in an ALIGN or REALIGN directive.
11	Since the length of each range-attr-list is the same as the rank of the ranger, each
12	range-attr, $R$ , in each range-distribution corresponds positionally to a dimension $D$ of the
13	ranger. This dimension $D$ in turn either corresponds (though not necessarily positionally)
14 15	to an axis $A$ of the template with which the ranger is ultimately aligned, or corresponds to no axis in that template.
16	With this notation, a RANGE attribute on a <i>ranger</i> is equivalent to the following restric-
17	tion:
18	For at least one range-distribution in the range-distribution-list, every range-attr, R,
19	must either
20	• be compatible with the distribution format of the corresponding axis $A$ , if such a
21	corresponding axis exists, or
22 23	- he sither the ATT if no such corresponding suists
24	• be either $*$ or ALL, if no such corresponding axis exists.
25	This compatibility must be maintained by any DISTRIBUTE or REDISTRIBUTE directive
26	in which the <i>ranger</i> appears as a <i>distributee</i> , or if the ranger has the <b>POINTER</b> attribute and
27	is transcriptively distributed, for any target with which the <i>ranger</i> becomes associated. A distribution format of
28 29	
30	1. BLOCK is compatible with a $range-dist-format$ of BLOCK, BLOCK() or CYCLIC();
31 32	2. BLOCK(n) is compatible with a <i>range-dist-format</i> of BLOCK(), or CYCLIC();
33	3. CYCLIC is compatible with a <i>range-dist-format</i> of CYCLIC or CYCLIC();
34 35	4. CYCLIC(n) is compatible with a <i>range-dist-format</i> of CYCLIC();
36	5. GEN_BLOCK(a) is compatible with a <i>range-dist-format</i> of GEN_BLOCK;
37 38	6. INDIRECT(a) is compatible with a <i>range-dist-format</i> of INDIRECT;
39 40	7. $*$ is compatible with a <i>range-dist-format</i> of $*$ .
41 42	All distribution formats are compatible with a <i>range-dist-format</i> of ALL. Note that the possibility of a RANGE directive of the form
43 44	!HPF\$ RANGE range-attr-stuff-list :: ranger-list
45 46 47 48	is covered by syntax rule H301 for a <i>combined-directive</i> using <i>combined-attribute-extended</i> as defined in rule H801. Examples:

 $1\,1$ 

12

13

14

15 16

17

18

19 20

21

22

23

 $^{24}$ 

25 26

 $^{27}$ 

28

29

30 31

32

33

34

35

36

37

38

39

48

!HPF\$	DISTRIBUTE T(BLOCK)	1
!HPF\$	ALIGN A(I,J) WITH T(I)	2
		3
	CALL SUB(A)	4
		5
		6
	SUBROUTINE SUB(X)	7
!HPF\$	INHERIT X	8
!HPF\$	RANGE X (BLOCK, *), (CYCLIC, *)	9

Since the ultimate align target of X, the inherited template T in this case, does not have a second dimension, only a \* or ALL can be used in the second dimension of each range-distribution for X.

```
REAL A(100, 100, 100)
!HPF$
         DISTRIBUTE A(BLOCK, *, CYCLIC)
         CALL SUB(A(:,:,1))
                                        ! Conforming
         CALL SUB(A(:,1,:))
                                        ! Nonconforming
         CALL SUB( A(1,:,:) )
                                         ! Nonconforming
            . . . .
         SUBROUTINE SUB(X)
         REAL X(:, :)
!HPF$
         INHERIT X
!HPF$
         RANGE X (BLOCK, *)
```

Given the range directive in the subroutine SUB, only the first call to SUB is conforming. However, all three calls can be made conforming if the range directive above is replaced by the following directive:

!HPF\$ RANGE (BLOCK, \*), (BLOCK, CYCLIC), (\*, CYCLIC) :: X

## 8.12 The SHADOW Directive

In compiling nearest-neighbor code—for example, in discretizing partial differential equations or implementing convolutions—a standard technique is to allocate storage on each processor for the local array section so as to include additional space for the elements that have to be moved in from neighboring processors. This additional storage is referred to as "shadow edges." There are conceptually two shadow edges for each array dimension: one at the low end of the local array section and the other at the high end.

In a single routine, the compiler can tell which arrays require shadow edges and allocate 40 this additional space accordingly. However, since the width of the shadow area is dependent 41on the size of the computational stencil being used, an array may require different shadow 42 widths in different routines. Thus, without interprocedural analysis, an array argument 43 may need to be copied into a space with the appropriate shadow width on each procedure 44 call. A similar data motion would be required to copy the data back to its original location 45on exit from the subroutine. This unnecessary data motion can be avoided by allowing the 46 user to specify the required shadow width when the array is declared. 47

The syntax for declaring shadow widths is as follows:

```
H817 shadow-directive
                                             is
                                                 SHADOW shadow-target shadow-attr-stuff
1
2
      H818 shadow-target
                                             \mathbf{is}
                                                 object-name
3
                                             or
                                                component-name
4
      H819 shadow-attr-stuff
                                                 ( shadow-spec-list )
                                             \mathbf{is}
5
6
      H820 shadow-spec
                                             is
                                                 width
7
                                                low-width : high-width
                                             or
8
      H821 width
                                             is
                                                 int-expr
9
10
      H822 low-width
                                             is
                                                 int-expr
1\,1
      H823 high-width
                                             is
                                                 int-expr
12
13
      Constraint: The int-expr representing a width, low-width, or high-width must be a constant
14
                   specification-expr with value greater than or equal to 0.
15
16
17
          A shadow-spec of width is equivalent to a shadow-spec of width: width. Thus, the direc-
18
      tives
19
20
        !HPF$
                   DISTRIBUTE (BLOCK) :: A
21
        !HPF$
                  SHADOW (w) :: A
22
23
      specify that the array A is distributed BLOCK and is to have a shadow width of w on both
24
      sides. If A is a dummy argument, this gives the compiler enough information to inhibit
^{25}
      unnecessary data motion at procedure calls.
26
          Alternatively, different shadow widths can be specified for the low end and high end of
27
      a dimension. For example:
28
29
                   REAL, DIMENSION (1000) :: A
30
        !HPF$
                  DISTRIBUTE(BLOCK), SHADOW(1:2) :: A
31
                   . . . .
32
                   FORALL (i = 2, 998)
33
                      A(i) = 0.25 * (A(i) + A(i-1) + A(i+1) + A(i+2))
34
                   END FORALL
35
36
      specifies that only one non-local element is needed at the lower end while two are needed
37
      at the high end.
38
39
40
             Equivalence and Partial Order on the Set of Mappings
      8.13
41
42
      Section 4.5 has to be changed to accommodate the new distributions, the SHADOW attribute,
43
      and mapping of components of derived types, all introduced as approved extensions. The
44
      relevant text now reads as follows; additions are in bold-face type.
45
```

First, we define a notion of equivalence for *dist-format* specifications:

1. Using the notation  $\equiv$  for the phrase "is equivalent to",

46 47

```
↑
```

<ul> <li>BLOCK ≡ BLOCK CYCLIC ≡ CYCLIC * ≡ * BLOCK(n) ≡ BLOCK(m) iff m and n have the same value CYCLIC(n) ≡ CYCLIC(m) iff m and n have the same value CYCLIC ≡ CYCLIC(1) GEN_BLOCK(v) ≡ GEN_BLOCK(w) iff the values of corresponding elements of v and w are equal INDIRECT(v) ≡ INDIRECT(w) iff the values of corresponding elements of v and w are equal</li> <li>2. Other than this, no two lexically distinct dist-format specifications are equivalent. This is an equivalence relation in the usual mathematical sense. Next we define a notion of equivalence for SHADOW attributes (see Section 8.12 for the syntax):</li> <li>1. The shadow-spec expressions w₁ and w₂ are equivalent iff they have the same value.</li> <li>2. The shadow-spec w is equivalent to the shadow-spec l₂:h₂ iff l₁ is equiv- alent to l₂ and h₁ is equivalent to h₂.</li> <li>4. Other than this, no two lexically distinct shadow-spec specifications are equivalent.</li> <li>We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec list of the other. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>1. G has the INHERIT attribute, and the following constraints all hold:</li> </ul>
<ul> <li>CYCLIC ≡ CYCLIC</li> <li>* ≡ *</li> <li>BLOCK(n) ≡ BLOCK(m) iff m and n have the same value</li> <li>CYCLIC(n) ≡ CYCLIC(m) iff m and n have the same value</li> <li>CYCLIC ≡ CYCLIC(1)</li> <li>GEN_BLOCK(v) ≡ GEN_BLOCK(w) iff the values of corresponding elements of v and w are equal</li> <li>INDIRECT(v) ≡ INDIRECT(w) iff the values of corresponding elements of v and w are equal</li> <li>2. Other than this, no two lexically distinct dist-format specifications are equivalent.</li> <li>This is an equivalence relation in the usual mathematical sense.</li> <li>Next we define a notion of equivalence for SHADOW attributes (see Section 8.12 for the syntax):</li> <li>1. The shadow-spec expressions w₁ and w₂ are equivalent iff they have the same value.</li> <li>2. The shadow-spec w is equivalent to the shadow-spec u:w.</li> <li>3. The shadow-spec l₁:h₁ is equivalent to the shadow-spec l₂:h₂ iff l₁ is equivalent to l₂ and h₁ is equivalent to h₂.</li> <li>4. Other than this, no two lexically distinct shadow-spec specifications are equivalent.</li> <li>We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec-list of the other.</li> <li>Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects.</li> <li>The mapping of S is a specialization of the mapping of G if and only if either</li> <li>1. G has the INHERIT attribute, and the following constraints all hold:</li> </ul>
<ul> <li>* ≡ *</li> <li>BLOCK(n) ≡ BLOCK(m) iff m and n have the same value CYCLIC(n) ≡ CYCLIC(m) iff m and n have the same value CYCLIC ≡ CYCLIC(1)</li> <li>GEN_BLOCK(v) ≡ GEN_BLOCK(w) iff the values of corresponding elements of v and w are equal INDIRECT(v) ≡ INDIRECT(w) iff the values of corresponding elements of v and w are equal</li> <li>2. Other than this, no two lexically distinct dist-format specifications are equivalent. This is an equivalence relation in the usual mathematical sense. Next we define a notion of equivalence for SHADOW attributes (see Section 8.12 for the syntax):</li> <li>1. The shadow-spec expressions w<sub>1</sub> and w<sub>2</sub> are equivalent iff they have the same value.</li> <li>2. The shadow-spec w is equivalent to the shadow-spec l<sub>2</sub>:h<sub>2</sub> iff l<sub>1</sub> is equiv- alent to l<sub>2</sub> and h<sub>1</sub> is equivalent to h<sub>2</sub>.</li> <li>4. Other than this, no two lexically distinct shadow-spec specifications are equivalent.</li> <li>We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec-list of the other. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>1. G has the INHERIT attribute, and the following constraints all hold:</li> </ul>
<ul> <li>CYCLIC(n) ≡ CYCLIC(m) iff m and n have the same value CYCLIC ≡ CYCLIC(1)</li> <li>GEN_BLOCK(v) ≡ GEN_BLOCK(w) iff the values of corresponding elements of v and w are equal INDIRECT(v) ≡ INDIRECT(w) iff the values of corresponding elements of v and w are equal</li> <li>2. Other than this, no two lexically distinct dist-format specifications are equivalent. This is an equivalence relation in the usual mathematical sense. Next we define a notion of equivalence for SHADOW attributes (see Section 8.12 for the syntax):</li> <li>1. The shadow-spec expressions w<sub>1</sub> and w<sub>2</sub> are equivalent iff they have the same value.</li> <li>2. The shadow-spec u is equivalent to the shadow-spec u:w.</li> <li>3. The shadow-spec l<sub>1</sub>:h<sub>1</sub> is equivalent to the shadow-spec l<sub>2</sub>:h<sub>2</sub> iff l<sub>1</sub> is equivalent to l<sub>2</sub> and h<sub>1</sub> is equivalent to h<sub>2</sub>.</li> <li>4. Other than this, no two lexically distinct shadow-spec specifications are equivalent. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>1. G has the INHERIT attribute, and the following constraints all hold:</li> </ul>
<ul> <li>CYCLIC ≡ CYCLIC(1)</li> <li>GEN_BLOCK(v) ≡ GEN_BLOCK(w) iff the values of corresponding elements of v and w are equal INDIRECT(v) ≡ INDIRECT(w) iff the values of corresponding elements of v and w are equal</li> <li>2. Other than this, no two lexically distinct dist-format specifications are equivalent. This is an equivalence relation in the usual mathematical sense. Next we define a notion of equivalence for SHADOW attributes (see Section 8.12 for the syntax):</li> <li>1. The shadow-spec expressions w₁ and w₂ are equivalent iff they have the same value.</li> <li>2. The shadow-spec w is equivalent to the shadow-spec u:w.</li> <li>3. The shadow-spec l₁:h₁ is equivalent to h₂.</li> <li>4. Other than this, no two lexically distinct shadow-spec l₂:h₂ iff l₁ is equivalent. We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec-list of the other. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>1. G has the INHERIT attribute, and the following constraints all hold:</li> </ul>
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<ul> <li>elements of v and w are equal iff the values of corresponding elements of v and w are equal iff the values of corresponding elements of v and w are equal</li> <li>2. Other than this, no two lexically distinct dist-format specifications are equivalent. This is an equivalence relation in the usual mathematical sense. Next we define a notion of equivalence for SHADOW attributes (see Section 8.12 for the syntax):</li> <li>1. The shadow-spec expressions w<sub>1</sub> and w<sub>2</sub> are equivalent iff they have the same value.</li> <li>2. The shadow-spec w is equivalent to the shadow-spec w:w.</li> <li>3. The shadow-spec l<sub>1</sub>:h<sub>1</sub> is equivalent to the shadow-spec l<sub>2</sub>:h<sub>2</sub> iff l<sub>1</sub> is equivalent to l<sub>2</sub> and h<sub>1</sub> is equivalent to h<sub>2</sub>.</li> <li>4. Other than this, no two lexically distinct shadow-spec specifications are equivalent. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>1. G has the INHERIT attribute, and the following constraints all hold:</li> </ul>
<ul> <li>INDIRECT(v) ≡ INDIRECT(w) iff the values of corresponding elements of v and w are equal</li> <li>2. Other than this, no two lexically distinct dist-format specifications are equivalent. This is an equivalence relation in the usual mathematical sense. Next we define a notion of equivalence for SHADOW attributes (see Section 8.12 for the syntax):</li> <li>1. The shadow-spec expressions w<sub>1</sub> and w<sub>2</sub> are equivalent iff they have the same value.</li> <li>2. The shadow-spec w is equivalent to the shadow-spec w:w.</li> <li>3. The shadow-spec l<sub>1</sub>:h<sub>1</sub> is equivalent to the shadow-spec l<sub>2</sub>:h<sub>2</sub> iff l<sub>1</sub> is equivalent to l<sub>2</sub> and h<sub>1</sub> is equivalent to h<sub>2</sub>.</li> <li>4. Other than this, no two lexically distinct shadow-spec specifications are equivalent. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>1. G has the INHERIT attribute, and the following constraints all hold:</li> </ul>
<ul> <li>elements of v and w are equal</li> <li>2. Other than this, no two lexically distinct dist-format specifications are equivalent. This is an equivalence relation in the usual mathematical sense. Next we define a notion of equivalence for SHADOW attributes (see Section 8.12 for the syntax):</li> <li>1. The shadow-spec expressions w<sub>1</sub> and w<sub>2</sub> are equivalent iff they have the same value.</li> <li>2. The shadow-spec w is equivalent to the shadow-spec w:w.</li> <li>3. The shadow-spec l<sub>1</sub>:h<sub>1</sub> is equivalent to the shadow-spec l<sub>2</sub>:h<sub>2</sub> iff l<sub>1</sub> is equivalent to l<sub>2</sub> and h<sub>1</sub> is equivalent to h<sub>2</sub>.</li> <li>4. Other than this, no two lexically distinct shadow-spec specifications are equivalent.</li> <li>We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec-list of the other. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>1. G has the INHERIT attribute, or</li> <li>2. S does not have the INHERIT attribute, and the following constraints all hold:</li> </ul>
<ol> <li>Other than this, no two lexically distinct dist-format specifications are equivalent. This is an equivalence relation in the usual mathematical sense. Next we define a notion of equivalence for SHADOW attributes (see Section 8.12 for the syntax):         <ol> <li>The shadow-spec expressions w<sub>1</sub> and w<sub>2</sub> are equivalent iff they have the same value.</li> <li>The shadow-spec w is equivalent to the shadow-spec w:w.</li> <li>The shadow-spec l<sub>1</sub>:h<sub>1</sub> is equivalent to the shadow-spec l<sub>2</sub>:h<sub>2</sub> iff l<sub>1</sub> is equivalent to l<sub>2</sub> and h<sub>1</sub> is equivalent to h<sub>2</sub>.</li> <li>Other than this, no two lexically distinct shadow-spec specifications are equivalent.</li> </ol> </li> <li>We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec-list of the other. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>G has the INHERIT attribute, and the following constraints all hold:</li> </ol>
<ol> <li>Other than this, no two lexically distinct dist-format specifications are equivalent.         This is an equivalence relation in the usual mathematical sense.             Next we define a notion of equivalence for SHADOW attributes (see Section 8.12             for the syntax):      </li> <li>The shadow-spec expressions w<sub>1</sub> and w<sub>2</sub> are equivalent iff they have the same value.         The shadow-spec w is equivalent to the shadow-spec w:w.         The shadow-spec l<sub>1</sub>:h<sub>1</sub> is equivalent to the shadow-spec l<sub>2</sub>:h<sub>2</sub> iff l<sub>1</sub> is equivalent to l<sub>2</sub> and h<sub>1</sub> is equivalent to h<sub>2</sub>.     </li> <li>Other than this, no two lexically distinct shadow-spec specifications are equivalent.         We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec-list of the other.             Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects.             The mapping of S is a specialization of the mapping of G if and only if either             I. G has the INHERIT attribute, and the following constraints all hold:      </li> </ol>
<ul> <li>This is an equivalence relation in the usual mathematical sense. Next we define a notion of equivalence for SHADOW attributes (see Section 8.12 for the syntax):</li> <li>1. The shadow-spec expressions w<sub>1</sub> and w<sub>2</sub> are equivalent iff they have the same value.</li> <li>2. The shadow-spec w is equivalent to the shadow-spec w:w.</li> <li>3. The shadow-spec l<sub>1</sub>:h<sub>1</sub> is equivalent to the shadow-spec l<sub>2</sub>:h<sub>2</sub> iff l<sub>1</sub> is equivalent to h<sub>2</sub>.</li> <li>4. Other than this, no two lexically distinct shadow-spec specifications are equivalent.</li> <li>We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec-list of the other. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>1. G has the INHERIT attribute, and the following constraints all hold:</li> </ul>
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<ol> <li>The shadow-spec expressions w<sub>1</sub> and w<sub>2</sub> are equivalent iff they have the same value.</li> <li>The shadow-spec w is equivalent to the shadow-spec w:w.</li> <li>The shadow-spec l<sub>1</sub>:h<sub>1</sub> is equivalent to the shadow-spec l<sub>2</sub>:h<sub>2</sub> iff l<sub>1</sub> is equivalent to l<sub>2</sub> and h<sub>1</sub> is equivalent to h<sub>2</sub>.</li> <li>Other than this, no two lexically distinct shadow-spec specifications are equivalent.</li> <li>We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec-list of the other. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>G has the INHERIT attribute, or</li> <li>S does not have the INHERIT attribute, and the following constraints all hold:</li> </ol>
<ol> <li>The shadow-spec expressions w<sub>1</sub> and w<sub>2</sub> are equivalent iff they have the same value.</li> <li>The shadow-spec w is equivalent to the shadow-spec w:w.</li> <li>The shadow-spec l<sub>1</sub>:h<sub>1</sub> is equivalent to the shadow-spec l<sub>2</sub>:h<sub>2</sub> iff l<sub>1</sub> is equivalent to l<sub>2</sub> and h<sub>1</sub> is equivalent to h<sub>2</sub>.</li> <li>Other than this, no two lexically distinct shadow-spec specifications are equivalent.</li> <li>We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec-list of the other. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>G has the INHERIT attribute, or</li> <li>S does not have the INHERIT attribute, and the following constraints all hold:</li> </ol>
<ul> <li>same value.</li> <li>2. The shadow-spec w is equivalent to the shadow-spec w:w.</li> <li>3. The shadow-spec l<sub>1</sub>:h<sub>1</sub> is equivalent to the shadow-spec l<sub>2</sub>:h<sub>2</sub> iff l<sub>1</sub> is equivalent to l<sub>2</sub> and h<sub>1</sub> is equivalent to h<sub>2</sub>.</li> <li>4. Other than this, no two lexically distinct shadow-spec specifications are equivalent.</li> <li>We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec-list of the other. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>1. G has the INHERIT attribute, or</li> <li>2. S does not have the INHERIT attribute, and the following constraints all hold:</li> </ul>
<ol> <li>2. The shadow-spec w is equivalent to the shadow-spec w:w.</li> <li>3. The shadow-spec l<sub>1</sub>:h<sub>1</sub> is equivalent to the shadow-spec l<sub>2</sub>:h<sub>2</sub> iff l<sub>1</sub> is equivalent to l<sub>2</sub> and h<sub>1</sub> is equivalent to h<sub>2</sub>.</li> <li>4. Other than this, no two lexically distinct shadow-spec specifications are equivalent.</li> <li>We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec-list of the other. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>2. S does not have the INHERIT attribute, and the following constraints all hold:</li> </ol>
<ol> <li>2. The shadow-spec w is equivalent to the shadow-spec w:w.</li> <li>3. The shadow-spec l<sub>1</sub>:h<sub>1</sub> is equivalent to the shadow-spec l<sub>2</sub>:h<sub>2</sub> iff l<sub>1</sub> is equivalent to l<sub>2</sub> and h<sub>1</sub> is equivalent to h<sub>2</sub>.</li> <li>4. Other than this, no two lexically distinct shadow-spec specifications are equivalent.</li> <li>We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec-list of the other. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>2. S does not have the INHERIT attribute, and the following constraints all hold:</li> </ol>
<ul> <li>3. The shadow-spec l<sub>1</sub>:h<sub>1</sub> is equivalent to the shadow-spec l<sub>2</sub>:h<sub>2</sub> iff l<sub>1</sub> is equivalent to l<sub>2</sub> and h<sub>1</sub> is equivalent to h<sub>2</sub>.</li> <li>4. Other than this, no two lexically distinct shadow-spec specifications are equivalent.</li> <li>We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec-list of the other. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>1. G has the INHERIT attribute, or</li> <li>2. S does not have the INHERIT attribute, and the following constraints all hold:</li> </ul>
<ul> <li>alent to l<sub>2</sub> and h<sub>1</sub> is equivalent to h<sub>2</sub>.</li> <li>4. Other than this, no two lexically distinct shadow-spec specifications are equivalent.</li> <li>We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec-list of the other. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>G has the INHERIT attribute, or</li> <li>S does not have the INHERIT attribute, and the following constraints all hold:</li> </ul>
<ul> <li>4. Other than this, no two lexically distinct shadow-spec specifications are equivalent.</li> <li>We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec-list of the other. Now we define the partial order on mappings: Let S ("special") and G ("general") be two data objects. The mapping of S is a specialization of the mapping of G if and only if either</li> <li>G has the INHERIT attribute, or</li> <li>S does not have the INHERIT attribute, and the following constraints all hold:</li> </ul>
equivalent.24We then say that two SHADOW attributes are equivalent iff the shadow-spec-list30of one is elementwise equivalent to the shadow-spec-list of the other.31Now we define the partial order on mappings: Let S ("special") and G ("general") be32two data objects.33The mapping of S is a specialization of the mapping of G if and only if either341. G has the INHERIT attribute, or362. S does not have the INHERIT attribute, and the following constraints all hold:34
We then say that two SHADOW attributes are equivalent iff the shadow-spec-list of one is elementwise equivalent to the shadow-spec-list of the other.actionNow we define the partial order on mappings: Let S ("special") and G ("general") be two data objects.actionThe mapping of S is a specialization of the mapping of G if and only if eitheractionI. G has the INHERIT attribute, oraction2. S does not have the INHERIT attribute, and the following constraints all hold:action
of one is elementwise equivalent to the shadow-spec-list of the other.       31         Now we define the partial order on mappings: Let S ("special") and G ("general") be       32         two data objects.       33         The mapping of S is a specialization of the mapping of G if and only if either       34         1. G has the INHERIT attribute, or       36         2. S does not have the INHERIT attribute, and the following constraints all hold:       37
Now we define the partial order on mappings: Let S ("special") and G ("general") betwo data objects.The mapping of S is a specialization of the mapping of G if and only if either1. G has the INHERIT attribute, or2. S does not have the INHERIT attribute, and the following constraints all hold:
The mapping of S is a specialization of the mapping of G if and only if either341. G has the INHERIT attribute, or362. S does not have the INHERIT attribute, and the following constraints all hold:37
1. G has the INHERIT attribute, or       34         2. S does not have the INHERIT attribute, and the following constraints all hold:       34
<ol> <li>G has the INHERIT attribute, or</li> <li>S does not have the INHERIT attribute, and the following constraints all hold:</li> </ol>
2. S does not have the INHERIT attribute, and the following constraints all hold: $38$
2. S does not have the INHERIT attribute, and the following constraints all hold: $38$
(a) S is a named object or structure component, and $\frac{36}{40}$
(b) The shapes of the ultimate align targets of S and G are the same, and $41$
(c) Corresponding dimensions of S and G are mapped to corresponding dimensions
of their respective ultimate align targets, and corresponding elements of S and G
are aligned with corresponding elements of their respective ultimate align targets,
and
(d) Either 46
i. The ultimate align targets of both $S$ and $G$ are not explicitly distributed, or $\frac{44}{44}$

1	ii. The ultimate align targets of both S and G are explicitly distributed. In this
2	case, the distribution directive specified for the ultimate align target of G
3	must satisfy one of the following conditions:
4 5	A. It has no <i>dist-onto-clause</i> , or
6	B. It has a <i>dist-onto-clause</i> of "ONTO *", or
7	C. It has a <i>dist-onto-clause</i> specifying a processor arrangement having the
8	same shape as that explicitly specified in a distribution directive for the ultimate align target of S.
9	
10	and must also satisfy one of the following conditions:
11 12	A. It has no <i>dist-format-clause</i> , or
13	B. It has a <i>dist-format-clause</i> of "*", or
14 15	C. Each <i>dist-format</i> is equivalent (in the sense defined above) to the <i>dist-format</i> in the corresponding position of the <i>dist-format-clause</i> in an explicit distribution directive for the ultimate align target of S.
16 17	(e) Either S and G both have no SHADOW attribute or they have equivalent
18	SHADOW attributes.
19	
20	8.14 Conditions for Omitting Explicit Interfaces
21	
22	The requirements in Section 4.6 are extended as follows to account for the possible presence
23 24	of the DYNAMIC attribute; the addition is in <b>bold-face</b> type:
25	An explicit interface is required <i>except</i> when all of the following conditions hold:
26	1. Fortran does not require one, and
27	
28 29	2. No dummy argument is distributed transcriptively or with the INHERIT attribute, and
30	3. No dummy argument has the DYNAMIC attribute, and
31 32	4. For each pair of corresponding actual and dummy arguments, either:
33 34	(a) They are both implicitly mapped, or
35	(b) They are both explicitly mapped and
36	i. The mapping of the actual argument is a specialization of the mapping of
37	the dummy argument, and
38 39	ii. If the ultimate align targets of the actual and dummy arguments are both ex-
40	plicitly distributed, then the <i>dist-onto-clause</i> of each must specify processor
41	arrangements with the same shape.
42	and
43	
44 45	5. For each pair of corresponding actual and dummy arguments, either:
46 47	(a) Both are sequential, or
48	(b) Both are nonsequential.

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8.15	Characteristics of Procedures	1
<b>T</b> 1 <b>c</b>		2
	HADOW and DYNAMIC attributes, if present, are HPF-characteristics of dummy argu-	3
	and procedure return values. To be precise, the definitions in Section 4.7 are rewritten ows; additions are in <b>bold-face</b> type:	4
as 101	ows, additions are in <b>bold-lace</b> type.	5
•	A processor arrangement has one HPF-characteristic: its shape.	6
		7 8
٠	A template has up to three HPF-characteristics:	9
	1 its shape	10
	1. its shape;	11
	2. its distribution, if explicitly stated;	12
	3. the HPF-characteristic (i.e., the shape) of the processor arrangement onto which it is distributed, if explicitly stated.	13 14
•	A dummy data object has the following HPF-characteristics:	15
•	A dummy data object has the following m r-characteristics.	16
	1. its alignment, if explicitly stated, as well as all HPF-characteristics of its align target;	17 18
	2. its distribution, if explicitly stated, as well as the HPF-characteristic (i.e., the	19 20
	shape) of the processor arrangement onto which it is distributed, if explicitly stated;	20 21 22
	3. its SHADOW attribute, if explicitly stated.	23
		24
	A sta DVNAMLC of the but of it over legitly atotod	
	4. its DYNAMIC attribute, if explicitly stated.	25
	4. Its DYNAMIC attribute, if explicitly stated. A function result has the same HPF-characteristics as a dummy data object. Specif- ically, it has the following HPF-characteristics:	26 27
	A function result has the same HPF-characteristics as a dummy data object. Specif-	26
	A function result has the same HPF-characteristics as a dummy data object. Specif- ically, it has the following HPF-characteristics: 1. its alignment, if explicitly stated, as well as all HPF-characteristics of its align	26 27 28 29 30 31 32 33
	<ul> <li>A function result has the same HPF-characteristics as a dummy data object. Specifically, it has the following HPF-characteristics:</li> <li>1. its alignment, if explicitly stated, as well as all HPF-characteristics of its align target;</li> <li>2. its distribution, if explicitly stated, as well as the HPF-characteristic (i.e., the shape) of the processor arrangement onto which it is distributed, if explicitly stated;</li> </ul>	26 27 28 30 31 32 33 34
	<ul> <li>A function result has the same HPF-characteristics as a dummy data object. Specifically, it has the following HPF-characteristics:</li> <li>1. its alignment, if explicitly stated, as well as all HPF-characteristics of its align target;</li> <li>2. its distribution, if explicitly stated, as well as the HPF-characteristic (i.e., the shape) of the processor arrangement onto which it is distributed, if explicitly stated;</li> <li>3. its SHADOW attribute, if explicitly stated.</li> </ul>	26 27 28 29 30 31 32 33 34 35
	<ul> <li>A function result has the same HPF-characteristics as a dummy data object. Specifically, it has the following HPF-characteristics:</li> <li>1. its alignment, if explicitly stated, as well as all HPF-characteristics of its align target;</li> <li>2. its distribution, if explicitly stated, as well as the HPF-characteristic (i.e., the shape) of the processor arrangement onto which it is distributed, if explicitly stated;</li> </ul>	26 27 28 30 31 32 33 34 35 36
	<ul> <li>A function result has the same HPF-characteristics as a dummy data object. Specifically, it has the following HPF-characteristics:</li> <li>1. its alignment, if explicitly stated, as well as all HPF-characteristics of its align target;</li> <li>2. its distribution, if explicitly stated, as well as the HPF-characteristic (i.e., the shape) of the processor arrangement onto which it is distributed, if explicitly stated;</li> <li>3. its SHADOW attribute, if explicitly stated.</li> </ul>	26 27 28 29 30 31 32 33 34 35
	<ul> <li>A function result has the same HPF-characteristics as a dummy data object. Specifically, it has the following HPF-characteristics:</li> <li>1. its alignment, if explicitly stated, as well as all HPF-characteristics of its align target;</li> <li>2. its distribution, if explicitly stated, as well as the HPF-characteristic (i.e., the shape) of the processor arrangement onto which it is distributed, if explicitly stated;</li> <li>3. its SHADOW attribute, if explicitly stated.</li> </ul>	26 27 28 29 30 31 32 33 34 35 36 37
	<ul> <li>A function result has the same HPF-characteristics as a dummy data object. Specifically, it has the following HPF-characteristics:</li> <li>1. its alignment, if explicitly stated, as well as all HPF-characteristics of its align target;</li> <li>2. its distribution, if explicitly stated, as well as the HPF-characteristic (i.e., the shape) of the processor arrangement onto which it is distributed, if explicitly stated;</li> <li>3. its SHADOW attribute, if explicitly stated.</li> </ul>	26 27 28 29 30 31 32 33 34 35 36 37 38
	<ul> <li>A function result has the same HPF-characteristics as a dummy data object. Specifically, it has the following HPF-characteristics:</li> <li>1. its alignment, if explicitly stated, as well as all HPF-characteristics of its align target;</li> <li>2. its distribution, if explicitly stated, as well as the HPF-characteristic (i.e., the shape) of the processor arrangement onto which it is distributed, if explicitly stated;</li> <li>3. its SHADOW attribute, if explicitly stated.</li> </ul>	26 27 28 29 30 31 32 33 34 35 36 37 38 39
	<ul> <li>A function result has the same HPF-characteristics as a dummy data object. Specifically, it has the following HPF-characteristics:</li> <li>1. its alignment, if explicitly stated, as well as all HPF-characteristics of its align target;</li> <li>2. its distribution, if explicitly stated, as well as the HPF-characteristic (i.e., the shape) of the processor arrangement onto which it is distributed, if explicitly stated;</li> <li>3. its SHADOW attribute, if explicitly stated.</li> </ul>	26 27 28 30 31 32 33 34 35 36 37 38 39 40
	<ul> <li>A function result has the same HPF-characteristics as a dummy data object. Specifically, it has the following HPF-characteristics:</li> <li>1. its alignment, if explicitly stated, as well as all HPF-characteristics of its align target;</li> <li>2. its distribution, if explicitly stated, as well as the HPF-characteristic (i.e., the shape) of the processor arrangement onto which it is distributed, if explicitly stated;</li> <li>3. its SHADOW attribute, if explicitly stated.</li> </ul>	26 27 28 30 31 32 33 34 35 36 37 38 39 40 41
	<ul> <li>A function result has the same HPF-characteristics as a dummy data object. Specifically, it has the following HPF-characteristics:</li> <li>1. its alignment, if explicitly stated, as well as all HPF-characteristics of its align target;</li> <li>2. its distribution, if explicitly stated, as well as the HPF-characteristic (i.e., the shape) of the processor arrangement onto which it is distributed, if explicitly stated;</li> <li>3. its SHADOW attribute, if explicitly stated.</li> </ul>	26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41
	<ul> <li>A function result has the same HPF-characteristics as a dummy data object. Specifically, it has the following HPF-characteristics:</li> <li>1. its alignment, if explicitly stated, as well as all HPF-characteristics of its align target;</li> <li>2. its distribution, if explicitly stated, as well as the HPF-characteristic (i.e., the shape) of the processor arrangement onto which it is distributed, if explicitly stated;</li> <li>3. its SHADOW attribute, if explicitly stated.</li> </ul>	26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42 43 44
	<ul> <li>A function result has the same HPF-characteristics as a dummy data object. Specifically, it has the following HPF-characteristics:</li> <li>1. its alignment, if explicitly stated, as well as all HPF-characteristics of its align target;</li> <li>2. its distribution, if explicitly stated, as well as the HPF-characteristic (i.e., the shape) of the processor arrangement onto which it is distributed, if explicitly stated;</li> <li>3. its SHADOW attribute, if explicitly stated.</li> </ul>	26 27 28 30 31 32 33 34 35 36 37 38 39 40 41 41 42 43 44

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## Section 9

# Approved Extensions for Data and Task Parallelism

Modern parallel machines achieve their best performance if operations are performed by many processors with each processor accessing its own data. As such, the highest-performing programs will be those for which the computation partitioning and data mapping work in synergy. Three approved extensions provide the means to exploit this symmetry:

- 1. The ON directive partitions computations among the processors of a parallel machine (much as the DISTRIBUTE directive partitions the data among the processors).
- 2. The **RESIDENT** directive asserts that certain data accesses do not require interprocessor data movement for their implementation.
- 3. The TASK\_REGION construct provides the means to create independent coarse-grain tasks, each of which can itself execute a data-parallel (or nested task-parallel) computation.

All three constructs are related to the concept of *active processors*, introduced in Section 9.1 below. By assigning computations to processors, the ON directive (Section 9.2) defines the active processors. The **RESIDENT** directive (Section 9.3) uses this set and the information given by mapping directives in its assertions of locality. Finally, the **TASK\_REGION** construct (Section 9.4) builds its tasks from active processor sets.

## 9.1 Active Processor Sets

Active processors are an extension of the idea of processors and processors arrangements as used in HPF 2.0. HPF 2.0 assumes that a (static) set of processors exists, and that the program uses these processors to store data (e.g., through the DISTRIBUTE directive) and perform computations (e.g., by execution of FORALL statements). Finer divisions of the pro-cessor set are seldom mentioned, although they do have uses (e.g., mapping onto processor subsets as in an approved extension, Section 8.7, or in explaining the performance of com-putations on subarrays). Features such as task parallelism, however, require considering a more dynamic set of processors. In particular, to answer the question "What processor(s) is (are) currently executing?" it is important to define these features. 

Simply put, an active processor is one that executes an HPF statement (or group of
 statements). Active processors perform all operations required to execute the statement(s)

*except* (perhaps) for the initial access of data and writing of results. Some operations require certain processors to be active, as described below, but for the most part any processor can be active in the execution of any statement. An HPF program begins execution with all processors active. As described in Section 9.2, the ON directive restricts the active processor set for the duration of execution of statements in its scope. Consider this simple example (which has a reasonably intuitive meaning):

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!HPF\$ ON HOME( Z(INDX) )
 X(INDX-1) = X(INDX-1) + Y(INDX) \* Z(INDX+1)

Let X, Y, and Z have the same distribution, which does not replicate data. Following the ON directive, the statement would be executed as follows:

- 1. The processor owning Z(INDX) is identified as the active processor. On different executions of this ON block, this may be a different processor.
- 2. The values of X(INDX-1), Y(INDX), and Z(INDX+1) are made available to the active processor. Because of the identical distributions, Y(INDX) is already stored there. Depending on the data distribution and the hardware running the program, retrieving the others might correspond to the active processor loading registers from memory, or it might mean one or two other processors sending messages to the active processor.
- 3. The active processor performs an addition and a multiplication, using the values sent in the last step.
- 4. The result is stored to X(INDX-1), which may be on another processor. Again, this may require synchronization or other cross-processor operations.

There are considerable subtleties of this scheme when one of the statements involved is a function or subroutine call. Section 9.2.4 deals with these cases. Advice on the implementation of the ON directive is given in Section 9.2.2 below.

A few additional terms are useful in conjunction with the concept of active processors. 31 If all processors in a set are active, then the set is called an *active processor set*. The set of 32 all active processors is sometimes called *the* active processor set. This set is dynamic, and 33 if a statement is executed repeatedly the active processor set may be different each time. In 34 general, an HPF construct can only restrict the active set, not enlarge it. However, if the 35 original active set is partitioned into several independent sets, all partitions may execute 36 simultaneously. This is exactly how the TASK\_REGION construct (described in Section 9.4) 37 works. 38

The *universal* processor set is the set of all processors available to the HPF program. It is precisely the set of processors that is active when execution of the main program begins.

A processor that is not in the active set is called *inactive*. (Note that a processor 41 may be inactive with respect to one statement, but active with respect to another. This is 42 common in TASK\_REGION constructs.) 43

It is sometimes necessary to query properties of the active processor set; this is accomplished by the approved extension intrinsics ACTIVE\_NUM\_PROCS and ACTIVE\_PROCS\_SHAPE described in Section 12.1.

The data mapped to a processor is said to be *resident* on it. A replicated object is 47 resident on all of the processors that have a copy of it. 48

Rationale. It may seem odd at first to concentrate on shrinking the active processor set. However, HPF's design assumes that all processors are available at the beginning of execution. For example, implementing **DISTRIBUTE** requires information about the number of processors (in order to determine block sizes, for example) and their identity (in order to allocate the memory and perform data motion). Therefore, the execution model uses a static set of processors that can be subdivided and reunited dynamically. (End of rationale.)

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# 9.1.1 The SUBSET Directive

This subsection explains the interaction of active and inactive processor sets with explicitly mapped data. The rule of thumb is that allocating memory must be done locally; that is, if a processor stores part of an array, then that processor must be active when the array is created. Implications of this rule include:

- Local objects must be stored on a set of active processors, either when their subprogram is invoked or when they are allocated.
  - Dummy arguments are always mapped to a set of active processors. Section 9.2.4 explains the mechanism that ensures this.
  - Global objects (i.e., objects in COMMON or MODULEs or objects accessed via host association) may be explicitly mapped to inactive processors. However, those processors must have been active when the globals were allocated, whether at program initialization (when all processors were active), or at on entry to another subprogram, or on execution of an ALLOCATE statement.

It should be clear from the treatment of local and global objects that declarations may 27 need to refer to two classes of processors arrangements. The first, used mainly for the 28 declaration of global data, consists of arrangements of the universal processor set. These 29 are known as *universal* processors arrangements. Since they always represent the same 30 processors, these serve as a fixed frame of reference, allowing consistent declarations. A 31 processors-directive (Rule H329) defines a universal processors arrangement by default. To 32 accommodate active processors, two slight changes to the rules in Section 3.6 need to be 33 made: 34

- An HPF compiler is required to accept any *universal* processors arrangement that is scalar, or whose size (i.e., product of the arrangement's dimensions) is equal to the size of the universal processor set.
- If two *universal* processors arrangements have the same shape, then corresponding elements of the two arrangements are understood to refer to the same abstract processor.
- <sup>43</sup> In both cases, the only change is the addition of the word "universal."

Restricted processors arrangements represent only processors that, at the time the arrangement is declared, are active. They are used for mapping local objects and dummy arguments. To declare a subset processors arrangement, one can use the SUBSET option of *combined-attribute-extended* (H801), defined on page 145. One can also use the statement form of the SUBSET attribute: H901 subset-directive

is SUBSET processors-name

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Examples of the two forms are

```
!HPF$ PROCESSORS, SUBSET :: P(NP/4,4)
!HPF$ PROCESSORS Q(ACTIVE_NUM_PROCS())
!HPF$ SUBSET Q
```

As for universal arrangements, there are some modified rules for the use of subset processors arrangements:

- An HPF compiler is required to accept any *subset* processors arrangement that is scalar, or whose size is equal to the number of active processors, i.e., the number that would be returned by the call ACTIVE\_NUM\_PROCS().
- If two *subset* processors arrangements are declared with the same shape and the active processor set has not changed between their declarations, then corresponding elements of the two arrangements are understood to refer to the same abstract processor.

It is important to note that a scalar subset processors arrangement is considered to represent a processor that is active at the time the arrangement is created.

Note that it is permitted for a subset processors arrangements to have fewer than NUMBER\_OF\_PROCESSORS() elements; this reflects the way that the active processor set can shrink. Also note that there is an added condition before two subset processors arrangements are considered identical; this reflects the dynamic nature of the active processor set. Finally, note that a local, subset processors arrangement will be an arrangement of the set of active processors until such time as the active processor set is further restricted by an ON directive.

# 9.1.2 Mapping Local Objects and Dummy Arguments

For explicitly mapped local objects without the SAVE attribute, the declarations must map all elements of the object onto active processors. This requirement gives rise to several cases:

- If the local object is mapped via a DISTRIBUTE directive, then it must be distributed onto a set of active processors. One way, but not the only way, is to use a local, subset processor arrangement as the *dist-target*. If there is no explicit ONTO clause, the implementation is free to choose any arrangement of active processors as the *dist-target*.
- A local, universal processors arrangement of size one is always identified with an active processor, and may occur as the *dist-target* for a local object.
- If the local object is mapped by an ALIGN directive, then the corresponding elements of the ultimate align target must be distributed exclusively onto active processors. This certainly occurs if the whole of the ultimate align target is distributed onto active processors. It also occurs if the local object is aligned to a section of a target that is distributed onto both active and inactive processors, provided the section that is "hit" by the aligned object is mapped only to active processors. If the align replicates the alignee over one or more axes of the align target, then the distribution of the align target must ensure that all copies of the alignee are mapped to active processors.

In any of these cases, the active processor set is determined at the time that the DISTRIBUTE Or ALIGN becomes instantiated. That is, the mapping directives for ALLOCATABLE variables are instantiated when the variable is allocated; other objects have their mapping instantiated when they are declared.

The declaration of subset processors arrangements does not cause processors to become active or inactive; only the execution of ON directives does that. In particular, if a program contains no ON directives or constructs that modify a program's active processor set, then all processors are always active and all DISTRIBUTE directives can use universal arrangements.

Explicitly mapped global objects must have consistent mappings wherever they appear.
 This will usually (for COMMON and USE associated objects) be accomplished by distribution
 onto universal processors arrangements. Notice that the interpretation of an implicit (i.e.,
 missing) ONTO clause differs for local and global objects; globals may be distributed onto
 all processors, while locals must use only active processors. Also note that, since universal
 processors arrangements are the default for the PROCESSORS directive, no modification to
 the mapping of global objects is needed when active processors are introduced.

Dummy arguments must be explicitly mapped in the same way as local objects, using the rules above. As Section 9.2.4 explains, the effect of this is that dummy arguments are always stored on the active processor set. Other data objects, particularly objects local to the subprogram, can therefore be aligned to the dummy arguments and allocated on the active processor set.

Objects with the SAVE attribute must be mapped consistently whenever they come into scope. They are not subject to the restriction of mapping to active processors; where mapping is concerned, they conform to the same rules as global objects.

<sup>25</sup> 9.1.3 Other Restrictions on Active Processors

In addition to the mapping of locals and dummy arguments, several other constructs are restricted when the active processor set does not match the universal processor set. In general, the intent of these restrictions is to ensure that all processors that are needed for an operation are active when it is performed. In particular, allocating or freeing memory mapped to a processor requires the cooperation of that processor.

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For a **REDISTRIBUTE** directive, the active processor set must include:

- All processors that stored any element of the *distributee* before the **REDISTRIBUTE** was encountered, and
- The processors that will store any element of the *distributee* after the **REDISTRIBUTE** is performed.

This implies that all elements of the redistributed object reside on active processors, both before and after the REDISTRIBUTE operation. Effectively, this means that all data movement for the REDISTRIBUTE will be among active processors. In addition, the processors that owned the *distributee* (or anything aligned to it) beforehand can free the memory, and processors that now own the *distributee* can allocate memory for it.

Similarly, for a REALIGN directive, the set of active processors must include all processors
 that stored elements of the *alignee* before the REALIGN and all processors that will store
 *alignee* elements after the REALIGN.

For an ALLOCATE statement that creates an explicitly mapped object, the set of active processors must include the processors used by the mapping directive for the allocated object. The allocated object's ultimate align target may fall into one of two classes:

• Distributed with no explicit ONTO clause. (This case includes ultimate align targets 1 with no DISTRIBUTE directive at all.) In this case, the compiler must choose a set of 2 active processors that the object will be stored on. з 4 • Distributed ONTO a section of a processors arrangement. In this case, the specified 5section must be an arrangement of an active processor set. 6 7 For example: 8 !HPF\$ PROCESSORS P(NUMBER\_OF\_PROCESSORS()) 10 !HPF\$ ON (P(1:4)) CALL OF\_THE\_WILD() 11 12 . . . 13 SUBROUTINE OF\_THE\_WILD() 14 15INTEGER, ALLOCATABLE, DIMENSION(:) :: A, B, C, D, E, F !HPF\$ PROCESSORS P(NUMBER\_OF\_PROCESSORS()), ONE\_P 16 !HPF\$ PROCESSORS, SUBSET :: Q(ACTIVE\_NUM\_PROCS()) 1718 !HPF\$ DISTRIBUTE (BLOCK) :: A, E 19 !HPF\$ DISTRIBUTE (BLOCK) ONTO P(1:4) :: B !HPF\$ DISTRIBUTE (\*) ONTO ONE\_P :: C 20 !HPF\$ DISTRIBUTE (BLOCK) ONTO Q :: D, F 2122 23 ! No explicit ONTO; block size is probably 25 ALLOCATE (A(100)) ALLOCATE (B(100)) ! Block size IS 25  $^{24}$  $^{25}$ ALLOCATE (C(100)) ! On one active processor 26 ALLOCATE (D(100)) ! On Q(1:4); block size 25 27 !HPF\$ ON HOME(B(1:50)) BEGIN ALLOCATE (E(100)) ! No ONTO; E is allocated on Q(1:2) 28 ALLOCATE (F(100)) ! Nonconforming since Q(3:4) are inactive 29 !HPF\$ END ON 30

For a DEALLOCATE statement that destroys an explicitly mapped object, the active processor set must include all processors that own any element of that object. Again, there are two cases for the deallocated object's ultimate align target:

- Distributed onto a section of a processors arrangement. In this case, the processors that store part of the object must be active when it is deallocated. One way to guarantee this is to ensure that any ON block enclosing the DEALLOCATE statement also encloses the corresponding ALLOCATE.
- Distributed with no explicit ONTO clause. (This case includes ultimate align targets with no DISTRIBUTE directive at all.) In this case, the active processor set must include all the processors that were active when the object was allocated in order to guarantee that the processors that store part of the object are active when it is deallocated. Again, this is ensured if all ON blocks that enclose the deallocation also enclose the allocation operation.

An example may be helpful:

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REAL, ALLOCATABLE :: X(:), Y(:)
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        !HPF$ PROCESSORS P(8)
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        !HPF$ DISTRIBUTE X(BLOCK) ONTO P(1:4)
       !HPF$ DISTRIBUTE Y(CYCLIC)
4
5
       !HPF$ ON ( P(1:6) )
6
       !HPF$
                   ON (P(1:5))
7
                       ALLOCATE( X(1000), Y(1000)
8
                       ON ( P(1:3) )
        !HPF$
9
                            ! Point 1
10
11
       !HPF$
                       END ON
                        ! Point 2
12
       !HPF$
                   END ON
13
                  ! Point 3
14
        !HPF$ END ON
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16
               . . .
        !HPF$ ON ( P(1:4) )
17
                   ! Point 4
18
        !HPF$ END ON
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              ! Point 5
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At point 1, neither X nor Y can be deallocated, since some of the processors that store their elements might not be active. If the innermost directive were

!HPF\$ ON ( P(1:4) )

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25then X could be safely deallocated because of its explicit ONTO clause; it would still be 26 incorrect to deallocate Y. At points 2 and 3, both X and Y can safely be deallocated. In 27 general, if the deallocation occurs at the same level of ON nesting or at an outer level and 28 the flow of control has not left the outer ON construct, then the deallocation is safe. At 29 point 4 it is correct to deallocate X because its ONTO clause matches the enclosing ON. It is 30 not, however, correct to deallocate Y, since some processors (e.g., P(5)) that were active at 31 the ALLOCATE statement are not active at point 4. This illustrates the care that must be 32 exercised if a DEALLOCATE statement is controlled by an ON clause. One can avoid potential 33 problems by performing the deallocation outside of any ON construct in the same procedure, 34 as at point 5. 35

It is possible that only of subset of the processors active at allocation time and named in the ONTO clause actually store part of the object:

```
!HPF$ DISTRIBUTE A(BLOCK(10)) ONTO P(1:4)
    INTEGER, ALLOCATABLE :: A(:)
    ALLOCATE A(10)
!HPF$ ON (P(1))
    DEALLOCATE(A) ! Correct, because only P(1) owns any part of A
```

### 9.2 The ON Directive

The purpose of the ON directive is to allow the programmer to control the distribution of computations among the processors of a parallel machine. In a sense, this is the computational analog of the DISTRIBUTE and ALIGN directives for data. The ON directive does this by specifying the active processor set for a statement or set of statements. This temporarily shrinks the active processor set.

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If the computations in two ON block executions are not related (for example, if the ON block executions are two iterations of an INDEPENDENT loop), their ON directives give the compiler clear instructions for exploiting this potential parallelism.

### 9.2.1 Syntax of the ON Directive

There are two flavors of the ON directive: a single-statement form and a multi-statement form. The syntax for these directives is

H902	on-directive	is	ON on-stuff	11
11002		10		12
H903	on-stuff	is	home [ , resident-clause ] [ , new-clause ]	13
II.oo.4		•		14
H904	on-construct	$\mathbf{is}$		15
			directive-origin block-on-directive block	16
				17
			$directive$ - $origin\ end$ - $on$ - $directive$	18
H905	block- $on$ - $directive$	is	ON on-stuff BEGIN	19
HOOG	and an direction	:-	END ON	20
п900	end- $on$ - $directive$	is	END ON	21
H907	home	is	HOME ( $variable$ )	22
		or	HOME ( template-elmt )	23
		or	( processors-elmt )	24
H908	template-elmt	is	template-name [ ( section-subscript-list ) ]	25
11500		10		26
H909	processors- $elmt$	$\mathbf{is}$	processors-name [ ( $section$ -subscript-list ) ]	27

The nonterminal *resident-clause* will be defined in Section 9.3. For the present, it suffices to say that this is a form of the **RESIDENT** directive mentioned in the introduction.

The *home* is often called the HOME clause, even in cases where the keyword HOME is not used. Note that *variable* is a Fortran syntax term that means (roughly) "a reference, including an array element, array section, or derived type field"; *variable* does not include template or processor elements because they are defined only in directives. Note also that *block* is a Fortran syntax term for "a series of statements treated as a group"—for example, the body of a DO construct.

The on-directive is a kind of executable-directive (see rule H205). This means that an on-directive can appear wherever an executable statement can.

An *on-construct* is a Fortran *executable-construct*. This syntax implies that such constructs can be nested, and if so they will be properly nested.

*Rationale.* Note the use of parentheses in the last option of the *home* rule (involving *processors-elmt*). This prevents the following ambiguity:

INTEGER X(4)	! X(I) will be on processor I	45
!HPF\$ PROCESSORS HOME(4)		46
!HPF\$ DISTRIBUTE X(BLOCK)		47
X = (/ 4, 3, 2, 1 /)		48

1	! HPF\$ ON HOME(X(2))
2 3	X(2) = X(1)
4	If the parentheses were not required, where should the computation be done?
6	1. Processor HOME(2) (i.e., the owner of X(2))?
7	2. Processor HOME(3) (i.e., use the value of $X(2)$ , before the assignment)?
8 9	3. Processor HOME(4) (i.e., use the value of X(2), after the assignment)?
10 11 12	The definition of $ON$ clearly indicates that interpretation 1 is correct. One can get the effect of interpretation 2 by the directive
13 14	!HPF\$ ON(HOME(X(2)))
15 16 17 18 19	There is no way to get the effect of interpretation 3. Introducing reserved keywords into Fortran was suggested as a better solution to this problem, but was seen as too large a change to the underlying language. ( <i>End of rationale.</i> )
20 21	9.2.2 Semantics of the ON Directive
22 23 24 25	The ON directive restricts the active processor set for a computation to those processors named in its <i>home</i> . The computation controlled is either the following Fortran statement (for a <i>on-directive</i> or the contained <i>block</i> for a <i>block-on-directive</i> . We refer to the controlled computation as the ON-block.
26 27 28 29 30	That is, it advises the compiler to use the named processor(s) to perform the ON block. Like the mapping directives ALIGN and DISTRIBUTE, this is advice rather than an absolute commandment; the compiler may override an ON directive. Also like ALIGN and DISTRIBUTE the ON directive may affect the efficiency of computation, but not the final results.
31 32 33 34 35 36 37	Advice to implementors. If the compiler may override the user's advice in an ON directive, then the compiler should also offer the user an option to force all directives to be obeyed. Because dummy arguments and local objects are required to be mapped onto active processors, an HPF compiler that fails to heed the programmer's advice with respect to the active processor set may also be required to ignore some of the programmer's advice concerning data mapping. (End of advice to implementors.)
38	The single-statement ON directive sets the active processor set for the first non-comment
39	statement that follows it. It is said to apply to that statement. If the statement is a
40	compound statement (e.g., a DO loop or an IF-THEN-ELSE construct), then the ON directive
41 42	also applies to all statements nested therein. Similarly, the ON construct applies the initial ON clause to—i.e., sets the active processor set for—all statements up to the matching END
43	ON directive.
44	The evaluation of any function referred to in the home expression is not affected by
45	the ON directive; these functions are called on all processors active when control reached the
46	directive. Thus,
47	
48	!HPF\$ ON HOME( P(1: (ACTIVE_NUM_PROCS() - 1)) )

is a reasonable way to idle one active processor, and is not paradoxically self-referential.

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The HOME clause can name a program object, a template, or a processors arrangement. For each of these possibilities, it can specify a single element or multiple elements. This is translated into the processor(s) executing the ON block as follows:

• If the HOME clause names a program object, then every processor owning any part of that object should execute the ON block. For example, if A is an explicitly mapped array, then

!HPF\$ ON HOME ( A(2:4) )

tells the compiler to perform the statement on the processors owning A(2), A(3), and A(4). If A were distributed BLOCK, this might be one processor; if it were distributed CYCLIC, it would be three processors (assuming that many processors were available). Extra copies of elements created by a SHADOW directive (H817) are not taken into consideration by the HOME clause.

- If the HOME clause names a template element or section, then every processor owning any element of the template element or section should execute the ON block. The example above applies here as well, if A is a template rather than an array.
- If the HOME clause names a processors arrangement, then the processor(s) referenced there should execute the ON block. For example, if P is a processors arrangement, then

!HPF\$ ON ( P(2:4) )

will execute the following statement on the three processors P(2), P(3), and P(4).

In every case, the ON directive specifies the processor(s) that should perform a computation. More formally, it sets the active processors for the statements governed by the ON directive, as described in Section 9.1. That section also describes how some statements (notably ALLOCATE and dynamic remapping directives) require that particular processors be included in the active set. If one of these constructs occurs in the ON block and the active processor set does not contain all the required processors, then the program is not standard-conforming.

Note that the ON directive only specifies how *computation* is partitioned among processors; it does not indicate processors that may be involved in data transfer. Also, the ON clause by itself does not guarantee that its body can be executed in parallel with any other operation. However, placing the computation can have a significant effect on data locality. As later examples will show, the combination of ON and INDEPENDENT can also provide control over the load balance of parallel computations.

Advice to implementors. If the HPF program is compiled into Single-Program-40 Multiple-Data (SPMD) code, then the ON clause can always be implemented (albeit 41inefficiently) by having all processors compare their processor id to an id (or list of ids) 42 generated from the HOME clause. (Similar naive implementations can be constructed 43 in other paradigms as well.) If the ON clause will be executed repeatedly, for example 44 in a DO loop, it is worthwhile to invert this process. That is, instead of all processors 45executing all the HOME clause tests, the compiler should determine the range of loop 46 iterations that will test true on the given processor. (See the "Advice to implementors" 47in Section 9.2.3 for more details.) For example, consider the following complex case: 48 1

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DO I = 1, N !HPF\$ ON HOME( A(MY\_FCN(I)) ) BEGIN ... !HPF\$ END ON END DO

Here, the generated code can perform an "inspector" (i.e., a skeleton loop that only evaluates the HOME clause of each iteration) to produce a list of iterations assigned to each processor. This list can be produced in parallel, since MY\_FCN must be side-effect free (at least, the programmer cannot rely on any side effects). However, distributing the computation of *home* to all processors may require unstructured communications patterns, possibly negating the advantage of parallelism. In general, more advanced compilers will be able to efficiently invert more complex HOME clauses. It is recommended that the abilities (and limitations) of a particular compiler be documented clearly for users.

16Note that processors "screened out" by the naive implementation may still be re-17quired to participate in data transfer. If the underlying architecture allows one-sided 18 communication (e.g., shared memory or GET/PUT), this is not a problem. On message-19 passing machines, a request-reply protocol may be used. This requires the inactive 20 processors to enter a wait loop until the ON block completes, or requires the runtime 21 system to handle requests asynchronously. Again, it is recommended that the docu-22 mentation tell programmers which cases are likely to be efficient and which inefficient 23 on a particular system. (End of advice to implementors.)

Advice to users. The form of the *home* in an ON directive can be arbitrarily complex. 25This is a two-edged sword; it can express very complicated computation partitioning, 26 but the implementation of these partitions may not be efficient. More concretely, it 27 may express a perfectly load-balanced computation, but force the compiler to serialize 28 the computation to implement the HOME clauses. Although the amount of overhead 29 for an ON clause will vary based on the HPF code, the compiler, and the hardware, one 30 can expect that compilers will generate very good code based solely on array mappings 31 or a named processors arrangement, and progressively worse code as the complexity 32 of the *home* increases. A rough measure of the complexity of an ON directive is the 33 amount of run-time data used to compute it; for example, a constant offset is fairly 34 simple, while a permutation array is very complex. See Section 9.2.3 below for more 35 concrete examples of this phenomenon. 36

It should also be noted that the ON clause does not change the semantics of a program, in the same sense that DISTRIBUTE does not change semantics. In particular, an ON clause *by itself* does not change sequential code into parallel code, because the code in the ON block can still interact with code outside the ON block. (To put it another way, ON does not spawn processes.) (*End of advice to users.*)

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It is legal to nest ON directives, if the set of active processors named by the inner ON directive is included in the set of active processors from the outer directive. The syntax of *onconstruct* automatically ensures that it is properly nested inside other compound statements, and that compound statements properly nest inside of it. As with other Fortran compound statements, transfer of control to the interior of an *on-construct* from outside the block is prohibited: an *on-construct* may be entered only by executing the (executable) ON directive.

### 180 SECTION 9. APPROVED EXTENSIONS FOR DATA AND TASK PARALLELISM

Transfers within a block may occur. However, HPF *also* prohibits transfers of control from the interior of an *on-construct* to outside the *on-construct*, except by "falling through" the END ON directive. Note that this is stricter than in ordinary Fortran. If ON clauses are nested, then the innermost *home* effectively controls execution of the statement(s). A programmer can think of this as successively restricting the set of processors at each level of ON nesting; clearly, the last restriction must be the strongest. Alternately, the programmer can think of this as a fork-join approach to nested parallelism.

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Rationale. The restrictions about control flow into and out of an ON block essentially make it a single-entry single-exit region, thus simplifying the semantics considerably. (End of rationale.)

If an ON directive includes a NEW clause, the meaning is the same as a NEW clause in 13 an INDEPENDENT directive. The operation of the program would be identical if the NEW 14variables were allocated anew, and distributed onto the active processors, on every entry 15to the ON directive's scope, and deallocated on exit from the ON block. That is, the NEW 16variables are undefined on entry (i.e., assigned before use in the ON block) and undefined 17on exit (i.e., not used after the ON block, unless first reassigned). In addition, NEW variables 18 cannot be remapped in the ON clause's scope, whether by REALIGN, REDISTRIBUTE, or by 19argument association (at subroutine calls). If a variable appears in a NEW clause but does 20 not meet these conditions, then the program is not HPF-conforming. NEW variables are not 21considered by any nested **RESIDENT** directives, as detailed in Section 9.3. 22

The NEW variables are implicitly reallocated and remapped onto the active processors on entry to the ON block. For this reason, there are restrictions on their explicit mappings.

- An ON block NEW variable may not occur as an alignee.
- An ON block NEW variable may occur as a distributee only if there is no ONTO clause.

!HPF\$ DISTRIBUTE X(BLOCK, \*)
!HPF\$ DISTRIBUTE Y ONTO P ! Nonconforming due to ONTO clause
!HPF\$ ALIGN WITH X :: Z ! Nonconforming; ALIGN forbidden
!HPF\$ ON (P(1:4), NEW(X, Y, Z), BEGIN
!HPF\$ END ON

*Rationale.* NEW clauses provide a simple way to create temporary variables. This ability is particularly important when **RESIDENT** directives come into play, as will be clear below. (*End of rationale.*)

Advice to implementors. Because they are not used outside of the ON blocks, NEW 40 variables need not be kept consistent before and after ON clauses. Therefore, no 41 communication outside of the active processor set, determined by the ON directive, 42 is required to implement them. Scalar NEW variables should be replicated over the 43 active processor set, or allocated in memory areas shared by the active processor 44 set. Note that memory must be dynamically allocated if there is a possibility that 45multiple instances of the ON block could be active concurrently. This is similar to the  $^{46}$ requirements for implementing NEW variables in INDEPENDENT loops. (End of advice 47to implementors.) 48

#### 9.2. THE ON DIRECTIVE

9.2.3**Examples of ON Directives** 1

The following are valid examples of ON directives. Most of them illustrate idioms that programmers might want to use, rather than contrived situations. For simplicity, the first several examples assume the following array declarations:

> REAL A(N), B(N), C(N), D(N)!HPF\$ DISTRIBUTE A(BLOCK), B(BLOCK), C(BLOCK), D(BLOCK)

9 One of the most commonly requested capabilities for HPF is to control how loop iterations 10 were assigned to processors. (Historically, the ON clause first appeared to perform exactly  $1\,1$ this role in the Kali FORALL construct.) This can be done by the ON directive, as shown 12 in the following examples:

```
14
              !HPF$ INDEPENDENT
15
              DO I = 2, N-1
16
                 !HPF$ ON HOME(A(I))
17
                A(I) = (B(I) + B(I-1) + B(I+1))/3
18
              END DO
19
20
              !HPF$ INDEPENDENT
21
              DO J = 2, N-1
22
                 !HPF$ ON HOME(A(J+1)) BEGIN
23
                 A(J) = B(J+1) + C(J+1) + D(J+1)
^{24}
                 !HPF$ END ON
25
              END DO
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The ON directive in the I loop sets the active processor for each iteration of the loop to 27 be the processor that stores A(I). In other words, it advises the compiler to have each 28 processor run over its own section of the A array (and therefore B as well). The references 29 to B(I-1) and B(I+1) must be fetched from off-processor for the first and last iterations 30 on each processor (except for the boundary processors); note that those processors are not 31 mentioned in the HOME clause. The ON directive in the J loop similarly sets the active set for 32 each iteration, but advises the compiler to shift computations. As a result, each processor 33 does a vector sum of its own sections of B, C, and D, stores the first element of the result on 34 the processor to its left, and stores the rest of the result (shifted by one) in A. It is worth 35 noting that the directives would still be valid (and minimize nonresident data accesses) if 36 the arrays were distributed CYCLIC, although the number of nonresident references would 37 be much higher. 38

Advice to implementors. It is highly recommended that compilers concentrate on optimizing DO loops with a single ON clause including the entire loop body. Schematically, the code will be:

```
DO i = lb, ub, stride
44
                       !HPF$ ON HOME(array(f(i))) BEGIN
45
                          body
46
                       !HPF$ END ON
47
                    END DO
48
```

where array has some data mapping. Assume the mapping gives processor p the elements myset(p). (In a BLOCK distribution, for example, myset(p) is a contiguous range of integers.) Then the generated code on processor p should be

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DO 
$$i \in [lb:ub:stride] \cap f^{-1}(myset(p))$$

body

END DO

(This schematic does not show where communication or synchronization must be placed; that must be derived from analysis of the body.) Moreover, f is likely to be the identity function or a linear function with integer coefficients, both of which can be inverted easily. Given this, techniques for iterating through the set can be found in several recent conferences. (*End of advice to implementors.*)

Advice to users. One can expect the I loop above to generate efficient code for the 14computation partitioning. In effect, the compiler will arrange for each processor to 15iterate over its own section of array A. The J loop is slightly more complex, since the 16compiler must find the inverse of the HOME clause's subscripting function. That is, 17the compiler must solve K=J+1 for J, where K ranges over the resident elements of 18 A. Of course, in this case J=K-1; in general, linear functions can be inverted by the 19compiler. (It should be pointed out, however, that complex combinations of ALIGN 20 and DISTRIBUTE may make the description of K unwieldy, and this may add overhead 21to the inversion process.) (End of advice to users.) 22

Sometimes it is advantageous to "split" an iteration between processors. The following case shows one example of this:

```
!HPF$ INDEPENDENT
DO I = 2, N-1
    !HPF$ ON HOME(A(I))
    A(I) = (B(I) + B(I-1) + B(I+1))/3
    !HPF$ ON HOME C(I+1)
    C(I+1) = A(I) * D(I+1)
END DO
```

Here, the active processor sets for the two statements in the loop body are different. Due to the first ON clause, the reference to A(I) is resident in the first statement. The second ON clause makes A(I) nonresident (for some values of I) there. This maximizes the data locality in both statements, but does require data movement between the two.

Advice to implementors. If there are several non-nested ON clauses in a loop, then the schematic above needs to be generalized. In essence, the iteration range for each individual ON clause must be generated. A processor will then iterate over the union of these ranges. Statements guarded by an ON directive must now be guarded by an explicit test. In summary, the code for

DO $i = lb$ ,	ub, stride	45
!HPF\$	ON HOME( $array_1(f_1(i))$ )	46
$stmt_1$		47
!HPF\$	ON HOME( $array_2(f_2(i))$ )	48

1	$stmt_2$
2	END DO
3	
4 5	on processor $p$ becomes
6	$set_1 = [lb:ub:stride] \cap f_1^{-1}(myset_1(p))$
7	$set_2 = [lb:ub:stride] \cap f_2^{-1}(myset_2(p))$
8 9	DO $i \in set_1 \cup set_2$
10	IF ( $i \in set_1$ ) THEN
11	stmt1
12 13	END IF
14	IF ( $i \in set_2$ ) THEN
15	stmt2
16	END IF
17 18	
19	END DO
20	where $myset_1(p)$ is the resident set for $array_1$ , and $myset_2(p)$ is the resident set
21	for $array_2$ . (Again, synchronization and communication must be handled by other means.) Code transformations such as leap distribution and leap peoling can be used
22 23	means.) Code transformations such as loop distribution and loop peeling can be used to eliminate the tests in many cases. They will be particularly profitable if there are
24	data dependences between the ON blocks. (End of advice to implementors.)
25	
26	Advice to users. Splitting an iteration like this is likely to require either additional
27 28	tests at runtime or additional analysis by the compiler. Even if the compiler can generate low-overhead scheduling for the individual ON clauses, combining them is not
29	necessarily low-overhead. The locality benefits must be rather substantial for this
30	to pay off, but there are cases where multiple ON clauses are valuable. (All these
31	statements are particularly true if one $ON$ block uses data computed in another one.)
32	(End of advice to users.)
33 34	Because ON clauses nest naturally, they can be useful for expressing parallelism along
35	different dimensions. Consider the following examples:
36	
37	REAL X(M,M)
38 39	!HPF\$ DISTRIBUTE X(BLOCK,BLOCK)
40	!HPF\$ INDEPENDENT, NEW(I)
41	DO J = 1, $M$
42	<pre>!HPF\$ ON HOME(X(:,J)) BEGIN</pre>
43	DO I = 2, M !HPF\$ ON HOME(X(I,J))
44 45	X(I,J) = (X(I-1,J) + X(I,J)) / 2
46	END DO
47	!HPF\$ END ON
48	END DO

The active processor set for each iteration of the J loop is a column of the (presumably universal) processors arrangement. The I loop further subdivides the computation, giving each processor responsibility for computing the elements it owns. Many compilers would have chosen this computation partitioning automatically for such a simple example. How-ever, the compiler might have attempted to fully parallelize the outer loop, executing each inner loop sequentially on one processor. (This might be attractive on a machine with very fast communications.) By inserting the ON clauses, the user has advised against this strategy, thus trading additional locality for restricted parallelism. Notice that the ON direc-tive neither requires nor implies the INDEPENDENT assertion. In both nests, each iteration of the I loop depends on the preceding iteration, but the ON directive can still partition the computation among processors. The ON directive does not automatically make a loop  $1\,1$ parallel. 

Advice to implementors. "Dimension-based" nesting, as above, will probably be a common case. The HOME clauses can be inverted at each level, treating indices from outer loops as run-time invariants. (End of advice to implementors.)

Advice to users. Nested ON directives will tend to have efficient implementations if their HOME clauses refer to different dimensions of the processors arrangements, as in the above example. This minimizes the interaction between the levels of the loops, simplifying the implementation. (End of advice to users.)

Consider the following variation on the above example:

```
!HPF$ DISTRIBUTE Y(BLOCK,*)
!HPF$ INDEPENDENT, NEW(I)
DO J = 1, M
    !HPF$ ON HOME(Y(:,J)) BEGIN
    DO I = 2, M
        !HPF$ ON HOME(Y(I,J))
        Y(I,J) = (Y(I-1,J) + Y(I,J)) / 2
        END DO
        !HPF$ END ON
END DO
```

Note that the ON clauses have not changed, except for the name of the array. The interpretation is similar to the above, except that the outer ON directive assigns each iteration of the J loop to all of the processors. The inner ON directive again implements a simple owner-computes rule. The programmer has directed the compiler to distribute a serial computation across all the processors. There are a few scenarios where this is more efficient than parallelizing the outer loop:

- 1. Parallelizing the outer loop will generate many nonresident references, since only a part of each column is on any processor. If nonresident references are very expensive (or if M is relatively small), this overhead may outweigh any gain from parallel execution.
- The compiler may take advantage of the INDEPENDENT directive to avoid inserting any synchronization. This allows a natural pipelined execution. A processor will execute

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its part of the I loop for one value of J, then immediately go on to the next J iteration. Thus, the first processor will start on J=2 while the second receives the data it needs (from processor one) for J=1. (A similar pipeline would develop in the X example above.)

Clearly, the suitability of these ON clauses will depend on the underlying parallel architecture.

Advice to users. This example points out how ON may improve software engineering. While the "value" of HOME(X(I)) will change if X's mapping changes, its intent will usually stay the same - run the loop "aligned with" the array X. Moreover, the form of the clauses is portable, and they simplify experimenting with alternative computation partitioning. Both qualities are similar to the advantages of DISTRIBUTE and ALIGN over low-level data layout mechanisms. (End of advice to users.)

ON directives are particularly useful when the compiler cannot accurately estimate data locality, for example when the computation uses indirection arrays. Consider three variations of the same loop:

19	REAL X(N), Y(N)
20	INTEGER IX1(M), IX2(M)
21	!HPF\$ DISTRIBUTE X(BLOCK), Y(BLOCK)
22	<pre>!HPF\$ DISTRIBUTE IX(BLOCK), IY(BLOCK)</pre>
23	
24	!HPF\$ INDEPENDENT
25	DO I = 1, N
26	!HPF\$ ON HOME( X(I) )
27	X(I) = Y(IX(I)) - Y(IY(I))
28	END DO
29	
30	!HPF\$ INDEPENDENT
31	DO J = 1, N
32	!HPF\$ ON HOME( IX(J) )
33	X(J) = Y(IX(J)) - Y(IY(J))
34	END DO
35	
36 37	!HPF\$ INDEPENDENT
	DO K = 1, N
38 39	!HPF\$ ON HOME( X(IX(K)) )
40	X(K) = Y(IX(K)) - Y(IY(K))
40	END DO
41	

In the I loop, each processor runs over its section of the X array. (That is, the active processor for iteration I is the owner of X(I).) Only the reference X(I) is guaranteed to be resident. (If  $M \neq N$ , then IX and IY have a different block size than X, and thus a different mapping.) However, if it is *usually* the case that X(I), Y(IX(I)), and Y(IY(I)) are located on the same processor, then this choice of active processors may be the best available. (If X(I) and one of the other references are *always* on the same processor, then the programmer should add the **RESIDENT** clause as explained in Section 9.3.) In the next loop, iteration J's

active processor is the owner of IX(J). Because IY has the same distribution as IX, reference 1 IY(J) is always resident as well as IX(J). This is the most common array reference class in 2 the loop, so it minimizes the number of nonresident data references in the absence of any 3 special properties of IX and IY. It may not evenly balance the load among processors; for 4 example, if /(N=M/2) then half the processors will be idle. As before, if the values in 5IX or IY ensure that one of the Y references is always resident, a RESIDENT assertion should 6 be added. In the K loop, only reference Y(IX(K)) is guaranteed to be resident (because Y 7and X have the same distribution). However, the values stored in IX and IY may ensure 8 that Y(IY(K)) and X(K) are always resident. Even if the three REAL values are not always, 9 but merely "usually" on the same processor, this may be a good computation partitioning 10 for both locality and parallelism. However, these advantages must be weighed against the  $1\,1$ cost of computing this partitioning. Since the HOME clause depends on a (presumably large) 12array of runtime values, substantial time may be required to determine which iterations 13 are assigned to each processor. It should be clear from this discussion that there is no 14magic solution for handling complex computation partitionings; the best answer is usually 15a combination of application knowledge, careful data structure design (including ordering 16of the elements), and efficient compilation methodology and runtime support. 17

Advice to implementors. The K loop is the situation that the inspector strategy described above was designed for. If there is an outer loop around any of these examples, and that loop does not modify the distribution of X or the values of IX, then a record of each processor's iterations can be saved for reuse. The cost is at worst linear in the sizes of the arrays. (End of advice to implementors.)

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Advice to users. It is unlikely that any current production compiler will generate low-overhead code for K loop. The difference from previous examples is that the HOME clause is not a function that can be easily inverted by the compiler. Some compilers may choose to execute every iteration on all processors, testing the HOME clause at run-time; others may pre-compute a list of iterations for every processor. Of course, the cost of computing the list will be substantial.

In practice, one would make all the arrays the same size to avoid some of the alignment problems above; the example was written this way for pedagogical reasons, not as an example of good data structure design. (*End of advice to users.*)

# 9.2.4 ON Directives Applied to Subprogram Invocations

The key rule about ON directives when applied to subprogram invocations is that the invocation does not change the active processor set. In effect, the callee inherits the caller's active processors. Thus,

```
!HPF$ PROCESSORS P(10)
!HPF$ DISTRIBUTE X(BLOCK) ONTO P
!HPF$ ON ( P(1:3) )
        CALL PERSON_TO_PERSON()
!HPF$ ON ( P(4:7) )
        CALL COLLECT( X )
```

calls PERSON\_TO\_PERSON on three processors, while it calls COLLECT on four. The actual 47 argument to COLLECT does not reside completely on the active set of processors. This is 48

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allowed, with appropriate declarations of the corresponding dummy argument as explained below.

The above rule has interesting implications for data distributions within the called routine. In particular, dummy arguments must be declared under the same restrictions as local objects, thus ensuring that the dummy is always stored on the active processor set. This does not imply that the corresponding actual argument is local, however. Consider the possibilities for how a dummy can be explicitly mapped:

- **Prescriptive mapping:** If the actual is not mapped on the active processor set, it will be remapped. This is exactly analogous to remapping a **BLOCK**-distributed array to **CYCLIC** via a prescriptive mapping.
- **Descriptive mapping:** The user is asserting that the actual is already mapped onto the set of active processors. If the assertion is true, then the dummy is already stored locally; if not, then the compiler inserts a remapping operation (and reports a warning, following the recommendations in Section 4).
- **Transcriptive mapping:** In this case, a new restriction must be made to allow efficient access to the dummy argument. If a dummy is transcriptively mapped, then the actual argument must be resident on the active processor set at the time of the invocation. This may be checked at run-time.

In summary, a dummy argument is always mapped to the set of active processors, although
 the actual argument need not be (except in the case of transcriptive mappings).

*Rationale.* The treatment of dummy arguments as local objects is consistent with all 24 previous Fortran (and FORTRAN) standards. Moreover, it has the advantage that 25it reflects the usual expectations and wishes of programmers. Dummy arguments are 26 not expected to create great inefficiencies in Fortran programs; ensuring that they are 27 always stored locally tends to reinforce that expectation. Also, programmers are used 28 to "pass by reference" behavior, in which arguments are not copied; the restrictions 29 on data mapping to active processor sets allow this implementation when the data 30 is not remapped on subprogram call. One case of this deserves special mention— 31 transcriptive mappings. If the programmer wants to keep the data in place (the usual 32 expectation of INHERIT and related features) and control which processors execute 33 the computation (the meaning of ON), then the basic principles of active sets (set 34 forth in Section 9.1) imply that the data must be resident before the call is made. 35 When remapping occurs due to explicit directives, then surely the user expects a 36 communication cost to accompany the remapping. 37

- It should also be noted that these rules do not invalidate any HPF programs written without using the ON directive. In those programs, the active processor set never changes (at least, not from the view of the language). Therefore, subset and universal processors arrangements can be used interchangeably, and the restriction on use of transcriptive mappings is obeyed automatically. (*End of rationale.*)
- Advice to implementors. These restrictions imply that accesses to dummy arguments never require one-sided communication if the argument is explicitly mapped and the ON clause is used. Of course, accesses to global data may still run into serious complications. If the compiler itself partitions the computation, it is not restricted by the N directive rules. (End of advice to implementors.)

Advice to users. The idea to remember in calling subprograms from an ON block is this: make sure that the actual arguments are stored on the active set. If the subroutine interface uses transcriptive ("take anything") mappings, then this is a requirement. If the subroutine uses any other type of mapping, then having resident actual arguments may avoid the expense of remapping data. (Of course, it does not by itself guarentee that remapping doesn't occur—a prescriptive interface can force a BLOCK-to-CYCLIC redistribution—but it does ensure that the remapping is between active processors. This allows simpler and more efficient collective communications operations to be generated in the runtime system.) (End of advice to users.)

Let us return to the previous example:

Let us return to the previous example:	11
!HPF\$ PROCESSORS P(10)	12
!HPF\$ DISTRIBUTE X(BLOCK) ONTO P	13
	14
!HPF\$ ON ( P(4:7) )	15
CALL COLLECT(X)	16
	17
If COLLECT were declared as	18
SUBROUTINE COLLECT( A )	19
!HPF\$ DISTRIBUTE A(CYCLIC)	20
	21
then the call will be executed as follows:	22
	23
1. X will be remapped from BLOCK on 10 processors (i.e., all of P) to CYC $(i.e., P(4,7))$ . This will be a mapped to mapped and pattern	LIC on 4 pro- <sup>24</sup> 25
cessors (i.e., P(4:7)). This will be a many-to-many exchange pattern.	25
2. COLLECT will be called on processors P(4), P(5), P(6), and P(7). Access	
the subroutine will be satisfied from the redistributed array on those p	
	20
3. A will be remapped back to the distribution of $X$ . This is the inverse of	step 1. 30
Note that the distribution of A is onto 4 processors (the active processor set in	uside the call). <sup>31</sup>
not onto the universal processor set. If the interface is	32
	33
SUBROUTINE COLLECT( A )	34
!HPF\$ DISTRIBUTE A(BLOCK)	35
then the process would be the same, except that there would be a remappin	g from BLOCK <sup>36</sup>
on 10 processors to BLOCK on 4 processors. That is, the block size would in	
times (with related shuffling of data) and then revert to the original. Again, i	
to note that the distribution of <b>A</b> is onto the active processor set rather than	-
The similar examples	40
	41
REAL X(100,100), Y(100,100)	42
!HPF\$ PROCESSORS P(4), Q(2,2)	43
!HPF\$ DISTRIBUTE X(BLOCK,*) ONTO P	44
!HPF\$ DISTRIBUTE Y(BLOCK,BLOCK) ONTO Q	45
	46
INTERFACE	47
SUBROUTINE A_CAB( B )	48

1	REAL B(:)
2	!HPF\$ DISTRIBUTE B *(BLOCK)
3	END INTERFACE
4	
5	!HPF\$ ON ( P(4:7) )
6	CALL A_CAB( X( 1:100, 1 )
7	!HPF\$ ON HOME(X(1:100,1))
8	CALL A_CAB( X(1:100,100) ) !HPF\$ ON HOME( Y(1:100,1) )
9 10	CALL A_CAB( Y(1:100,1) )
11	!HPF\$ ON HOME( Y(99,1:100) )
12	CALL A_CAB( Y(99,1:100) )
13	
14	can be explained as follows. Calling A_CAB(1:100,1) on P(4:7) will produce a remapping
15	from 10 processors to 4, as in the example above. (The compiler would be expected to
16	produce a warning in this case, as explained in Section 4.) Calling A_CAB(X(1:100,100)) on
17	HOME(X(1:100,1)) produces no such remapping (or warning), because the active processor
18	set does not change; therefore, the descriptive mapping correctly asserts that the data is
19	already on the right processors. The last two examples, calling $A\_CAB(Y(1:100,1))$ and $A\_CAB(Y(00,1:100))$ on the homes of their computer are also accomplished without
20	A_CAB(Y(99,1:100)) on the homes of their arguments, are also accomplished without remapping. In both cases, the actual arguments are mapped BLOCK-wise onto a subset of
21	the processors (a column of Q in the first case, a row of Q in the second). Some compilers
22	may not be able to generate code for these more complex examples, however.
23	Two examples of transcriptive mapping are also useful:
24	1 no champios of champeriperio mapping are also aberali
25	! Assume
26 27	! PROCESSORS P(4)
28	! is declared in a module
29	REAL X(100)
30	!HPF\$ DISTRIBUTE X(CYCLIC(5)) ONTO P
31	INTERFACE
32	SUBROUTINE FOR_HELP( C )
33	REAL C(:)
34	HPF\$ INHERIT C
35	END INTERFACE
36	
37	!HPF\$ ON HOME( X(11:20) )
38	CALL FOR_HELP( X(11:20) )
39	!HPF\$ ON ( P(1) )
40	CALL FOR_HELP( X(51:60) ) ! Nonconforming
41	
42	The first example is valid—the actual argument is (trivially) distributed on the active
43	processor set. The second example is invalid—for example, element X(51) is stored on P(3),

which is not in the active processor set for the call. The second example would be valid if the ON directive specified P(3:4) or HOME(X(11:20)), both of which map to the same processor set.

Calls to EXTRINSIC subprograms also deserve mention. The "standard" HPF 2.0 description of calling an EXTRINSIC (Section 6) says in part:

1 A call to an extrinsic procedure must be semantically equivalent to a call of 2 an ordinary HPF procedure. Thus a call to an extrinsic procedure must behave as if the following actions occur... 4 1. Exactly the same set of processors are visible to the HPF environment 5 before and after the subprogram call. 6 7 This constraint is changed to read 8 9 • Exactly the same set of *active* processors are available to the HPF environment before 10 and after the subprogram call. 11 • Exactly the same *universal* processor set is visible to the HPF environment before 12and after the subprogram call. 13 14 The intent is the same as in the original language design. Processors where data is stored can 15 neither appear not disappear; nor may the set of processors executing the program change 16 without notice to the program. Similarly, some extrinsic kinds specify "all processors must 17be synchronized" or "execution of a local procedure on each processor"; such language is 18 understood to mean "all active processors must be synchronized" or "execution of a local 19 procedure on each *active* processor." 20  $^{21}$ Rationale. This gives the combination of EXTRINSIC procedures and ON directives a 22 fork-join model of parallelism, which seems to be both natural and semantically clean. 23 (End of rationale.)  $^{24}$  $^{25}$ If a procedure uses alternate return, then the target of the return must be have the 26 same active processor set as the CALL statement. In effect, this means that labels passed as 27 arguments must refer to statements in the same ON block as the CALL statement. 28 29 Rationale. This constraint is similar to the prohibition against jumping out of an ON 30 block, and has the same justification. (*End of rationale.*) 31 32 Explicit use of CALLs in ON directives is often associated with task parallelism. Several 33 examples can be found in Section 9.4. The following example illustrates how processors can 34 be used for a one-dimensional domain decomposition algorithm: 35 !HPF\$ PROCESSORS PROCS(NP) 36 !HPF\$ DISTRIBUTE X(BLOCK) ONTO PROCS 37 38 ! Compute ILO(IP) = lower bound on PROCS(IP) 39 ! Compute IHI(IP) = upper bound on PROCS(IP) 40 DONE = .FALSE. 41DO WHILE (.NOT. DONE) 42 !HPF\$ INDEPENDENT 43 DO IP = 1, NP 44 !HPF\$ ON (PROCS(IP)) 45 CALL SOLVE\_SUBDOMAIN( IP, X(ILO(IP):IHI(IP)) ) 46 END DO 47

!HPF\$ ON HOME(X) BEGIN

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CALL SOLVE\_BOUNDARIES( X, ILO(1:NP), IHI(1:NP) ) DONE = CONVERGENCE\_TEST( X, ILO(1:NP), IHI(1:NP) ) !HPF\$ END ON END DO

The algorithm divides the entire computational domain (array X) into NP subdomains, one 6 for each processor. The INDEPENDENT IP loop performs a computation on each subdomain's 7 interior. The ON directive tells the compiler which processors to use in executing these (con-8 ceptually) parallel operations. This can increase data locality substantially, particularly 9 if the compiler could not otherwise analyze the data access patterns in SOLVE\_SUBDOMAIN. 10 The subroutine SOLVE\_SUBDOMAIN can use a transcriptive or descriptive mapping for its 11 array argument, placing it on a single processor. In the next phase, the processors collab-12orate to update the boundaries of the subdomains and test for convergence. Subroutines 13 SOLVE\_BOUNDARIES and CONVERGENCE\_TEST may well have their own loops similar to the IP 14 loop, with similar **RESIDENT** clauses. Note that only the lower and upper bounds of each sub-15domain is recorded; this allows different processors to process different-sized subdomains. 16However, each subdomain must "fit" into one processor's section of the X array. 17

Advice to implementors. The IP loop above is likely to be a common idiom among programmers doing block-structured codes. In general, it can be implemented by inverting the HOME clause as was done above. In the one-to-one case shown here (probably very popular with programmers), it can be implemented by assigning the processor id to the loop index variable and testing the range of the loop (once). (End of advice to implementors.)

*Rationale.* Some compilers will propagate the ON information from the caller to the callee at compile time, and some at run time. Repeating the ON clause in the caller and callee will tend to give the compiler better information, resulting in better generated code. (End of rationale.)

#### 9.3The **RESIDENT** Clause, Directive, and Construct

32 The purpose of the **RESIDENT** clause is to promise that data accessed by a computation 33 are mapped to active processors. That is, **RESIDENT** asserts that certain references (or all 34 references) in its scope are stored on the active processor set. The compiler can use this 35 information to avoid generating communication or to simplify array address calculations. 36 Note that whether a given data element is resident depends on two facts:

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- Where the data is stored (i.e., **DISTRIBUTE** and **ALIGN** attributes for the object)
- Where the computation is executed (i.e., its active set, as specified by an ON directive)

For these reasons, the **RESIDENT** clause is added to the **ON** directive, which is usually the 41 earliest point in the program where the needed facts might be available. The **RESIDENT** 42 clause can also appear as a stand-alone directive; this is useful when the locality information 43 is not true for an entire ON region. Note that changing the ON directive may invalidate some 44 **RESIDENT** clauses, or may make more **RESIDENT** clauses true. 45

46 47	H910	resident-clause	is	RESIDENT resident-stuff
48	H911	resident-stuff	is	[ ( res-object-list ) ]

H912	resident- $directive$	$\mathbf{is}$	RESIDENT resident-stuff	1
H913	resident-construct	is		2
			directive- $origin$ $block$ - $resident$ - $directive$	3
			block	4
			directive-origin end-resident-directive	5
			airective-origin ena-resident-airective	6
H914	$block\-resident\-directive$	is	RESIDENT resident-stuff BEGIN	7
H915	$end\-resident\-directive$	is	END RESIDENT	8
HOLC	1	•	1	9
H910	res- $object$	15	object	10

A resident-directive is a kind of executable-directive. Similarly, a resident-construct is a kind of executable-construct.

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Any top-level objects in the **RESIDENT** clause must be explicitly mapped. Similarly, the **RESIDENT** clause must appear at a point in the program with a declared active processor set (i.e., inside an ON block). Otherwise, the assertion (see below) is a statement about how the compiler works, not about the program.

Advice to implementors. RESIDENT removes the need for inactive processors to participate in communication into/out of an ON clause. (End of advice to implementors.)

The RESIDENT directive is an assertion to the compiler that certain object references made within the ON are stored on the active processors if the computation is performed by the specified active processor set. The scope of the assertion is the next Fortran statement if the *resident-directive* form is used and the enclosed *block* of code if the *resident-construct* form is used. If RESIDENT appears as a clause in an ON directive, then the ON and RESIDENT apply to the same statements.

**RESIDENT**(var) means the *lexical expression var*, when encountered in the execution 27of statements in the scope of the **RESIDENT** directive, accesses only data resident on the set 28 of active processors. (That is, the set of processors named by the innermost available ON 29 directive.) If var is accessed by the statement (e.g., it appears on the right-hand side of an 30 assignment statement, or in the evaluation of a conditional expression), then at least one 31 copy of the object and any subobject of the object must be mapped to the active processor 32 set. If var is assigned to by the statement (e.g., it appears on the left hand side of an 33 34 assignment statement, or in the variable list of a **READ** statement), or in any other context that may cause its value to change) then all copies of the variable and all subobjects of 35 the variable must reside in the active processor set. The application of **RESIDENT** to **CALL** 36 statements and function invocations introduces some complexity into this interpretation; 37 38 these issues will be dealt with in Section 9.3.2.

Note that **RESIDENT** is always an assertion relative to the surrounding **ON** directive. Therefore, if the compiler does not implement the **ON** directive then it must be careful in interpreting **RESIDENT**. Similarly, if the compiler overrules the programmer-specified **ALIGN** and **DISTRIBUTE** directives, then it may not rely on the **RESIDENT** clause in general.

Finally, NEW variables are not considered by any nested **RESIDENT** directives, as detailed below.

Rationale. The different treatment of variable reads and writes is due to the implementation requirements. If a variable's value is read (but not written), then it can be taken from any consistent copy. Therefore, **RESIDENT** only asserts that one of those copies is available. Conversely, all copies of a replicated variable must be consistent, so **RESIDENT** asserts that all copies are available if it is updated.

The RESIDENT assertion is always relative to the declared data mappings and ON clauses because both pieces of information are necessary to determine the locality of data references. Data mapping determines where the data is stored, while ON clauses determine where they are used; in essence they determine the endpoints of a data path. RESIDENT itself says that the path length is very short; obviously, one cannot measure a path without knowing both endpoints. (*End of rationale.*)

Consider the following:

 $^{24}$ 

!HPF\$ ON HOME(Z(I)), RESIDENT(X,Y,RECORD(I))
X(I) = Y(I+1) + RECORD(I)%FIELD1 + RECORD(I+1)%FIELD2

- The following facts are asserted by the directive:
- Z(I) would be local if it appeared, due to its use in the HOME directive.

• All copies of X(I) are stored on processors that also store a copy of Z(I), due to the RESIDENT clause. This may be true because X and Z have the same mapping, or because Z is replicated over a set of processors that contains the set of processors that store X(I).

- At least one copy of Y(I+1) is on the same processor as Z(I), due to the RESIDENT clause. This may be true because Y is replicated on all processors, because Z(I) and Y(I+1) are the only elements of their arrays that are mapped to the same processor, or because the directive
  - !HPF\$ ALIGN Y(J) WITH Z(J-1)

appears elsewhere in the program. (Other situations also make the **RESIDENT** assertion true.)

• At least one copy of all subobjects of RECORD(I) is mapped on the same processor as Z(I). In particular, the reference RECORD(I)%FIELD1 (i.e., a subobject consisting of one component) can be accessed locally. The situations in which this is true are similar to those for X(I). No information is available in this example regarding RECORD(I+1)%FIELD2.

If there is no *res-object-list*, then *all* references to *all* variables referenced during ex-ecution of the RESIDENT directive's body except those declared NEW in a surrounding ON directive are local in the sense described above. That is, for every usage of any variable's value, at least one copy of the variable will be mapped to the ON processor set. Likewise, for every operation that assigns to a variable, all copies of that variable are mapped to the ON processor set. References and assignments to NEW variables are always considered resident. If there are no function or subroutine invocations, this is syntactic sugar for list-ing all variable references within the directive's scope. (See Section 9.3.2 for a discussion of **RESIDENT** clauses applied to subprogram calls.) It might well have been named the ALL\_RESIDENT clause; the present form, however, does not add yet another keyword to the directive sublanguage. 

#### 194 SECTION 9. APPROVED EXTENSIONS FOR DATA AND TASK PARALLELISM

Note that if the active set includes more than one processor, then **RESIDENT** only 1 asserts that the variables are stored on one of the processors. For example, if a statement 2 is executed on a section of the processors arrangement, then communication within that 3 section may be needed for some variables in the **RESIDENT** clause. Communication with 4 processors outside of the section will not be needed for those variables, however. 5

Rationale. The alternative to this interpretation would be that any variable named in the **RESIDENT** clause would be local to all processors, i.e., replicated. While that certainly allows more extensive optimizations, it is a less common case. In addition, it does not seem to capture the intent of **ON** directives applied to **CALL** statements or compound statements. For example,

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would presumably call FOO on a row of the processors arrangement, passing elements of X in place. This is what the current definition does; if **RESIDENT** meant "resident on every processor", the call would force X to be replicated. (*End of rationale.*)

It is not correct to assert that an unmapped object, including of necessity any sequential object, is mapped exclusively to the active processors, unless the programmer knows that all of the processors are active. Thus, when a proper subset of processors is active, no such object can occur in a *res-object-list* or in the scope of a **RESIDENT** directive with no *res-object-list*.

27The RESIDENT directive is similar to the INDEPENDENT directive, in that if it is cor-28 rect it does not change the meaning of the program. If the **RESIDENT** clause is incorrect, 29 the program is not standard-conforming (and is thus undefined). Like the INDEPENDENT 30 directive, the compiler may use the information in the **RESIDENT** clause, or ignore it if it is 31 insufficient for the compiler's purposes. If the compiler can detect that the **RESIDENT** clause 32 is incorrect (i.e., that a **RESIDENT** variable is definitely nonlocal), it is justified in producing 33 a warning. Unlike the INDEPENDENT directive, however, the truth of the RESIDENT clause 34 depends on the mapping of computations (specified with the ON clause) and the mapping 35 of data (specified with DISTRIBUTE and ALIGN clauses); if the compiler overrides either of 36 these, then it may not be able to use information in the **RESIDENT** directive.

*Rationale.* Knowing that a reference is local is valuable information for the optimizer. It is in keeping with the spirit of HPF to phrase this as an assertion of fact, which the compiler can use as it pleases. Expressing it as advice to the compiler seems to have disadvantages. Some possible ways this advice could be phrased, and the counter-arguments, are

"Don't generate communication for this reference" has great potential for changing the meaning of the program. Some programmers want this capability, but it violates the "correct directives should not change the meaning of a program"
principle of HPF. Also, once communication is "turned off" for a reference, it's not clear how to turn it back on.

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• "Generate communication for this reference" is not a useful directive, since the
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                compiler has to do this anyway.
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             • "Generate communication for this reference, and place it here" is useful, since
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                it can override the default placement by the compiler. It still has potential
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                for changing program meaning. It also has the potential to create programs as
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                complex as message-passing, as programmers try to move communication out of
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                loops.
8
           (End of rationale.)
9
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             Examples of RESIDENT Clauses
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     9.3.1
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     As in Section 9.2.3, our aim here is to suggest idioms that may be generally useful to
13
     programmers. We begin by expanding on two earlier examples.
14
          RESIDENT is most useful in cases where the compiler cannot detect access patterns.
15
     Often this arises due to the use of indirection, as in the following examples:
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17
              REAL X(N), Y(N)
18
              INTEGER IX1(M), IX2(M)
19
              !HPF$ PROCESSORS P(NP)
20
              !HPF$ DISTRIBUTE (BLOCK) ONTO P :: X, Y
21
               !HPF$ DISTRIBUTE (BLOCK) ONTO P :: IX, IY
22
23
              !HPF$ INDEPENDENT
^{24}
              DO I = 1, N
25
                 !HPF$ ON HOME( X(I) ), RESIDENT( Y(IX(I)) )
26
                 X(I) = Y(IX(I)) - Y(IY(I))
27
              END DO
28
29
              !HPF$ INDEPENDENT
30
              DO J = 1, N
31
                 !HPF$ ON HOME( IX(J) ), RESIDENT( Y )
32
                 X(J) = Y(IX(J)) - Y(IY(J))
33
              END DO
34
35
              !HPF$ INDEPENDENT
36
              DO K = 1, N
37
                 !HPF$ ON HOME( X(IX(K)) ), RESIDENT( X(K) )
38
                 X(K) = A(IX(K)) - A(IA(K))
39
              END DO
40
          As we saw in Section 9.2.3, X(I) is always local in the I loop and IX(I) and IY(I)
41
```

rarely are. The RESIDENT directive above ensures that Y(IX(I)) is local as well. This would
most likely be due to some property of the algorithm that generated IX (for example, if
IX(I)=I for all I). Note that it is possible for an expression (e.g., Y(IX(I))) to be local
even though one of its subexpressions (IX(I)) is not.

The directive gives no information about Y(IY(I)); it might have only one nonlocal value, or all its values might be nonlocal. (We assume that if there were no nonlocal values, then the RESIDENT clause would include Y(IY(I)) as well.) If there are many local elements referenced by this expression, and they can easily be separated from the local elements, then it may be worthwhile to restructure the loop to make this clear to the compiler. For example, suppose that we knew that only the "first" and "last" X elements on each processor were nonlocal. The loop could then be split thus:

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!HPF$ INDEPENDENT, NEW(LOCALI)
DO I = 1, N
  !HPF$ ON HOME( X(I) ), RESIDENT( Y(IX(I)), Y(IY(I)) ) BEGIN
    LOCALI = MOD(I, N/NP)
    IF (LOCALI\=1 .AND. LOCALI\=0) THEN
      X(I) = Y(IX(I)) - Y(IY(I))
    END IF
  !HPF$ END ON
END DO
!HPF$ INDEPENDENT, NEW(LOCALI)
DO I = 1, NP
  !HPF$ ON (P(I)), RESIDENT( X(LOCALI), Y(IX(LOCALI)) ) BEGIN
    LOCALI = (I-1)*N/NP
    X(LOCALI) = Y(IX(LOCALI)) - Y(IY(LOCALI))
    LOCALI = I * N / NP
    X(LOCALI) = Y(IX(LOCALI)) - Y(IY(LOCALI))
  !HPF$ END ON
END DO
```

The first loop (inefficiently) processes the local elements of Y(IY(I)), while the second (more efficiently) handles the rest. On most machines, it would pay to rewrite both loops to avoid the division operations, for example by creating a logical mask *a priori*.

In the J loop, the **RESIDENT** clause asserts that all accessed elements of Y are local. In this case, that is equivalent to the assertion

!HPF\$ RESIDENT( Y(IX(J)), Y(IY(J)) )

Although the original **RESIDENT** clause only referred to the lexical expression Y, the compiler can infer that the subexpressions are also local. This is because it is impossible for a subobject to be on a different processor than the "parent" object is. This observation can often shorten **RESIDENT** clauses substantially.

In the K loop, the following references are local:

- Y(IX(K)), because Y has the same distribution as X and X(IX(K)) is local (due to the ON clause).
- X(K), because of the RESIDENT clause.

Note that a reference may be local even if it does not appear explicitly in a **RESIDENT** clause. One mark of a good compiler will be that it aggressively identifies these elements.

Because it is an assertion of act, the compiler can draw many inferences from a single **RESIDENT** clause. For example, consider the following case:

!HPF\$ ALIGN Y(I) WITH X(I)
!HPF\$ ALIGN Z(J) WITH X(J+1)

1 !HPF\$ ON HOME( X(K) ), RESIDENT( X(INDX(K)) ) 2 з X(K) = X(INDX(K)) + Y(INDX(K)) + Z(INDX(K))4 The compiler is justified in making the following assumptions in compiling the assignment 5 statement (assuming it honors both the ALIGN directives and the ON directive): 6 7 • X(K) requires no communication (because of the HOME clause) 8 9 • X(INDX(K)) requires no communication (because of the RESIDENT clause) 10  $1\,1$ • Y(INDX(K)) requires no communication (because Y has the same mapping as X, and 12INDX(K) clearly cannot change values between its use in the two references X(INDX(K)) and Y(INDX(K))) 13 14 The compiler cannot make any assumption about INDX(K) or Z(INDX(K)) from the above 15code. There is no indication how INDX is mapped relative to X, so the ON directive gives 16 no guidance. Note that the fact that an expression (here, X(INDX(K))) is local does not 17imply that its subexpressions (here, INDX(K)) are also local. Similarly, Z's mapping does not 18 determine if Z(INDX(K)) would be local; it indicates that Z(INDX(K)-1) is local, but that 19 isn't a great help. If the compiler has additional information (for example, X is distributed 20 by BLOCK and INDX(K) is not near a block boundary), it might be able to make additional 21 deductions. 22 23 One mark of a good compiler will be that it aggressively Advice to implementors. 24 propagates **RESIDENT** assertions. This is likely to significantly reduce communication 25costs. Note the cases under "Advice to users" below. (End of advice to implementors.) 26 27 Advice to users. One can expect compilers to differ in how aggressive they are 28 in drawing these deductions. Higher-quality compilers will be able to identify more 29 references as local, and use this information to eliminate data movement. All compilers 30 should recognize that if an element of one array is local, then the same element of 31 any other arrays with the same static mapping (i.e., arrays aligned together, or with 32 the same **DISTRIBUTE** pattern and array size) will also be local. That is, any compiler 33 should recognize Y(INDX(K)) in the above example as local. Dynamically changing 34 array mappings (i.e., REALIGN and REDISTRIBUTE) will tend to limit such information 35 and information propagation. Also, assignments that might change subexpressions 36 (for example, an assignment to K or any element of INDX in the above example) will 37 force the compiler to be conservative in its deductions. (End of advice to users.) 38 39 40 **RESIDENT** Directives Applied to Procedure Reference 9.3.241If a **RESIDENT** directive applies to procedure reference, then the assertion is more subtle. 42 43 • If a *res-object-list* appears in the **RESIDENT** directive, then no assertion is made about 44 behavior within the called procedure. For example, consider the statements: 45 46 !HPF\$ RESIDENT( A(I), B ) 47

A(I) = F(A(I), B(LO:HI))

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The directive declares all variable referces in the statement (including the actual arguments) to be local to the current ON processor set. However, the execution of F itself could access elements of arrays named A and B stored on arbitrary processors.

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*Rationale.* Propagating assertions about the behavior of lexical entities is difficult to define consistently and usefully. For example, consider the following function called from the code fragment above:

REAL FUNCTION F( X, Y ) USE MODULE\_DEFINING\_A REAL X, Y(:), B(I) !HPF\$ INHERIT Y !HPF\$ ALIGN B(:) WITH Y(:) INTEGER I Z = 0.0 DO I = 1, SIZE(Y)

Z = Z + A(I) \* X + B(I) \* Y(I)END DO F = Z END FUNCTION

Assume A is defined as a distributed, global array in module MODULE\_DEFINING\_A. 22 What should the **RESIDENT** clause mean regarding operations in F? The expres-23 sion A(I) in the **RESIDENT** directive might reasonably mean references only to  $^{24}$ the array A that is visible in the caller, or it might mean references to any array  $^{25}$ named A. Note that the A in the caller may be local, the same global array as the 26 A in F (if the caller used MODULE\_DEFINING\_A), or a different global array (if the 27caller used a different module). Perhaps a limiting case is array B. The array B 28 in function F is local, and thus different from the caller; however, because of the 29 restrictions on ON clauses it is certain that the local B will be mapped to the ON 30 processor set. Thus, the **RESIDENT** assertion is trivially true. To further confuse 31 matters, **RESIDENT** variables might seem to apply to dummy arguments that be-32 come associated with those variables. Unfortunately, this implies that the lexical 33 expression B in the caller refers to the lexical expression Y in F, which stretches 34 the definition of "lexical" beyond the breaking point. For all these reasons, it 35 was decided to limit the meaning of named variables in **RESIDENT** clauses to the 36 lexical scope of the directive. (End of rationale.) 37

• If the **RESIDENT** directive does not contain a *res-object-list*, then the directive asserts that all references in the caller *and the called procedures* are local as defined above. For example, consider the statements:

!HPF\$ RESIDENT
A(I) = F( A(I), B(LO:HI) )

The directive declares all variable referces in the statement (including the actual arguments) to be local to the current ON processor set, and that F itself does not reference or update any nonlocal variables.

Rationale. The **RESIDENT** assertion is always true for data local to the called 1 procedure. This is true because the called procedure must use a declarative 2 3 ON clause, which in turn limits the set of processors that can store any local explicitly mapped variables. The above definition extends this assertion to all 4 global explicitly mapped data, producing a very powerful directive. This is sim-5 ilar to the meaning of INDEPENDENT, in that it also makes an assertion about 6 variable accesses in any called procedure in the loop. An alternative semantics 7 for **RESIDENT** would have been to avoid propagating the assertion interprocedu-8 rally (i.e., treat both the variable-list version and the no-list version the same). 9 However, this would not provide enough information to optimize code on certain 10 machines. In particular, it would have made task parallelism guite difficult on 11 message-passing machines. (End of rationale.) 1213 Advice to implementors. **RESIDENT** without a variable list guarantees that no one-14 sided communication outside of the ON processor set will be generated by the callee. 15Such a procedure can be called only on the "active" processors, unless the runtime 16system has additional constraints (for example, if the runtime system requires all 17processors to participate in collective communications). 18 19 The other forms of **RESIDENT** provide information that could be propagated inter-20 procedurally. For example, if the actual argument to a subprogram is asserted to be 21 **RESIDENT** and is passed transcriptively, then anything that is aligned to it in the callee 22 will also be **RESIDENT**. If the information is not propagated, the only result will be 23 less optimization. (End of advice to implementors.) 24 25Advice to users. Although the **RESIDENT** assertion applies interprocedurally, it is by 26 no means certain that all compilers will make use of this information. In particular, 27 separate compilation limits the propagation that can take place. It is therefore good 28 practice to include a RESIDENT clause both in the caller's ON directive and in the 29 callee. (This assumes that the assertion is true, of course!) This ensures that the 30 compiler has the **RESIDENT** information available when it is compiling both ends of 31 the procedure call. This is especially useful for **RESIDENT** clauses without a variable 32 list; knowing that all data accessed is local allows many optimizations that are not 33 otherwise possible. (End of advice to users.) 34 35 Locality information is particularly critical interprocedurally. Here, the **RESIDENT** di-36 rective without a res-object-list can be used to good advantage. Consider the following 37 extension of the block-structured example from Section 9.2.4: 38 !HPF\$ PROCESSORS PROCS(NP) 39 !HPF\$ DISTRIBUTE X(BLOCK) ONTO PROCS 40 4142 ! Compute ILO(IP) = lower bound on PROCS(IP) ! Compute IHI(IP) = upper bound on PROCS(IP) 43 DONE = .FALSE. 44 DO WHILE (.NOT. DONE) 45 !HPF\$ INDEPENDENT 46

DO IP = 1, NP

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!HPF\$ ON (PROCS(IP)), RESIDENT

```
CALL SOLVE_SUBDOMAIN( IP, X(ILO(IP):IHI(IP)) )

END DO

!HPF$ ON HOME(X) BEGIN

CALL SOLVE_BOUNDARIES( X, ILO(1:NP), IHI(1:NP) )

!HPF$ RESIDENT

DONE = CONVERGENCE_TEST( X, ILO(1:NP), IHI(1:NP) )

!HPF$ END ON

END DO
```

Recall that the INDEPENDENT IP loop performs a computation on each subdomain's interior, 10 where a subdomain is mapped to a particular processor. The first **RESIDENT** clause addition-11 ally informs the compiler that no subdomain uses data from another processor. Without 12this information, the compiler would have to assume a worst-case scenario in which each 13 subdomain performed its updates based on non-local read-only data. Any nonlocal data 14 could not be written by another processor without violating the tt INDEPENDENT direc-15tive; however, if the data were not updated (for example, a large lookup table) it could be 16stored remotely. Particularly on nonshared-memory machines, access to this remote data 17would be difficult. The **RESIDENT** clause ensures that this possibility need not be considered. 18 All data required by SOLVE\_SUBDOMAIN is stored locally. The second RESIDENT clause asserts 19 that all data for CONVERGENCE\_TEST is stored on the same processors that store X. The same 20 cannot be said for SOLVE\_BOUNDARIES, which does not fall in the scope of the RESIDENT  $^{21}$ directive. For example, there might be a processors arrangement other than PROCS with 22 necessary data. Accessing this data might well cause a bottleneck in the computation as 23 described above.  $^{24}$ 

Again, note the usefulness of **RESIDENT** clauses in giving the compiler information. Few compilers would be able to unravel nontrivial assignments to ILO and IHI, and no current compiler would even attempt to understand the comments in the above code fragment. End of advice to programmers.

### 9.4 The TASK\_REGION Construct

Task parallelism is expressed in HPF by mapping data objects onto subsets of processors and adding assertions that allow concurrent execution of different code blocks on different processor subsets. A data object is mapped to a processor subset by distribution onto a subsection of a processors arrangement. Execution on a subset of processors is specified by using an ON directive. This section introduces a TASK\_REGION directive that allows the user to specify that disjoint processor subsets can execute blocks of code concurrently.

A TASK\_REGION directive is used to assert that a block of code satisfies the following set of constraints. All lexically outermost ON blocks inside a task region must have a RESIDENT attribute implying that all data accessed inside them is mapped to the corresponding active processor subset. Further, the code inside two such ON blocks must not have interfering I/O. Under these constraints, two such ON blocks can safely execute concurrently if they execute on disjoint processor subsets.

### 9.4.1 Syntax of the TASK\_REGION Construct

A task region is a single entry region delimited by two structured comments:

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1	H917	task-region-construct	$\mathbf{is}$	
2				$directive\mbox{-}origin\ block\mbox{-}task\mbox{-}region\mbox{-}directive$
3				block
4				directive- $origin$ $end$ - $task$ - $region$ - $directive$
5 6	H918	$block\-task\-region\-directive$	is	TASK_REGION
7	H919	$end\-task\-region\-directive$	is	END TASK_REGION

A task-region-construct is a kind of executable-construct.

There must not be a transfer of control from outside the *task-region-construct* to inside the *task-region-construct*. Transfer of control out of the *task-region-construct* is allowed provided that the transfer does not originate inside an ON block. (The reason for this will be apparent later.)

9.4.2 Semantics of the TASK\_REGION Construct

<sup>16</sup> We will refer to a block of code enclosed by a TASK\_REGION ... END TASK\_REGION pair as <sup>17</sup> a *task region*. The TASK\_REGION directives are a way for the programmer to assert that a <sup>18</sup> section of code satisfies a set of conditions. The compiler is expected to use these assertions <sup>19</sup> to generate task-parallel code.

A task region can contain blocks of code that are directed to execute ON processor subsets. All other code executes on a subset that contains all active processors. Every ON block at the outermost nesting level (i.e., not inside another ON block or another task region) inside a task region is defined as a *lexical task*. Every execution instance of a lexical task is defined as an *execution task* and will also be referred to as just *task* when the distinction is clear from the context. An execution task is associated with a set of *active processors* discussed earlier in this chapter.

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The following restrictions must hold inside a task region:

• Every ON block corresponding to a lexical task must have the **RESIDENT** attribute. This means that, for reading a variable inside an execution task, the corresponding active processors must own at least one copy of the variable, and for writing, they must own all copies of that variable.

- An I/O operation inside an execution task may interfere with an I/O operation inside another execution task if and only if the two tasks execute on identical subsets of processors. Note that two execution tasks can be instances of the same or different lexical tasks. In general, two I/O operations interfere if they access the same file or unit. The conditions for interference of I/O operations are detailed in Section 5.1 in the context of the INDEPENDENT directive.
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## 9.4.3 Execution Model and Usage

A task region does not introduce a fundamentally new execution model. However, the assertions implicit in a task region imply that only the specified active processors of an execution task need to participate in its execution, and that other processors can skip its execution. A processor executing a task region participates in the execution of all tasks executing on a processor subset that it belongs to, and does not participate in the execution of tasks executing on processor subsets that it does not belong to. Code outside lexical tasks is executed as normal data parallel code by all active processors of the task region. The access restrictions for a task region guarantee that the results obtained by this execution paradigm will be consistent with pure data parallel execution of a task region.

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A task region presents a simple yet powerful model to write integrated task and data parallel programs. We illustrate three basic computation structures in which task parallelism can be effectively exploited with this model.

- 1. *Parallel sections:* A task region can be used to divide the available processors into disjoint sets for performing independent computations, simulating what is often referred to as *parallel sections*. This form of task parallelism is relatively straightforward and useful in many application scenarios, an example being multiblock applications. The task region simply contains a sequence of **RESIDENT ON** blocks on disjoint processor subsets. Note that the division of processors among subsets can be dynamic, that is, it can be in terms of other variables computed during execution.
- 2. Nested parallelism: Task regions can be nested, and in particular, a subroutine call made from an execution task can further subdivide the active processors using another task region directive. This allows the exploitation of nested parallelism. An example is the implementation of dynamic tree structured divide and conquer computations. As a specific example, quicksort can be implemented by recursively partitioning the input array of keys around a pivot, and assigning proportionate number of processors 20 to the two new arrays obtained as a result of partitioning.  $^{21}$
- 3. Data parallel pipelines: Task regions can be used to implement pipelined data parallel computations. We will illustrate this with a 2 dimensional fast Fourier transform (2D FFT) computation. The first stage of a 2D FFT reads a two dimensional matrix and performs a 1 dimensional FFT on each row of the matrix. The second stage performs a 1 dimensional FFT on each column of the matrix and generates the final output. In a pipelined data parallel implementation of this form of 2D FFT, the two stages are mapped on to disjoint subsets of processors. Task and data parallel code for a 2D FFT, along with a brief description, is included in Section 9.4.5.

#### Implementation 9.4.4

A task region is simply an assertion about a code block and the exploitation of task parallelism is, at least partially, dependent on the compilation scheme. While the specifics of how task parallelism is exploited will be strongly dependent on the parallel system architecture, the compiler, and the underlying communication model, we will point out some important considerations and illustrate task parallel code generation with an example. We primarily address distributed memory machines using a message passing communication and synchronization model, but will point out some of the important issues relating to shared memory implementations.

#### Localized computation and communication 9.4.4.1

It is of central importance that computation and communication inside an executing task should not involve any processors other than those directed to execute the task in the relevant ON clause.

On entry to a lexical task, the compiler has to insert checks so that the inactive pro-47 cessors jump to the code following the task. Since an execution task cannot access data 48 outside of the active processor set, no communication needs to be generated between the relevant active processors and other processors. In a message passing model, a communication generation algorithm that only generates necessary messages will naturally achieve the desired results. However, some communication schemes can involve generation of empty messages between processors that do not communicate and it is important to ensure that empty messages are not generated between active processors of an executing task and other processors.

A communication model that uses barriers for synchronization (in shared or distributed 8 memory machines) must ensure that all barriers inside an executing task are subset barriers 9 that only span the active processors. An implementation may also need to include a subset 10 barrier, on entry to and on exit from, an executing task for consistency of data accesses  $1\,1$ inside and outside an executing task. In general, the compilation framework has to ensure 12the consistency of data accesses inside and and outside an executing task and this can 13 be done in the context of virtually any synchronization scheme in a shared or distributed 14 memory environment. 15

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## 9.4.4.2 Replicated computations

All computations exclusively involving replicated variables should be replicated on all executing processors. A simple alternative is that one processor performs the computation and broadcasts the results to all processors. While such replication is generally profitable in HPF anyway, it has additional importance in a task region since the communication generated by a broadcast can cause additional synchronization that may interfere with task parallelism.

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# 9.4.4.3 Implications for I/O

In some parallel system implementations, I/O is performed through a single processor of the system. Task parallelism in the presence of I/O assumes that all processors can perform I/O independently and this paradigm has to be supported, although it is not necessary that each processor be able to physically perform all I/O operations independently. One simple solution is to have a single designated I/O processor that performs all I/O but is not considered an executing processor and hence does not have any execution related dependences.

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# 9.4.4.4 SPMD or MIMD code generation

Another issue for the compiler is whether or not the same code image should execute on all processors. Since different processor groups may need different variables, a naive SPMD implementation is likely to be wasteful of memory since it must allocate all variables on all processors. This can be addressed by dynamic memory allocation, but at the cost of added complexity. Using different code images for different processor subsets is another solution that also leads to significant added complexity.

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## 9.4.5 Example: 2-D FFT

This section shows the use of task parallelism to build a pipelined data-parallel 2dimensional FFT and illustrates the compilation of task parallelism by showing SPMD code generated from the HPF program.

The basic sequential 2DFFT code is as follows: REAL, DIMENSION(n,n) :: a1, a2 DO WHILE(.true.) READ (unit = 1, end = 100) a1 CALL rowffts(a1) a2 = a1 CALL colffts(a2) WRITE (unit = 2) a2 10 CYCLE 11100 CONTINUE 12EXIT 13 END DO 14

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To write a pipelined task and data parallel 2D FFT in HPF, the code is slightly modified and several HPF directives are added. First, variables a1 and a2 are distributed onto disjoint subsets of processors, and then a task region is used to create two lexical tasks to perform rowffts and colffts on different subsets of processors. The assignment a2 = a1 in the task region specifies the transfer of data between the tasks. A new variable done1 is introduced to store the termination condition. The modified code is as follows:

```
REAL, DIMENSION(n,n) :: a1,a2
                                                                                     22
        LOGICAL done1
                                                                                     23
        PROCESSORS procs(8)
!HPF$
                                                                                     ^{24}
                                                                                     ^{25}
!HPF$
        DISTRIBUTE a1(block,*) ONTO procs(1:4)
                                                                                     26
        DISTRIBUTE a2(*,block) ONTO procs(5:8)
!HPF$
                                                                                     27
                                                                                     28
        TEMPLATE, DIMENSION(4), DISTRIBUTE(BLOCK) ONTO procs(1:4) :: td1
!HPF$
                                                                                     29
!HPF$
        ALIGN WITH td1(*) :: done1
                                                                                     30
                                                                                     31
!HPF$
        TASK_REGION
                                                                                     32
        done1 = .false.
                                                                                     33
        DO WHILE (.true.)
                                                                                     34
!HPF$
             ON (procs(1:4)) BEGIN, RESIDENT
                                                                                     35
               READ (unit = iu, end=100) a1
                                                                                     36
               CALL rowffts(a1)
                                                                                     37
               GOTO 101
                                                                                     38
    100
               done1 = .true.
                                                                                     39
               CONTINUE
    101
                                                                                     40
!HPF$
             END ON
                                                                                     41
                                                                                     42
             IF (done1) EXIT
                                                                                     43
             a2 = a1
                                                                                     44
                                                                                     45
             ON (procs(5:8)) BEGIN, RESIDENT
!HPF$
                                                                                     46
                CALL colffts(a2)
                                                                                     47
                WRITE(unit = ou) a2
                                                                                     48
```

1	!HPF\$	END ON
2		END DO
3	!HPF\$	END TASK_REGION
4		
5	Finall	y, we show simplified SPMD code generated for each processor. We assume a
6	message p	assing model where sends are asynchronous and nonblocking and receives block
7	until the d	ata is available. We use a simple memory model where variable declarations are
8	identical a	cross all processors even though some variables will be referenced only on subsets
9	of the pro	cessors. A shadow variable done1_copy is created by the compiler to transfer
10	informatio	n from processor subset 1 to processor subset 2 about termination of processing.
11	The code i	is as follows:
12		
13		REAL DIMENSION(n/4,n) :: a1
14		REAL DIMENSION(n,n/4) :: a2
15		LOGICAL done1
16		
17	С	Following are compiler generated variables
18		LOGICAL done1_copy
19		LOGICAL inset1, inset2
20	С	
21	С	Following magic compiler function call is to set the variables
22	С	inset1 and inset2 to .true. for subset 1 and subset 2 processors
23	С	respectively, and .false. otherwise.
24	С	
25		CALL initialize_tasksets(inset1,inset2)
26		
27	С	Code for processor subset 1
28		IF (inset1)
29		done1 = .false.
30		DO WHILE (.true.)
31		
32	С	Read is left unchanged as the code depends on the I/O model
33		READ (unit = 1, $end=100$ ) a1
34		
35		CALL rowffts(a1)
36		GOTO 101
37	100	
38	101	
39		_send(done1,procs(5:8))
40		IF (done1) EXIT
41		_send(a1,proces(5:8))
42		END DO
43		END IF
44		
45	C	Code for processor subset 2
46		IF (inset2)
47		DO WHILE(.true.)
48		_receive(done1_copy,procs(1:4))

```
IF (local_done1) EXIT
_receive(a2,procs(1:4))
CALL colffts(a2)
C Write is left unchanged as the code depends on the I/O model.
        WRITE (unit = 2) a2
        END DO
        END IF
```

\_send and \_receive are communication calls to transfer variables between subsets of processors. Program execution until the end of input is as follows. Subset 1 processors repeatedly read input, compute rowffts, and send the computed output as well as done1 flag, which normally has the value .false., to subset 2 processors. The subset 2 processors receive the flag and the data set, compute colffts and write the results to the output. When the end of input is reached, subset 1 processor set the value flag done1 to .true., send it and terminate execution. Subset 2 processors receive the flag, recognize that the end of input has been reached, and terminate execution.

# Section 10

# Approved Extension for Asynchronous I/O

This section defines a mechanism for performing Asynchronous I/O from an HPF or Fortran program. These are presented as changes to the Fortran 95 proposed draft standard, X3J3/96-007r1. This extension is a subset of the proposed X3J3 Asynchronous I/O extension, paper X3J3/96-158r2. Briefly, this extension allows direct unformatted data transfers to be performed asynchronously with program execution. The WAIT statement can be used to wait for the data transfers to complete. The INQUIRE statement can be used to determine if the data transfers are complete.

To section 9.3.4, rule R905 connect-spec, add

#### or ASYNCHRONOUS

Add a new section after 9.3.4.10, entitled "ASYNCHRONOUS specifier in the OPEN statement", containing the following paragraphs:

If the ASYNCHRONOUS specifier is specified for a unit in an OPEN statement, then READ and WRITE statements for that unit may include the ASYNCHRONOUS specifier in the control information list.

The presence of an ASYNCHRONOUS specifier in a READ or WRITE statement permits (but does not require) a processor to perform the data transfer asynchronously. The WAIT statement is used to wait for or inquire as to the status of asynchronous input/output operations.

 $^{37}_{38}$  To section 9

To section 9.4.1, rule R912 io-control-spec, add

or ID = scalar-default-int-variable
or ASYNCHRONOUS

- and also add the constraints
- 44 Constraint: If either an ASYNCHRONOUS or an ID= specifier is present, then both 45 shall be present.
- 46 Constraint: If an ASYNCHRONOUS specifier is present, the REC= specifier shall
   47 appear, a *format* shall not appear, and a *namelist-group-name* shall
   48 not appear.

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Constraint:	If an ASYNCHRONOUS specifier is present, then no function reference
	may appear in an expression anywhere in the data transfer state-
	ment.

At the end of section 9.4.1, add the following paragraphs:

The addition of the ID= specifier results in the initiation of an asynchronous data transfer. Execution of the data transfer statement shall be eventually followed by execution of a WAIT statement specifying the same ID value that was returned to the ID variable in a data transfer statement. This WAIT statement is called the *matching* WAIT statement. Note that asynchronous data transfer shall be direct and unformatted.

The matching WAIT statement shall be executed in the same instance of the same subprogram in which the asynchronous data transfer statement was executed.

Advice to implementors. The above restriction is to prevent the compiler from performing code motion optimizations across WAIT statements. Any operations involving variables listed in asynchronous input/output lists must be performed after the matching WAIT statement is executed. (End of advice to implementors.)

No ASYNCHRONOUS specifier nor any ID= specifier shall be specified if the io-unit was not opened with the ASYNCHRONOUS specifier.

In section 9.4.1, in the fourth and fifth paragraphs after the constraints, change both instances of "IOSTAT= or a SIZE=" to "IOSTAT=, SIZE=, or an ID=".

Insert the following text at the end of section 9.4.3 before the final paragraph:

For an asynchronous data transfer, errors may occur either during execution of the data transfer statement or during subsequent data transfer. If these errors occur during the data transfer statement and do not result in termination of the program, then they will be detectable using ERR= and IOSTAT= specifiers in the data transfer statement. If these error conditions occur during subsequent data transfer and do not result in termination of the program, then they will be detectable using ERR= and IOSTAT= specifiers in the matching WAIT statement.

In the paragraph at the end of section 9.4.3, change the first occurrence of "execution" to read "execution or subsequent data transfer."

To section 9.4.4, add the following paragraphs:

For asynchronous data transfers steps 1-8 correspond to both the asynchronous data transfer statement and the matching WAIT statement. Steps 4-7 may occur asynchronously with program execution. If an implementation does not support asynchronous data transfers then steps 1-8 may be performed by the asynchronous data transfer statement. The matching WAIT statement shall still be executed, the only effect being to return status information.

Any variable that appears as an *input-item* or *output-item* in an asynchronous data transfer statement, or that is associated with such a variable, shall not be referenced, become defined, or become undefined until the execution of the

## 10.1. THE WAIT STATEMENT

1 2	matching WAIT statement. ciated with such a variable.	However, it is allowed for a pointer to become asso-
3 4 5 6		hronous data transfer operations (READ or WRITE) are RITE operations may use the same unit and record ning WAIT.
7 8	-	ermits left-to-right definition of the $I/O$ list on a READ, ronous. This means that a statement such as
9 10 11	READ(10,ID=IDNU	M,REC=10) I,A(I)
12 13 14	is conforming and has the store to users.)	ame input behavior as a synchronous READ. ( $End \ of \ advice$
15 16 17	, 0	QUIRE statements" to ", INQUIRE, and WAIT statements". Collowing sentence as the last sentence of the paragraph:
18 19 20 21	assigned to the $NEXTREC=$ s	a transfer operations for the specified unit, the value specifier is computed as if all the outstanding data leted, in the order in which they were issued.
22	To section 9.6.1, rule R924 $i$	nquire-spec, add
23 24 25		<pre>or ID = scalar-default-int-variable or PENDING = scalar-default-logical-variable</pre>
26	and also add the constraints	
27 28 29		PENDING= specifiers shall not appear in an INQUIRE ne FILE= specifier is present.
30 31	Constraint: If either an II both shall be p	<b>D</b> = specifier or a <b>PENDING=</b> specifier is present, then present.
32 33 34 <b>1</b>		5.1.22, entitled "ID= and PENDING= specifiers in ontaining the following paragraph:
35 36 37 38 39	in the <b>PENDING=</b> specifier is by the <b>ID=</b> specifier for the	in an INQUIRE statement, then the variable specified assigned the value true if the data transfer identified e specified unit has not yet completed. In all other in the PENDING= specifier is set to false.
40 41 42	10.1 The WAIT Statemer	nt
43	H1001 wait-stmt	is WAIT ( $wait$ -spec-list )
44	H1002 wait-spec	is UNIT = io-unit or ID = scalar-default-int-expr or ERR = label or IOSTAT = label

Constraint: A *wait-spec-list* shall contain exactly one UNIT= specifier, exactly one ID= specifier, and at most one of each of the other specifiers.

The WAIT statement terminates an asynchronous data transfer. The IOSTAT= and ERR= specifiers are optional and are described in sections 9.4.1.4 and 9.4.1.5, respectively.

Advice to implementors. Implementors may choose to implement any or all asynchronous I/O synchronously. This essentially means using the ID= clause on the data transfer statement to record the results of the transfer, then supplying the results to the matching WAIT statement. (End of advice to implementors.)

# Section 11

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# Approved Extensions for HPF Extrinsics

This section builds on Section 6 by defining specific interfaces for HPF with different models of parallelism, LOCAL and SERIAL (Section 11.1), and different languages, HPF (Section 11.3), C (Section 11.4), Fortran (Section 11.5), and Fortran 77 (Section 11.6). Library routines useful in the extrinsic models are defined in Section 11.7. These are defined with Fortran bindings. An implementation may chose to define similar routines for use with C. The intent of the HPF extrinsic mechanism is to generalize. These definitions may serve as a model for other interfaces. Some additional extrinsic interfaces are given after Annex E.

In HPF 1.1, specific extrinsic types HPF, HPF\_LOCAL, and HPF\_SERIAL were defined. There was also a short discussion of F90\_LOCAL. In this more general setting, the HPF keywords are retained for compatibility. As defined in Section 6, a model of HPF refers to the global language, HPF\_LOCAL is the same as LANGUAGE='HPF', MODEL='LOCAL', and HPF\_SERIAL is the same as LANGUAGE='HPF', MODEL='SERIAL'. The F90\_LOCAL extrinsic is now addressed by the LANGUAGE='FORTRAN', MODEL='SERIAL' extrinsic kind.

From the caller's standpoint, an invocation of an extrinsic procedure from a "global" 30 HPF program has the same semantics as an invocation of a regular procedure. The callee 31 may see a different picture. The following sections describe sets of conventions for coding 32 callees in the various extrinsic options. The set of extrinsic options supported by a partic-33 ular HPF compiler is implementation dependent. They are not limited to those described 34 in this chapter. Furthermore, the language processor used to compile the actual extrinsic 35 subprogram need not be an HPF compiler. More specifically, it need not actually be a com-36 piler for the language noted in the LANGUAGE specification, as long as the executing extrinsic 37 code conforms to the conventions defined for the language. We define these interfaces to 38 promote portability and interoperability, but a given implementation and the programmer 39 are free to create other combinations of models and languages. 40

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#### 11.1 Alternative Extrinsic Models: LOCAL and SERIAL

A global HPF program may be thought of as a set of processors cooperating in a loosely synchronous fashion on a single logical thread of program control. Section 6 defines two additional models that may be invoked from global HPF: LOCAL, where the model is singleprocessor "node" code, in which all active processors participate, but with only the data that is mapped to a given physical processor directly accessible, and SERIAL, where the

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model is a single-processor operating alone, with all necessary data aggregated by the caller before the serial subprogram is invoked. As examples of use, the LOCAL model is useful for programs that drop down into code with explicit message passing for data communications, while the SERIAL model may be needed for calls to system libraries or specialized I/O routines.

Both the LOCAL and the SERIAL models can be invoked from a global HPF program, but in general these models may not be mixed. Calling global HPF from local or serial procedures is not allowed. Furthermore, calling a serial procedure from a local one is not allowed. Section 6.3.1 gives more detail about how various models can interact with each other.

Some additional restrictions are placed on all local and serial subprograms invoked from global HPF:

- The subprogram directly invoked by global HPF must not be recursive.
- The subprogram directly invoked by global HPF must not use any alternate return mechanism.

The behavior of I/O statements in local and serial subprograms is implementation dependent.

#### 11.1.1 The LOCAL Model

An extrinsic procedure can be defined as explicit SPMD code by specifying the local pro-23 cedure code that is to execute on each processor. In this section, we describe the contract  $^{24}$ between the caller and an EXTRINSIC (MODEL='LOCAL') callee. It is important not to con- $^{25}$ fuse the extrinsic procedure, which is conceptually a single procedural entity called from 26 the HPF program, with the individual local procedures that are executed on each node, 27one apiece. An *invocation* of an extrinsic procedure results in a separate invocation of a 28 local procedure on each processor. The *execution* of an extrinsic procedure consists of the 29 concurrent execution of a local procedure on each executing processor. Each local proce-30 dure may terminate at any time by executing a **RETURN** statement. However, the extrinsic 31 procedure as a whole terminates only after every local procedure has terminated; in effect, 32 the processors are synchronized before return to a global HPF caller. 33

With the exception of returning from a local procedure to the global caller that initi-34 ated local execution, there is no implicit synchronization required of the locally executing 35 processors. Variables declared in a local procedure are held in local storage, private to each 36 processor. To access data outside the processor requires either preparatory communication 37 to copy data into the processor before running the local code, or explicit communication 38 operations between the separately executing copies of the local procedure. Individual imple-39 mentations may provide implementation-dependent means for communicating, for example, 40 through a message-passing library or a shared-memory mechanism. Such communication 41mechanisms are beyond the scope of this specification. Note, however, that many useful 42 portable algorithms that require only independence of control structure can take advantage 43 of local routines, without requiring a communication facility. 44

The LOCAL model assumes only that nonsequential array axes are mapped independently to axes of a rectangular processor grid, each array axis to at most one processor axis (no "skew" distributions) and no two array axes to the same processor axis. This restriction suffices to ensure that each physical processor contains a subset of array elements that can 48 be locally arranged in a rectangular configuration. (Of course, to compute the global indices
of an element given its local indices, or vice versa, may be quite a tangled computation—but
it will be possible.) In the case of cyclic distributions, multiple sections of the array may
be mapped to the local processors.

It is recommended that if, in any given implementation, an extrinsic type does not obey the conventions described in this section, then its model name or keyword should not contain the word LOCAL.

11.1.1.1 Conventions for Calling LOCAL Subprograms

The default mapping of scalar dummy arguments, of scalar function results, and of sequential arrays is such that the argument is replicated on each physical processor. These mappings may, optionally, be explicit in the interface (except in the case of sequential arrays, which may not be explicitly mapped), but any other explicit mapping is not HPF conforming. Dummy arguments may not be of explicitly mapped derived types or have explicitly mapped structure components.

As in the case of non-extrinsic subprograms, actual arguments may be mapped in any way; if necessary, they are copied automatically to correctly mapped temporaries before invocation of and after return from the extrinsic procedure.

It should be noted that the conventions for calling local subprograms apply only at the interface between the GLOBAL and LOCAL models. The conventions do not propagate to further subprograms called from within the LOCAL model.

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# 11.1.1.2 LOCAL Calling Sequence

Execution of an extrinsic local procedure must be performed as if the actions detailed below occur prior to the invocation of the local procedure on each processor (see Section 6.3.2 for a related list of conditions and for the meaning of *as if*). Any actions thus required are enforced by the compiler of the calling routine, and are not the responsibility of the programmer, nor do they impact the local procedure.

- 1. The processors are synchronized. In other words, all actions that logically precede the call are completed.
- 2. Each actual argument is remapped, if necessary, according to the directives (explicit or implicit) in the declared interface for the extrinsic procedure. Thus, HPF mapping directives appearing in the interface are binding—the compiler must obey these directives in calling local extrinsic procedures. (The reason for this rule is that data mapping is explicitly visible in local routines). Actual arguments corresponding to sequential arrays and scalar dummy arguments are replicated (by broadcasting, for example) in all processors. Scalars of derived types with explicitly mapped structure components or of an explicitly mapped derived type cannot be passed from global HPF to an extrinsic local procedure.
- 3. If a variable accessible to the called routine has a replicated representation, then all copies are updated prior to the call to contain the correct current value according to the sequential semantics of the source program.
- After these actions have occurred, the local procedure is invoked on each processor. The information available to the local invocation is described below in Section 11.1.1.3.

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When control is transferred back to the caller at the conclusion of the extrinsic local procedure, execution must resume as if the following actions have already been performed.

- 1. All processors are synchronized after the call. In other words, global computation procedes as if the execution of every copy of the local routine is completed before execution in the caller is resumed.
- 2. The original distribution of arguments (and of the result of an extrinsic function) is restored, if necessary.

Advice to implementors. An implementation might check, before returning from the local subprogram, to make sure that replicated variables have been updated consistently by the subprogram. However, there is certainly no requirement—perhaps not even any encouragement—to do so. This is the responsibility of the local subprogram, and any checks in the caller are a tradeoff between speed and, for instance, debuggability. (End of advice to implementors.)

#### 11.1.1.3 Information Available to the Local Procedure

The local procedure invoked on each processor is passed a *local argument* for each *global argument* passed by the caller to the (global) extrinsic procedure interface. Each global argument is an HPF array or scalar. The corresponding local argument is the part of the global array stored locally, or a local copy of a scalar argument or sequential array replicated across processors. Note that if the HPF array is replicated, each local procedure receives a copy of the entire actual. An array actual argument passed by an HPF caller is called a *global array*; the subgrid of that global array passed to one copy of a local routine (because it resides in that processor) is called a *local array*.

If the extrinsic procedure is a function, then the local procedure is also a function. Only scalar-valued extrinsic functions are allowed. All local functions must return the same value.

If a global HPF program calls local subprogram A with an actual array argument X, and A receives a portion of array X as dummy argument P, then A may call another local subprogram B and pass P or a section of P as an actual argument to B.

The run-time interface must provide enough information that each local function can discover for each local argument the mapping of the corresponding global argument, translate global indices to local indices, and vice-versa. A specific set of procedures that provide this information is listed in the HPF Local Library Section 11.7.1. The manner in which this information is made available to the local routine depends on the implementation and the programming language used for the local routine.

#### 11.1.2 The SERIAL Model

This section defines a set of conventions for writing code in which an instance of a subprogram executes on only one processor (of which there may be more than one).

If a program unit has extrinsic model SERIAL, an HPF compiler should assume that the subprogram is coded to be executed on a single processor. From the point of view of a global HPF caller, the SERIAL procedure behaves the same as an identically coded HPF procedure would. Differences might only arise in implementation-specific behavior (such as the performance). There is currently no manner in which to specify which processor is to execute an HPF\_SERIAL procedure.

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# 11.1.2.1 SERIAL Calling Sequence

Prior to invocation of a SERIAL procedure from global HPF, the behavior of the program will be as if the following actions take place:

- 1. The processors are synchronized. All actions that logically precede the call are completed.
- 2. All actual arguments are remapped to the processor that will actually execute the SERIAL procedure. Each argument will appear to the SERIAL procedure as a sequential argument.

The behavior of the SERIAL procedure will be as if it was executed on only one processor. After the instance of the SERIAL procedure invoked from global HPF has completed, the behavior will be as if the following happens:

- 1. All processors are synchronized after the call.
- 2. The original mappings of actual arguments are restored.
- 11.2 Extrinsic Language Bindings

The previous section lays out the rules and considerations for execution models defined for HPF extrinsics. The HPF extrinsic interface is also used to tell the compiler what the language conventions of a called subprogram are. Four language bindings are defined here: HPF, Fortran, F77, and C. A given implementation may support additional interfaces or allow a user to construct custom interfaces. Taken together, these sections define the special extrinsics HPF\_LOCAL and HPF\_SERIAL.

The key feature of the language interface is an extended set of attributes for dummy arguments in explicit extrinsic interfaces, which can give the programmer control over aspects of argument passing between procedures of different extrinsic types. This mechanism is used extensively in the interfaces to C and Fortran 77, but it is defined in this more general context because it can also apply to other language interfaces.

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# 11.2.1 Control of Arguments

The special data attributes for dummy arguments in routines of certain extrinsic types are MAP\_TO, LAYOUT, and PASS\_BY. These may only appear in data types statements declaring dummy arguments within explicit interfaces to procedures of appropriate extrinsic types. In particular, all of these attributes have been defined for extrinsic interfaces of type LANGUAGE = 'C' (Section 11.4), and the latter two have been defined for extrinsic interfaces of type LANGUAGE = 'F77' (Section 11.6).

The purpose of this language extension is to increase the flexibility of the EXTRINSIC interface mechanism to facilitate argument passing between procedures written in different programming languages. These three attributes provide a convenient way to pass data between nearly equivalent data type representations and array layouts, as well as to allow for different data passing conventions and options. It should be noted, however, that these

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mechanisms are by no means expected to address the problems of full equivalence between data types and implementations of different languages.

In particular, the MAP\_TO attribute is designed to provide a standard, portable mecha-3 nism for passing arguments between data types (in different languages) that have substantial 4 overlap but not necessarily identical ranges of values or an identical machine representation. 5The programmer, not the language implementer, retains the responsibility for determining 6 whether or not any actual argument's value will be adequately represented in the new data 7type, or whether that value may be altered in successive operations involved in conversion 8 first to the new language, in operations within the extrinsic procedure, and then potentially 9 in conversion back to the original language. The LAYOUT attribute is used in cases when 10 the ordering of array elements within one or more processors may need to be changed when 11passing them as arguments between procedures of different languages. Finally, the PASS\_BY 12attribute is designed to offer more detailed control of passing mechanisms for arguments to 13 allow for differences between language implementations, to choose between distinct passing 14options offered in the non-HPF language, and to enable passing implementation-specific 15data structures when it is desired to convey HPF mapping information along with data 16values to non-HPF procedures. 17

These attributes are specified by an extension of the syntax rule R503 for *attr-spec* in the Fortran standard. Rule R501 for *type-declaration-stmt* is not changed except to refer for the extended *attr-spec*. The first two constraints below are repeated without change from the Fortran standard for clarity, since they apply generally to all attributes. The remaining constraints in the Fortran standard following rules R501 through R506 are specific to attributes already defined in the standard and will also be assumed but not repeated here.

These changes to Fortran syntax are made in anticipation of such extensions being considered for addition to the standard language in the next revision, as language interoperability is an area of active interest to the full Fortran community.

H1101 type-declaration-stmt-extended is type-spec [[, attr-spec-extended]...:] enfity-decl-list

			30
H1102 attr-spec-extended	$\mathbf{is}$	PARAMETER	31
	or	access-spec	32
	or	ALLOCATABLE	33
	or	DIMENSION ( $array$ -spec )	34
	or	EXTERNAL	35
	or	INTENT ( <i>intent-spec</i> )	36
	or	INTRINSIC	37
	or	OPTIONAL	38
	or	POINTER	39
	or	SAVE	40
	or	TARGET	40
	or	MAP_TO ( $map-to-spec$ )	42
	or	LAYOUT ( layout-spec )	43
	or		43
II 1102 man to an e	•		44
H1103 map-to-spec	$\mathbf{is}$	scalar-char-initialization- $expr$	
H1104 layout-spec	$\mathbf{is}$	scalar- $char$ - $initialization$ - $expr$	46
		· · · · · · · · · · · · ·	47
$ m H1105\ pass-by-spec$	$\mathbf{is}$	scalar- $char$ - $initialization$ - $expr$	48

#### 11.2. EXTRINSIC LANGUAGE BINDINGS

- Constraint: The same attr-spec-extended shall not appear more than once in a given typedeclaration-stmt.
  - Constraint: An entity shall not be explicitly given any attribute more than once in a scoping unit.

Constraint: The attributes MAP\_TO, LAYOUT, and PASS\_BY may be specified only for dummy arguments within a scoping unit of an extrinsic type for which these attributes have been explicitly defined.

The definition of *characteristics of a dummy data object* as given in F95:12.2.1.1 and extended in Section 8.15 is further extended to include the dummy data object's MAP\_TO, LAYOUT, and PASS\_BY attributes.

<sup>15</sup> In the MAP\_TO attribute, values of *map-to-spec* are intended to describe how the data <sup>16</sup> type of the named actual argument is mapped to the data type of the dummy argument in <sup>17</sup> the extrinsic procedure. An example is given in Section 11.4.2.1

18 For a given extrinsic type that allows the MAP\_TO attribute, the set of permitted values 19 for the *map-to-spec* will be specified. If the range of permitted values of the type and 20 mapped-to type differ, and the value of the actual argument or some subobject of the 21 actual argument is not within the permitted range of the mapped-to type, the value of the 22 associated dummy argument or subobject becomes undefined. Conversely, if the value of 23 the dummy argument or some subobject of the dummy is not within the permitted range of  $^{24}$ values of the associated actual argument, and the associated actual argument is a variable,  $^{25}$ the value of the associated actual argument or subobject of the actual becomes undefined.

If there is a mismatch between the precision, representation method, range of permitted values, or storage sequence between the type of the dummy argument and the permitted mapped-to type of the dummy argument, the compiler shall ensure that, for the duration of the reference to the extrinsic, the dummy argument is represented in a manner that is compatible with the expectations of the callee for an object of the permitted mapped-to type. Upon return from the procedure, the compiler shall ensure that the value of an actual argument that is a variable is restored to the specified type and kind.

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> Advice to users. This rule was created to ensure the portability of interoperability. However, it should be noted that for large objects, a significant overhead may be incurred if there is a mismatch between the representation method used for the data type versus the representation method used for the permitted mapped-to type. (End of advice to users.)

In the LAYOUT attribute, any permitted values of *layout-spec* for a given extrinsic interface are intended to describe how the data layout of the named actual argument is mapped to the data layout of the dummy argument in the extrinsic procedure. An example is given in Section 11.6.3. If no LAYOUT attribute is specified for a dummy array argument, the data layout shall be the same as if it were being passed to an HPF procedure of the same model, unless another default layout is defined for the given extrinsic type.

In the PASS\_BY attribute, any permitted values of *pass-by-spec* for a given extrinsic interface indicate a choice of mechanism used to associate the named actual argument with the dummy argument in the extrinsic procedure. Examples are given in Sections 11.6.3

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and 11.4.2.1. If no PASS\_BY attribute is specified, the argument association mechanism is implementation dependent, based on the compiler's knowlege of the extrinsic language processor.

#### 11.3 HPF Bindings

HPF is the default language assumption. It requires no MAP\_TO, LAYOUT, or PASS\_BY attributes in explicit interface definitions. All required subprogram binding information can be accomplished via the standard Fortran explicit interface.

The rules stated in section 14.7 of the Fortran standard will apply to variables defined in Fortran-based SERIAL and LOCAL scoping units. In particular, if the definition status, association status, or allocation status of a variable is defined upon execution of a RETURN statement or an END statement in a Fortran subprogram, such a variable in an SERIAL or LOCAL subprogram will be defined upon execution of a RETURN statement or an END statement.

Any I/O performed within an extrinsic subprogram of a different model, and the correspondence between file names and unit numbers used in global HPF and those used in local or serial subprogram code will be implementation defined.

#### 11.3.1 Additional Special Considerations for HPF\_LOCAL

There are some considerations about what HPF features may be used in writing a local, per-processor procedure.

Local program units can use all HPF constructs except for REDISTRIBUTE and REALIGN.  $^{24}$ Moreover, DISTRIBUTE, ALIGN, and INHERIT directives may be applied only to dummy  $^{25}$ arguments; that is, every *alignee* and *distributee* must be a dummy argument, and every 26 align-target must be a template or a dummy argument. Mapping directives in local HPF 27program units are understood to have global meaning, as if they had appeared in global 28 HPF code, applying to the global array of which a portion is passed on each processor. 29 (The principal use of such mapping directives is in an HPF\_LOCAL module that is used by a 30 global HPF module.) 31

HPF\_ALIGNMENT, HPF\_TEMPLATE, and HPF\_DISTRIBUTION, the distribution query library subroutines, may be applied to non-sequential local arrays. Their outcome is the same as for a global array that happens to have all its elements on a single node.

As introduced in Section 6.3.1, a local HPF program unit may not access global HPF <sup>35</sup> data other than data that is accessible, either directly or indirectly, via the actual arguments. <sup>36</sup> In particular, a local HPF program unit does not have access to global HPF COMMON blocks; <sup>37</sup> COMMON blocks appearing in local HPF program units are not identified with global HPF <sup>38</sup> COMMON blocks. The same name may not be used to identify a COMMON block both within a <sup>39</sup> local HPF program unit and an HPF program unit in the same executable program. <sup>40</sup>

Like local variables in local subprograms, COMMON blocks in local subprograms contain 41 local data, held in local storage on each processor. This storage is only accessible locally, 42 and it will in general contain data that is different on each processor. Indeed, the size of a 43 local COMMON block can be different on each processor. 44

According to the Fortran specification, a COMMON block that goes out of scope is not 45 preserved, unless it has the SAVE attribute. This is true of local COMMON blocks, which should 46 be given the SAVE attribute if they are intended to convey information between calls to the 47 local subprogram. 48

Scalars of an explicitly mapped derived type cannot be passed to an HPF\_LOCAL subprogram. 2

3 The attributes (type, kind, rank, optional, intent) of the dummy arguments must match the attributes of the corresponding dummy arguments in the explicit interface. A dummy 4 argument of an EXTRINSIC('HPF', 'LOCAL') routine may not be a procedure name. 5

A dummy argument of an EXTRINSIC('HPF', 'LOCAL') routine may not have the 6 **POINTER** attribute. 7

A nonsequential dummy array argument of an EXTRINSIC ('HPF', 'LOCAL') routine 8 must have assumed shape. Note that, in general, the shape of a dummy array argument 9 differs from the shape of the corresponding actual argument, unless there is a single exe-10  $1\,1$ cuting processor.

Explicit mapping directives for dummy arguments may appear in a local procedure. 12Such directives are understood as applying to the global array whose local sections are passed 13 as actual arguments or results on each processor. If such directives appear, corresponding 14mapping directives must be visible to every global HPF caller. This may be done by 1516providing an interface block in the caller, or by placing the local procedure in a module of extrinsic kind HPF\_LOCAL that is then used by the global HPF program unit that calls the 1718 local procedure.

An EXTRINSIC ('HPF', 'LOCAL') routine may not be invoked, either directly or indi-19rectly, in the body of a FORALL construct or in the body of an INDEPENDENT loop. 20

A local procedure may have several ENTRY points. A global HPF caller must contain a separate extrinsic interface for each entry point that can be invoked from the HPF program.

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#### **Argument Association** 11.3.2

If a dummy argument of an EXTRINSIC('HPF', 'LOCAL') routine is a scalar, then the cor-26 responding dummy argument of the local procedure must be a scalar of the same type and 27 type parameters. Only scalars of intrinsic types, or of derived types that are not explicitly 28 mapped, may be passed from a global to an HPF\_LOCAL routine. When the extrinsic proce-29 dure is invoked, the local procedure is passed an argument that consists of a local copy of 30 the scalar. This copy will be a valid HPF scalar. 31

If a dummy argument of an EXTRINSIC ('HPF', 'LOCAL') routine is an array, then the 32 corresponding dummy argument in the specification of the local procedure must be an array 33 of the same rank, type, and type parameters. 34

If the array is sequential in the extrinsic interface, the corresponding actual argument 35 will be passed by replicatation on all processors, just as scalar arguments are passed. Each 36 local dummy argument will be associated with a full copy of the actual array argument. 37 The dummy argument in the extrinsic interface and the corresponding dummy argument 38 in the specification of the local procedure may be declared with the same explicit shape. 39 All sequential dummy arguments passed by replication to an EXTRINSIC('HPF', 'LOCAL') 40 procedure must either be INTENT(IN) arguments or should be updated consistently across 41 processors. 42

If the dummy argument is a nonsequential array, then, when the extrinsic procedure is 43 invoked, the local dummy argument is associated with the local array that consists of the 44 subgrid of the global array that is stored locally. This local array will be a valid HPF array. 45If an EXTRINSIC ('HPF', 'LOCAL') routine is a function, then the local procedure is a 46 function that returns a scalar of the same type and type parameters as the HPF extrinsic 47function. The value returned by each local invocation must be the same. 48

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Each physical processor has at most one copy of each HPF variable. Consider the following extrinsic interface:

```
INTERFACE
EXTRINSIC('HPF','LOCAL') FUNCTION MATZOH(X, Y) RESULT(Z)
REAL, DIMENSION(:,:) :: X
REAL, DIMENSION(:) :: Y
REAL Z
!HPF$ ALIGN WITH X(:,*) :: Y(:)
! note that this asserts that size(Y) = size(X,1)
!HPF$ DISTRIBUTE X(BLOCK, CYCLIC)
END FUNCTION
END INTERFACE
```

The corresponding local HPF procedure is specified as follows.

```
EXTRINSIC('HPF', 'LOCAL') FUNCTION MATZOH(XX, YY) RESULT(ZZ)

REAL, DIMENSION(:,:) :: XX

REAL, DIMENSION(5:) :: YY ! assumed shape with lower bound of 5

REAL ZZ

NX1 = SIZE(XX, 1)

LX1 = LBOUND(XX, 1)

UX1 = UBOUND(XX, 1)

NX2 = SIZE(XX, 2)

LX2 = LBOUND(XX, 2)

UX2 = UBOUND(XX, 2)

NY = SIZE(YY, 1)

LY = LBOUND(YY, 1)

UY = UBOUND(YY, 1)

...

END FUNCTION
```

Assume that the function is invoked with an actual (global) array X of shape  $3 \times 3$  and an actual vector Y of length 3 on a 4-processor machine, using a  $2 \times 2$  processor arrangement (assuming one abstract processor per physical processor).

Then each local invocation of the function MATZOH receives the following actual arguments:

Processor $(1,1)$	Processor $(1,2)$	37
X(1,1) X(1,3) X(2,1) X(2,3)	X(1,2) X(2,2)	38 39
Y(1) Y(2)	Y(1) Y(2)	40 41
		42
Processor $(2,1)$	Processor $(2,2)$	43
X(3,1)  X(3,3)	X(3,2)	44
Y(3)	Y(3)	45
		46

Here are the values to which each processor would set NX1, LX1, UX1, NX2, LX2, UX2, NY, LY, and UY:

1	Processor $(1,1)$	Processor $(1,2)$
2	NX1 = 2 $LX1 = 1$ $UX1 = 2$	NX1 = 2 $LX1 = 1$ $UX1 = 2$
3	NX2 = 2 $LX2 = 1$ $UX2 = 2$	NX2 = 1 $LX2 = 1$ $UX2 = 1$
4	NY = 2 $LY = 5$ $UY = 6$	NY = 2 $LY = 5$ $UY = 6$
5	<b>Processor</b> $(2,1)$	$\mathbf{P}_{\mathrm{reconstant}}(2,2)$
6	Processor $(2,1)$	Processor $(2,2)$
7	NX1 = 1 $LX1 = 1$ $UX1 = 1$	$\mathbf{NX1} = 1  \mathbf{LX1} = 1  \mathbf{UX1} = 1$
8	NX2 = 2  LX2 = 1  UX2 = 2	NX2 = 1 $LX2 = 1$ $UX2 = 1$
9	NY = 1 $LY = 5$ $UY = 5$	NY = 1 $LY = 5$ $UY = 5$
10	An actual argument to an extrinsic procedu	re may be a pointer. Since the correspond-
11 12	ing dummy argument may not have the POINTE	R attribute, the dummy argument becomes
12	associated with the target of the HPF global poi	
14	pointer associated with a global HPF target. The	,
15	a derived-type containing a pointer component.	
16	Rationale. It is expected that global point	nter variables will have a different represen-
17		, at least on distributed memory machines,
18	1	ormation for global addressing. This restric-
19	tion could be lifted in the future. ( <i>End of</i>	
20	· · · ·	,
21	1 0 ,	or PRESENT, should also behave as expected.
22	Note that when a global array is passed to a loc	cal routine, some processors may receive an
23	empty set of elements.	
24		
25	11.3.3 Special Considerations for HP	F_SERIAL
26 27	There are restrictions that apply to an HPF_SER	IAL subprogram.
28	No $specification-directive, realign-directive,$	or <i>redistribute-directive</i> is permitted to be
29	appear in an HPF_SERIAL subprogram or interfa	.ce body.
30	Rationale. An HPF mapping directive we	ould likely be meaningless in an HPF_SERIAL
31		<i>lent-directive</i> may appear in an HPF_SERIAL
32		gful information to a compiler about a DO
33	loop or a FORALL statement or construct.	
34		()
35		result variables of an HPF_SERIAL procedure
36	will be considered to be sequential.	
37	. 0	ain a definition of a common block that has
38	the same name as a common block defined in	
39	addition, an HPF_SERIAL subprogram must not	
40	block if an HPF or HPF_LOCAL program unit ha	
41		riable of an HPF_SERIAL procedure that is
42	referenced in global HPF must not have the PC	
43	argument or function result of an HPF_SERIAL p must not have the POINTER attribute.	procedure that is referenced in global HPF,
44 45		cedure that is referenced in global HPF and
40	A dummy argument of all REF DERIAL PLOC	coure that is referenced in global III I allo

as A dummy argument of an HPF\_SERIAL procedure that is referenced in global HPF a any subobject of such a dummy argument must not have the TARGET attribute.

A dummy procedure argument of an HPF\_SERIAL procedure must be an HPF\_SERIAL
 procedure.

```
PROGRAM MY_TEST
                                                                                    1
        INTERFACE
                                                                                    2
          EXTRINSIC('HPF', 'SERIAL') SUBROUTINE GRAPH_DISPLAY(DATA)
             INTEGER, INTENT(IN) :: DATA(:, :)
          END SUBROUTINE GRAPH_DISPLAY
                                                                                    5
        END INTERFACE
                                                                                    6
        INTEGER, PARAMETER :: X_SIZE = 1024, Y_SIZE = 1024
                                                                                    8
                                                                                    9
        INTEGER DATA_ARRAY(X_SIZE, Y_SIZE)
                                                                                    10
!HPF$
        DISTRIBUTE DATA_ARRAY(BLOCK, BLOCK)
                                                                                    1\,1
                                                                                   12
i
   Compute DATA_ARRAY
                                                                                   13
        . . .
                                                                                    14
        CALL DISPLAY_DATA(DATA_ARRAY)
                                                                                   15
      END PROGRAM MY_TEST
                                                                                    16
                                                                                    17
! The definition of a graphical display subroutine.
                                                                                    18
! In some implementation-dependent fashion,
                                                                                   19
! this will plot a graph of the data in DATA.
                                                                                   20
                                                                                   21
      EXTRINSIC('HPF', 'SERIAL') SUBROUTINE GRAPH_DISPLAY(DATA)
                                                                                   22
        INTEGER, INTENT(IN) :: DATA(:, :)
                                                                                   23
        INTEGER :: X_IDX, Y_IDX
                                                                                   ^{24}
                                                                                   ^{25}
        DO Y_{IDX} = LBOUND(DATA, 2), UBOUND(DATA, 2)
                                                                                   26
          DO X_IDX = LBOUND(DATA, 1), UBOUND(DATA, 1)
                                                                                   27
           . . .
                                                                                   28
          END DO
                                                                                   29
        END DO
                                                                                   30
      END SUBROUTINE GRAPH_DISPLAY
                                                                                   31
                                                                                   32
                                                                                   33
```

### 11.4 C Language Bindings

A common problem faced by Fortran users is the need to call procedures written in other languages, particularly those written in C or ones that have interfaces that can be described by C prototypes. Although many Fortran implementations provide methods that solve this problem, these solutions are rarely portable. 34

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This section defines a method of specifying interfaces to procedures defined in C that removes most of the common obstacles to interoperability, while retaining portability.

### 11.4.1 Specification of Interfaces to Procedures Defined in C

If a user wishes to specify that a procedure is defined by a C procedure, this is specified with an *extrinsic-spec-arg* of LANGUAGE = 'C', or an *extrinsic-kind-keyword* of C, as specified in Section 6.

For C subprograms for which EXTRINSIC (LANGUAGE = 'C') has been specified, the 47 constraints associated with the syntax for *attr-spec-extended* (H1102) are extended as fol-48

1	lows:	
2 3 4	Constraint:	A LANGUAGE = 'C' function shall have a scalar result of type integer, real or double precision.
5 6 7 8	Constraint:	A dummy argument of a LANGUAGE = 'C' procedure shall not be an assumed- shape array, shall not have the POINTER attribute, shall not have the TARGET attribute, nor shall it have a subobject that has the POINTER attribute.
9 10 11 12	Constraint:	The bounds of a dummy argument shall not be specified by specification expressions that are not constant specification expressions, nor shall the character length parameter of a dummy argument of such a procedure be specified by a specification expression that is not a constant specification expression.
13 14 15	Constraint:	A $dummy$ - $arg$ - $list$ of a LANGUAGE = 'C' subroutine shall not have a $dummy$ - $arg$ that is $*$ or a dummy procedure.
16 17 18 19 20 21	name of the name specif	ue of the <i>scalar-char-initialization-expr</i> in the EXTERNAL_NAME specifier gives the procedure as defined in C. This value need not be the same as the procedure ied by the <i>function-stmt</i> or <i>subroutine-stmt</i> . If EXTERNAL_NAME is not specified, were specified with a value that is the same as the procedure name in lower case
22 23 24 25 26	the na the pr compi	e to users. Note that the EXTERNAL_NAME specifier does not necessarily specify ame by which a binder knows the procedure. It specifies the name by which cocedure would be known if it were referenced by a C program, and the HPF ler is required to perform any transformations of that name that the C compiler perform.
27 28 29 30 31	be per as a w	XTERNAL_NAME specifier also allows the user to specify a name that might not mitted by an HPF compiler, such as a name beginning with an underscore, or vay of enforcing the distinction between upper and lower case characters in the (End of advice to users.)
32 33 34 35	is defined in	trinsic-spec-arg of LANGUAGE = 'C' helps a compiler identify a procedure that C so that it can take appropriate steps to ensure that the procedure is invoked her required by the C compiler.
36 37 38 39 40 41 42	than c procec compi it mag	e to implementors. A vendor may feel compelled to provide support for more one C compiler, if different C compilers available for a system provide different dure calling conventions or different data type sizes. For instance, a vendor's ler may provide support for a value of GNU_C in the LANGUAGE= specifier, or y provide support through the use of compiler switches. (End of advice to mentors.)
43 44	11.4.2 S	pecification of Data Type Mappings for C
45 46		ic dummy argument feature, consisting of the MAP_TO, LAYOUT, and PASS_BY s the principal feature that facilitates referencing procedures defined in C from

attributes, is the principal feature that facilitates referencing procedures defined in C from
 within Fortran programs. Together, these attributes allow the user to specify conversions
 required to associate the actual arguments specified in the procedure reference with the

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formal arguments defined by the referenced procedure. In particular, the MAP\_TO attribute indicates the type of the C data to which the HPF data shall be converted by the compiler; the PASS\_BY attribute indicates whether a C pointer to the dummy argument needs to be passed; the LAYOUT attribute indicates for an array whether the array element order needs to be changed from Fortran's array element ordering to C's.

For C, the constraints associated with *attr-spec-extended*, *map-to-spec*, *layout-spec*, and *pass-by-spec* (H1102–H1105) are further extended as follows.

- Constraint: The MAP\_TO attribute shall be specified for all dummy arguments and function result variables of a LANGUAGE = 'C' explicit interface.
- Constraint: The *map-to-spec* associated with a dummy argument shall be compatible with the type of the dummy argument. (See below for compatibility rules.)
- Constraint: A LAYOUT attribute shall only be specified for a dummy argument that is an array.
- Constraint: A LAYOUT attribute shall not be specified for an assumed-size array.

If the compiler is capable of representing letters in both upper and lower case, the value specified for a *map-to-spec*, *layout-spec* or *pass-by-spec* is without regard to case. Any blanks specified for a *map-to-spec*, *layout-spec* or *pass-by-spec* shall be ignored by the compiler for the purposes of determining its value.

An implementation shall provide a module, ISO\_C, that shall define a derived type, C\_VOID\_POINTER. The components of the C\_VOID\_POINTER type shall be private.

Advice to users. The C\_VOID\_POINTER type provides a method of using void \* pointers in a program, but does not give the user any way of manipulating such a pointer in the Fortran part of the program, since I/O cannot be performed on an object with private components outside the module that defines the type, neither can the components or structure constructor of such a structure be used outside of the module that defines the type. (End of advice to users.)

The values permitted for a *map-to-spec* for LANGUAGE = 'C' are 'INT', 'LONG', 'SHORT', 'SIGNED\_CHAR', 'FLOAT', 'DOUBLE', 'LONG\_DOUBLE', 'CHAR', 'CHAR\_PTR', 'VOID\_PTR', or a comma-separated list, delimited by parentheses, of any of these values. The HPF types with which these are compatible are shown in the table below.

A *map-to-spec* that is a parenthesized list of values is compatible with a dummy argument of derived type if each value in the list is compatible with the corresponding component of the derived type.

When the PASS\_BY attribute is used, the values permitted for a *pass-by-spec* are 'VAL', <sup>39</sup> '\*', or '\*\*'. If no PASS\_BY attribute is specified, then PASS\_BY ('VAL') is assumed. If a <sup>40</sup> *pass-by-spec* of VAL is specified, the dummy argument shall not have the INTENT(OUT) or <sup>41</sup> INTENT(INOUT) attribute specified. If a value of '\*' or '\*\*' is specified for the *pass-by-spec*, <sup>42</sup> an associated actual argument shall be a variable. <sup>43</sup>

The value of the *map-to-spec* specified for a dummy argument in the interface body of a procedure for which a LANGUAGE= specifier whose value is C appears shall be such that at least one of the permitted mapped-to types is the same as the C data type of the corresponding formal argument in the C definition of the procedure (or a type that is compatible with one of the permitted mapped-to types). The C data type of a function in the C definition 48

#### 11.4. CLANGUAGE BINDINGS

of a procedure shall be one of the permitted mapped-to types (or a type that is equivalent to the permitted mapped-to types) specified for the function result variable in the interface body of a function with the LANGUAGE= specifier whose value is C. If a subroutine has been specified with a LANGUAGE= specifier whose value is C, the C definition of the procedure shall be specified with a data type of void.

The permitted mapped-to types for scalar dummy arguments of intrinsic type or of the derived type C\_VOID\_POINTER, are shown in the following table.

MAP_TO	Compatible		C Type if PASS.	BY
	With	'VAL'	'*'	'**'
'INT'	INTEGER	int	int*	int**
'LONG'	INTEGER	long	long*	long**
'SHORT'	INTEGER	short	short*	short**
'SIGNED_CHAR'	INTEGER	signed char	signed char*	signed char**
'FLOAT'	REAL	float	float*	float**
'DOUBLE'	REAL	double	double*	double**
'LONG_DOUBLE'	REAL	double	double*	double**
'CHAR'	CHARACTER(1)	char	char*	char**
'CHAR_PTR'	CHARACTER	char*	char**	char***
'VOID_PTR'	C_VOID_POINTER	void*	void**	void***

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> The permitted mapped-to types of an array are the same as the permitted mappedto types of a scalar variable of that type followed by a left bracket ([), followed by the extent of the corresponding dimension of the dummy argument, followed by a right bracket (]), for each dimension of the array. If no value is specified for the LAYOUT attribute, the corresponding dimensions of the dummy argument are determined from right to left; if the value C\_ARRAY is specified for the LAYOUT attribute, the corresponding dimensions of the dummy argument are determined from left to right.

29 30

The value permitted for a LANGUAGE = 'C' layout-spec is C\_ARRAY.

The permitted mapped-to types of a scalar variable of derived type are the structures whose corresponding members are of one of the permitted mapped-to types of the components of the derived type.

If there is a mismatch between the precision, representation method, range of permitted 34 values or storage sequence between the type of the dummy argument and the permitted 35 mapped-to type of the dummy argument, the compiler shall ensure that, for the duration 36 of the reference to a procedure defined with a LANGUAGE= specifier whose value is C, the 37 dummy argument is represented in a manner that is compatible with the expectations of 38 the C processor for an object of the permitted mapped-to type. Upon return from the 39 procedure, the compiler shall ensure that the value of an actual argument that is a variable 40 is restored to the specified type and kind. 41

If the range of permitted values of the type and mapped-to type differ and the value of the actual argument or some subobject of the actual argument is not within the permitted range of the mapped-to type, the value of the associated dummy argument or subobject becomes undefined. Conversely, if the value of the dummy argument or some subobject of the dummy is not within the permitted range of values of the associated dummy argument, and the associated actual argument is a variable, the value of the associated actual argument or subobject of the actual becomes undefined.

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Advice to users. These rules were created to ensure the portability of interoperability. However, it should be noted that for large objects, a significant overhead may be incurred if there is a mismatch between the representation method used for the data type versus the representation method used for the permitted mapped-to type. (*End* of advice to users.)

Advice to users. In some cases, this may cause the value of the actual argument to change without the value being modified by the procedure referenced. For example,

```
PROGRAM P
          INTERFACE
            EXTRINSIC(LANGUAGE='C') SUBROUTINE C_SUB(R,I)
              REAL(KIND(1.0D0)), MAP_TO('FLOAT'), PASS_BY('*') :: R
              INTEGER, MAP_TO('INT'), PASS_BY('*') :: I
            END SUBROUTINE C_SUB
          END INTERFACE
          REAL(KIND(0.0D0)) RR
          RR = 1.0D0 + 1.0D-10
          I = 123456789
          PRINT *, RR
          CALL C_SUB(RR, I)
          PRINT *, RR
       END PROGRAM P
       void c_sub(float *r, int *i)
        {
       }
might print
   1.000000001000000
   although the value of *r is not modified in c_sub. Similarly, the value of I might
become undefined after the reference to c_sub, although *i is not modified.
```

Although it is good practice to avoid specifying a mapped-to type of float for a dummy argument of any type other than default real, or a mapped-to type of double for a dummy argument of any type other than double precision real, selecting an appropriate dummy argument type for objects requiring a mapped-to type int or long might not be so simple. (*End of advice to users.*)

If no *layout-spec* is specified for a dummy array argument, the array element order shall <sup>45</sup> be the same as that specified by Fortran. If the value of *layout-spec* specified is C\_ARRAY, <sup>46</sup> the array element order of the array shall be transposed for the duration of the reference <sup>47</sup> to the procedure. <sup>48</sup>

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1 11.4.2.1 Examples of Data Type Mappings

Some examples should help to clarify what sorts of C procedure definitions would be permitted given an interface body in a Fortran program. For example, the following interface body

```
6
                INTERFACE
7
                  EXTRINSIC('C') SUBROUTINE C_SUB(I, R, DARR, STRUCT)
8
                    INTEGER, MAP_TO('INT') :: I
9
                     REAL, MAP_TO('FLOAT'), PASS_BY('*') :: R
10
                    REAL(KIND(1.0D0)), MAP_TO('DOUBLE') :: DARR(10)
11
                    TYPE DT
12
                       SEQUENCE
13
                       INTEGER :: I, J
14
                    END TYPE DT
15
                    TYPE(DT), MAP_TO('(INT, LONG)'), PASS_BY('*') :: STRUCT
16
                  END SUBROUTINE C_SUB
17
                END INTERFACE
18
         could correspond to a C procedure that has the prototype
19
20
          void c_sub(int i, float r*, double darr[10], struct {int i, long j} *)
21
22
         In the following example of the LAYOUT attribute,
23
              PROGRAM P
24
                INTERFACE
25
                  EXTRINSIC('C') SUBROUTINE C_SUB(A, B)
26
                    INTEGER, MAP_TO('INT') :: A(2,2)
27
                     INTEGER, MAP_TO('INT'), LAYOUT('C_ARRAY') :: B(2,2)
28
                  END SUBROUTINE C_SUB
29
                END INTERFACE
30
31
                INTEGER :: AA(2,2), BB(2,2)
32
                CALL C_SUB(AA, BB)
33
              END PROGRAM P
34
35
              void c_sub(int a[2][2], b[2][2])
36
37
         the correspondence between elements of AA and a, and elements of BB and b is
38
                                                             b[0][0]
                AA(1,1)
                           a[0][0]
                                                   BB(1,1)
39
                AA(2,1)
                           a[0][1]
                                                   BB(2,1)
                                                              b[1][0]
40
                AA(1,2)
                           a[1][0]
                                                   BB(1,2)
                                                             b[0][1]
41
42
                AA(2,2)
                           a[1][1]
                                                  BB(2,2)
                                                             b[1][1]
43
44
            Fortran Language Bindings
     11.5
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```

When the language specified in an extrinsic definition is Fortran the rules are basically the
same as those for HPF because HPF is based on the Fortran standard. There are a few
issues to consider in this case:

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- Only Fortran constructs should be used. Features such as asynchronous I/O or the HPF library may not be supported.
- It is recommended that Fortran language processors to be used for this purpose be extended to support the HPF\_LOCAL distribution query routines and the associated HPF\_LOCAL\_LIBRARY
- Assuming the intent is to compile the extrinsic routines with a Fortran processor, these routines should be in separate files rather than incorporated into files with HPF source code.
- The programmer should expect any HPF directives may be ignored.

### 11.6 Fortran 77 Language Bindings

For language interface purposes, Fortran 77 is still essentially a subset of ANSI/ISO standard Fortran, so most considerations relating to HPF calling Fortran also apply to HPF calling Fortran 77 extrinsic procedures. However, two chief differences between Fortran and Fortran 77 complicate the specification of any EXTRINSIC(LANGUAGE='F77'), interface from HPF, especially for the local model:

- Arguments are usually passed differently. Fortran 77 implementations typically pass arguments between subprograms by address (reference). That is, no other information about the actual argument is passed for example, data type, size, distribution, etc. In contrast, HPF implementations often pass by variables by descriptor in order to make such information available to the subprogram.
- Very different information about how array elements are to be assigned to specific memory locations is available to Fortran 77 and HPF programmers.

In Fortran 77, the declaration of an array prescribes exactly the mapping of array elements to the linear sequence of storage locations. In HPF, the mapping of ar-ray elements to different processors may be controlled (e.g., with DISTRIBUTION and ALIGN directives) and queried (e.g., with HPF\_ALIGNMENT, HPF\_DISTRIBUTION, and HPF\_TEMPLATE) but there is absolutely no information about how array elements on a given processor are organized within local, serial memory. Even in Fortran 90, as-sumed shape dummy arrays, for example, do not have to follow the same storage and sequence association rules as Fortran 77 arrays do. 

Indeed, different HPF compilers may organize the data locally in different manners — perhaps including border cells for "stencil" optimizations, or global padding to ensure equal-size subgrids on all processors. Certainly, different HPF compilers are not *bound* to organize local data in any particular manner, and some may choose imaginative orderings in such cases as SMP's, for example.

### 11.6.1 Special Considerations for F77\_LOCAL

The EXTRINSIC (F77 LOCAL) interface extends the HPF LOCAL and FORTRAN\_LOCAL extrinsic interfaces to meet the needs of Fortran 77 programmers.

This EXTRINSIC type uses the syntax for calling extrinsic subprograms described above. 47 It can be described more precisely as an EXTRINSIC(LANGUAGE='F77', MODEL='LOCAL') 48

interface. The basic conventions for transferring control between global and local routines 1 described previously in Section 11.1 also apply. 2

3 However, the differences in argument passing and data distribution between these two languages, as well as the different possible motivations for using such an interface, can be better addressed by allowing additional options for passing data and distribution information. These options are provided with the help of LAYOUT and PASS\_BY attributes. 6

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#### 11.6.2Argument Passing to F77\_LOCAL Procedures

A typical Fortran 77 implementation passes arguments by reference, usually by passing 10 the base address of the location of the first data element, and such arguments may also 11 be assumed to be sequence associated. These facts make it most practical for the default 12 method of passing a distributed data structure from HPF to an F77\_LOCAL procedure by 13 passing the base address of that section of local memory that has been allocated to it. 14 To allow for sequence association of actual and dummy arguments, data should also be 15 reordered or compressed or both, if necessary, on all processors. This is the safest method 16 of passing distributed data to an EXTRINSIC (F77\_LOCAL) procedure, and hence it should 17be the default one. However, it tends to have the greatest performance costs. 18

A second argument passing option is to pass distributed array data "as is" from a 19 global HPF procedure to the local F77 ones, not guaranteeing sequence association of the 20 dummy arguments in order to avoid unwanted local data motion that might be required 21 to compress or reorder the elements of an array local to a processor. In other words, it 22 should be possible to do no more data motion than if the same argument were being passed 23 to another HPF procedure. The guarantee of a sequence associated dummy argument is  $^{24}$ sacrificed for the possible gains in performance available because the local components of  $^{25}$ the actual argument are not reordered or compressed. The local programmer must be able 26 to use the implementation-dependent ordering created by the global HPF program. 27

A third option that can be useful to permit HPF\_LOCAL-style local programming from 28 an EXTRINSIC (F77\_LOCAL) procedure call is to pass an array via a descriptor or handle, as 29 is typically done in HPF implementations or for Fortran 90 assumed shape arrays. The local 30 procedure may not access elements of this dummy argument directly but may only pass it 31 on to special utility routines, perhaps to obtain local or global distribution information. 32

The following attributes suffice to support the above three alternate form of passing data to an EXTRINSIC(F77\_LOCAL)x procedure:

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• LAYOUT('F77\_ARRAY') indicates that the rectangular configuration should be FOR-TRAN 77 sequence associated in local, serial memory.

For example, many compilers add border elements for "stencil" optimizations or pad array allocations on particular processors so that all processors allocate equal amounts of memory for each array. Local reordering eliminates such padding and provides FORTRAN 77 sequence association for actual data values.

Any local reordering is in addition to any global remapping that may be dictated by 43 DISTRIBUTION or ALIGN directives in the INTERFACE block.

- 45If no LAYOUT attribute is specified, then LAYOUT ('F77\_ARRAY') is assumed.
- LAYOUT('HPF\_ARRAY') indicates that an array argument is passed just as it would be 47 to a global HPF procedure, with no local reordering of the data. 48

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This option is desirable when the programmer decides that the overhead of local 1 reordering should be eliminated or that certain characteristics of the global HPF 2 compiler's ordering (border cells, equal-size allocations among processors, etc.) should 3 be preserved at the local level. It forces the local programmer to access the local data 4 in whatever implementation-dependent style the global HPF compiler employs. 5

Furthermore, each argument in the INTERFACE block can also have its PASS\_BY attribute specified to indicate whether the data is passed by reference, for Fortran 77-style access, or via a special handle, perhaps a descriptor used for HPF variable passing, that permits the global HPF caller to pass special mapping information for use within the local Fortran 77 procedure.

- PASS\_BY('\*') indicates that the local procedure should be able to access the dummy argument locally as an F77-style variable, passed by reference.
- PASS\_BY('HPF\_HANDLE') indicates that the local procedure should receive a reference to a global descriptor that can be used with special inquiry routines to obtain useful distribution information.

Thus, the default dummy argument attributes are LAYOUT('F77\_ARRAY'), a guarantee of sequence association, and PASS\_BY('\*'), an indication that data is being passed via a pointer to its location.

#### Advice to implementors.

 $^{24}$ In addition to providing argument passing and data reordering options, a good  $^{25}$ EXTRINSIC (F77 LOCAL) implementation should address the problem of declaring arbi-26 trary sized local subgrids and accessing their elements without being able to describe  $^{27}$ them as assumed-shape arrays, as in HPF. Dealing with the local results of global 28 data distributions within each local procedure initiated by an extrinsic procedure call 29 can also be difficult without Fortran 90 array inquiry functions and the inquiry sub-30 routines in the HPF Library. Special inquiry routines, callable globally or locally, 31 such as the proposed library of Fortran 77 function interfaces in Annex G are recom-32 mended as supplements to the EXTRINSIC(F77\_LOCAL) procedure interface in order 33 to permit more flexible and efficient use of a broad range of possible global HPF data 34 distributions. 35

(End of advice to implementors.)

#### 11.6.3 F77\_LOCAL Programming Examples

# 11.6.3.1 LAYOUT('F77\_ARRAY') and PASS\_BY('\*')

This example illustrates F77\_LOCAL programming using the default LAYOUT('F77\_ARRAY') and PASS\_BY('\*') attributes, and the use of inquiry routines from the local level using the LAYOUT('HPF\_ARRAY') attribute.

• HPF caller

PROGRAM EXAMPLE

```
! Declare the data array and a verification copy
1
                  INTEGER, PARAMETER :: NX = 100, NY = 100
2
3
                  REAL, DIMENSION(NX,NY) :: X, Y
            !HPF$ DISTRIBUTE(BLOCK, BLOCK) :: X, Y
4
5
            ! The global sum will be computed
6
            ! by forming partial sums on the processors
7
                  REAL PARTIAL_SUM(NUMBER_OF_PROCESSORS())
8
            !HPF$ DISTRIBUTE PARTIAL_SUM(BLOCK)
9
10
11
            ! Local subgrid parameters are declared per processor
            ! for a rank-two array
12
                  INTEGER, DIMENSION(NUMBER_OF_PROCESSORS(),2) ::
13
                 & LB, UB, NUMBER
14
            !HPF$ DISTRIBUTE(BLOCK,*) :: LB, UB, NUMBER
15
16
            ! Define interfaces
17
                  INTERFACE
18
19
                    EXTRINSIC(F77_LOCAL) SUBROUTINE LOCAL1
20
                      (LB1, UB1, LB2, UB2, X, X_DESC)
                 &
21
                     INTEGER, DIMENSION(:) :: LB1, UB1, LB2, UB2
22
                     REAL,DIMENSION(:,:),LAYOUT('HPF_ARRAY') :: X
23
                     REAL,DIMENSION(:,:),LAYOUT('HPF_ARRAY'),
                                                                     &
24
                                                             :: X_DESC
                            PASS_BY('HPF_HANDLE')
^{25}
26
                    DISTRIBUTE(BLOCK) :: LB1, UB1, LB2, UB2
            !HPF$
27
            !HPF$
                    DISTRIBUTE(BLOCK, BLOCK) :: X, X_DESC
28
                     END
29
30
                     EXTRINSIC(F77_LOCAL) SUBROUTINE LOCAL2(N,X,R)
31
                     INTEGER N(:)
32
                    REAL X(:,:), R(:)
33
            ! Defaults:
34
                    LAYOUT('F77_ARRAY')
                                              sequential, column-major storage
            1
35
            !
                    PASS_BY('*')
                                              pass by reference (local address)
36
            !HPF$
                  DISTRIBUTE N(BLOCK)
37
            !HPF$ DISTRIBUTE X(BLOCK, BLOCK)
38
            !HPF$
                    DISTRIBUTE R(BLOCK)
39
                    END
40
41
42
                  END INTERFACE
43
            ! Determine result using only global HPF
44
45
                   ! Initialize values
46
                  FORALL (I=1:NX, J=1:NY) X(I, J) = I + (J-1) * NX
47
48
```

```
! Determine and report global sum
                                                                                 1
          PRINT *, 'Global HPF result: ',SUM(X)
                                                                                 2
                                                                                 3
    ! Determine result using local subroutines
                                                                                 4
                                                                                 5
          ! Initialize values ( assume stride = 1 )
                                                                                 6
          CALL HPF_SUBGRID_INFO( Y, IERR, LB=1b, UB=UB )
                                                                                 7
          IF (IERR.NE.O) STOP 'ERROR!'
                                                                                 8
          CALL LOCAL1( LB(:,1), UB(:,1), LB(:,2), UB(:,2), Y , Y )
                                                                                 9
                                                                                 10
          ! DETERMINE AND REPORT GLOBAL SUM
                                                                                 11
          NUMBER = UB - LB + 1
                                                                                 12
          CALL LOCAL2 ( NUMBER(:,1) * NUMBER(:,2) , Y , PARTIAL_SUM )
                                                                                 13
          PRINT *, 'F77_LOCAL result #1 : ',SUM(PARTIAL_SUM)
                                                                                 14
                                                                                 15
          END
                                                                                 16
                                                                                 17
• FORTRAN 77 callee
                                                                                 18
                                                                                 19
          SUBROUTINE LOCAL1 (LB1, UB1, LB2, UB2, X, DESCRX)
                                                                                 20
                                                                                21
          REAL X ( LB1 : UB1 , LB2 : UB2 )
                                                                                22
          INTEGER DESCRX ( * )
                                                                                23
                                                                                 ^{24}
    ! Get the global extent of the first axis
                                                                                 ^{25}
    ! This is an HPF_LOCAL type of inquiry routine with an 'F77_' prefix
                                                                                 26
          CALL F77_GLOBAL_SIZE ( NX , DESCRX , 1 )
                                                                                 27
                                                                                 28
    ! Initialize elements of the array
                                                                                29
          DO J = LB2, UB2
                                                                                 30
            DO I = LB2, UB2
                                                                                 31
              X(I,J) = I + (J-1) * NX
                                                                                 32
            END DO
                                                                                 33
          END DO
                                                                                 34
                                                                                35
          END
                                                                                 36
                                                                                 37
                                                                                 38
          SUBROUTINE LOCAL2(N,X,R)
                                                                                 39
                                                                                 40
    ! Here, the correspondence to the global indices is not important
                                                                                41
    ! Only the total size of the subgrid is passed in
                                                                                 42
          REAL X(N)
                                                                                 43
                                                                                 44
          R = 0.
                                                                                 45
          DO I = 1, N
                                                                                 46
            R = R + X(I)
                                                                                 47
          END DO
                                                                                 48
```

1 END 2 з 4 LAYOUT('HPF\_ARRAY') and PASS\_BY('HPF\_HANDLE') 11.6.3.25 This example performs only the initialization of the above example. It illustrates use of the 6 LAYOUT ('F77\_ARRAY') attribute to pass an HPF distributed array without remapping, as 7 well as use of PASS\_BY('HPF\_HANDLE') to pass an HPF-style descriptor or handle for use 8 in the F77\_LOCAL subgrid inquiry function. It also illustrates the addressing of data in 9 terms of "embedding arrays." 10 11 • HPF caller 1213 PROGRAM EXAMPLE 14 15INTEGER, PARAMETER :: NX = 100, NY = 100 16REAL, DIMENSION(NX,NY) :: Y 17!HPF\$ DISTRIBUTE(BLOCK, BLOCK) :: Y 18 19 ! Local subgrid parameters are declared per processor 20 ! for a rank-two array 21 INTEGER, DIMENSION(NUMBER\_OF\_PROCESSORS(),2) :: 22 & LB, UB, LB\_EMBED, UB\_EMBED 23 !HPF\$ DISTRIBUTE(BLOCK,\*) :: LB, UB, LB\_EMBED, UB\_EMBED  $^{24}$ 2526 ! Define interfaces 27 28 INTERFACE 29 30 EXTRINSIC(F77\_LOCAL) SUBROUTINE LOCAL1( 31 LB1, UB1, LB\_EMBED1, UB\_EMBED1, 32 & LB2, UB2, LB\_EMBED2, UB\_EMBED2, X, X\_DESC ) & 33 INTEGER, DIMENSION(:) :: 34 LB1, UB1, LB\_EMBED1, UB\_EMBED1, & 35 LB2, UB2, LB\_EMBED2, UB\_EMBED2 & 36 i By default, X is passed by reference 37 REAL, DIMENSION(:,:), LAYOUT('HPF\_ARRAY') :: X 38 i X\_DESC is passed by its descriptor or 'handle' 39 REAL, DIMENSION(:,:), LAYOUT('HPF\_ARRAY'), 40 & PASS\_BY('HPF\_HANDLE') :: X\_DESC 41 !HPF\$ DISTRIBUTE(BLOCK) :: LB1, UB1, LB\_EMBED1, UB\_EMBED1 42 !HPF\$ DISTRIBUTE(BLOCK) :: LB2, UB2, LB\_EMBED2, UB\_EMBED2 43 !HPF\$ DISTRIBUTE(BLOCK,BLOCK) :: X 44 END 45 46 END INTERFACE 47 48

```
! Initialize values
                                                                                      1
       ! ( Assume stride = 1 and no axis permutation )
                                                                                      2
                                                                                      3
             CALL HPF_SUBGRID_INFO( Y, IERR,
                                                                                      4
            & LB=LB, LB_EMBED=LB_EMBED,
                                                                                     5
            & UB=UB, UB_EMBED=UB_EMBED)
                                                                                      6
             IF (IERR.NE.O) STOP 'ERROR!'
                                                                                     8
             CALL LOCAL1(
                                                                                      9
            & LB(:,1), UB(:,1), LB_EMBED(:,1), UB_EMBED(:,1),
                                                                                     10
            & LB(:,2), UB(:,2), LB_EMBED(:,2), UB_EMBED(:,2), Y, Y)
                                                                                     11
                                                                                     12
             END
                                                                                     13
                                                                                     14
                                                                                     15
   • Fortran 77 callee
                                                                                     16
                                                                                     17
             SUBROUTINE LOCAL1(
                                                                                     18
            & LB1, UB1, LB_EMBED1, UB_EMBED1,
                                                                                     19
            & LB2, UB2, LB_EMBED2, UB_EMBED2, X, X_DESC )
                                                                                     20
                                                                                     21
       ! The subgrid has been passed in its 'embedded' form
                                                                                     22
             REAL X ( LB_EMBED1 : UB_EMBED1 , LB_EMBED2 : UB_EMBED2 )
                                                                                     23
                                                                                     ^{24}
       ! This argument is used only as input to inquiry functions
                                                                                     ^{25}
             INTEGER X_DESC
                                                                                     26
                                                                                     27
       ! Get the global extent of the first axis
                                                                                     28
       ! This is an HPF_LOCAL type of inquiry routine with an 'F77_' prefix
                                                                                     29
             CALL F77_GLOBAL_SIZE(NX,X_DESC,1)
                                                                                     30
                                                                                     31
       ! Otherwise, initialize elements of the array
                                                                                     32
       ! Loop only over actual array elements
                                                                                     33
             DO J = LB2, UB2
                                                                                     34
               DO I = LB2, UB2
                                                                                     35
                  X(I,J) = I + (J-1) * NX
                                                                                     36
               END DO
                                                                                     37
             END DO
                                                                                     38
                                                                                     39
             END
                                                                                     40
                                                                                     41
                                                                                     42
11.7
       The Extrinsic Library
                                                                                     43
Following are Fortran bindings for routines useful in intrinsic subprograms.
                                                                                     44
                                                                                     45
                                                                                     46
11.7.1 HPF Local Routine Library
                                                                                     47
Local HPF procedures can use any HPF intrinsic or library procedure.
                                                                                     48
```

Advice to implementors. The arguments to such procedures will be local arrays.
Depending on the implementation, the actual code for the intrinsic and library routines used by local HPF procedures may or may not be the same code used when called from global HPF code. (End of advice to implementors.)

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In addition, local library procedures GLOBAL\_ALIGNMENT, GLOBAL\_DISTRIBUTION, and GLOBAL\_TEMPLATE are provided to query the global mapping of an actual argument to an extrinsic function. Other local library procedures are provided to query the size, shape, and array bounds of an actual argument. These library procedures take as input the name of a dummy argument and return information on the corresponding global HPF actual argument. They may be invoked only by a local procedure that was directly invoked by global HPF code. If module facilities are available, they reside in a module called HPF\_LOCAL\_LIBRARY; a local routine that calls them should include the statement

13 14 15

#### USE HPF\_LOCAL\_LIBRARY

<sup>16</sup> or some functionally appropriate variant thereof.

<sup>17</sup> The HPF local routine library identifies each physical processor by an integer in <sup>18</sup> the range 0 to n - 1, where n is the value returned by the global HPF\_LIBRARY func-<sup>19</sup> tion NUMBER\_OF\_PROCESSORS. Processor identifiers are returned by ABSTRACT\_TO\_PHYSICAL, <sup>20</sup> which establishes the one-to-one correspondence between the abstract processors of an <sup>21</sup> HPF processors arrangement and the physical processors. Also, the local library function <sup>22</sup> MY\_PROCESSOR returns the identifier of the calling processor.

In all cases, when an argument of one of the procedures of the local HPF library is required to be a local dummy argument associated with a global HPF actual argument, such association is not considered to be transitive. That is, the local dummy argument must be a dummy argument of a procedure which was referenced from global HPF, not from another local subprogram.

28 29

30

# 11.7.1.1 Accessing Dummy Arguments by Blocks

The mapping of a global HPF array to the physical processors places one or more blocks, 31 which are groups of elements with consecutive indices, on each processor. The number 32 of blocks mapped to a processor is the product of the number of blocks of consecutive 33 indices in each dimension that are mapped to it. For example, a rank-one array X with 34 a CYCLIC(4) distribution will have blocks containing four elements, except for a possible 35 last block having  $1 + SIZE(X) \mod 4$  elements. On the other hand, if X is first aligned to a 36 template or an array having a CYCLIC(4) distribution, and a non-unit stride is employed (as 37 is !HPF\$ ALIGN X(I) WITH T(3\*I), then its blocks may have fewer than four elements. 38 In this case, when the align stride is three and the template has a block-cyclic distribution 39 with four template elements per block, the blocks of X have either one or two elements each. 40 If the align stride were five, then all blocks of **X** would have exactly one element, as template 41 42 blocks to which no array element is aligned are not counted in the reckoning of numbers of blocks. 43

The portion of a global array argument associated with a dummy argument in an HPF\_LOCAL subprogram may be accessed in a block-by-block fashion. Three of the local library routines, LOCAL\_BLKCNT, LOCAL\_LINDEX, and LOCAL\_UINDEX, allow easy access to the local storage of a particular block. Their use for this purpose is illustrated by the following example, in which the local data are initialized one block at a time:

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37 38 39

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43 44 45

46

```
EXTRINSIC(HPF_LOCAL) SUBROUTINE NEWKI_DONT_HEBLOCK(X)
REAL X(:,:,:)
INTEGER BL(3)
INTEGER, ALLOCATABLE LIND1(:), LIND2(:), LIND3(:)
INTEGER, ALLOCATABLE UIND1(:), UIND2(:), UIND3(:)
BL = LOCAL_BLKCNT(X)
ALLOCATE LIND1(BL(1))
ALLOCATE LIND2(BL(2))
ALLOCATE LIND3(BL(3))
ALLOCATE UIND1(BL(1))
ALLOCATE UIND2(BL(2))
ALLOCATE UIND3(BL(3))
LIND1 = LOCAL_LINDEX(X, DIM = 1)
UIND1 = LOCAL_UINDEX(X, DIM = 1)
LIND2 = LOCAL_LINDEX(X, DIM = 2)
UIND2 = LOCAL_UINDEX(X, DIM = 2)
LIND3 = LOCAL_LINDEX(X, DIM = 3)
UIND3 = LOCAL_UINDEX(X, DIM = 3)
DO IB1 = 1, BL(1)
  DO IB2 = 1, BL(2)
    DO IB3 = 1, BL(3)
      FORALL (I1 = LIND1(IB1) : UIND1(IB1), &
              I2 = LIND2(IB2) : UIND2(IB2), \&
              I3 = LIND3(IB3) : UIND3(IB3) ) &
                X(I1, I2, I3) = IB1 + 10*IB2 + 100*IB3
    ENDDO
  ENDDO
ENDDO
END SUBROUTINE NEWKI_DONT_HEBLOCK
```

### GLOBAL\_ALIGNMENT(ARRAY, ...)

This has the same interface and behavior as the HPF inquiry subroutine HPF\_ALIGNMENT, but it returns information about the *global* HPF array actual argument associated with the local dummy argument ARRAY, rather than returning information about the local array.

## GLOBAL\_DISTRIBUTION(ARRAY, ...)

This has the same interface and behavior as the HPF inquiry subroutine HPF\_DISTRIBUTION, 47 but it returns information about the *global* HPF array actual argument associated with the 48

1	local dummy argument ARRAY, rather than returning information about the local array.
2 3	GLOBAL_TEMPLATE(ARRAY,)
4 5 6 7	This has the same interface and behavior as the HPF inquiry subroutine HPF_TEMPLATE, but it returns information about the <i>global</i> HPF array actual argument associated with the local dummy argument ARRAY, rather than returning information about the local array.
8 9	GLOBAL_SHAPE(SOURCE)
10 11	<b>Description.</b> Returns the shape of the global HPF actual argument associated with an array or scalar dummy argument of an HPF_LOCAL procedure.
12 13	Class. Inquiry function.
14 15	Argument.
16 17 18 19	SOURCE may be of any type. It may be array valued or a scalar. It must be a dummy argument of an HPF_LOCAL procedure which is argument associated with a global HPF actual argument.
20 21 22	<b>Result Type, Type Parameter and Shape.</b> The result is a default integer array of rank one whose size is equal to the rank of SOURCE.
23 24	<b>Result Value.</b> The value of the result is the shape of the global actual argument associated with the actual argument associated with SOURCE.
25 26	Examples. Assuming A is declared by the statement
27	INTEGER A(3:100, 200)
28 29 30 31	and is argument associated with B, the value of GLOBAL_SHAPE(B) is $\begin{bmatrix} 98 & 200 \end{bmatrix}$ . If B is argument associated with the section, A(5:10, 10), the value of GLOBAL_SHAPE(B) is $\begin{bmatrix} 6 \end{bmatrix}$ .
32 33	GLOBAL_SIZE(ARRAY, DIM)
34 35	Optional argument. DIM
36 37 38	<b>Description.</b> Returns the extent along a specified dimension of the global HPF actual array argument associated with a dummy array argument of an HPF_LOCAL procedure.
39 40	Class. Inquiry function.
41 42	Argument.
43 44 45	ARRAY may be of any type. It must not be a scalar. It must be a dummy argument of an HPF_LOCAL procedure which is argument associated with a global HPF actual argument.
46 47 48	DIM (optional) must be scalar and of type integer with a value in the range $1 \leq DIM \leq n$ , where n is the rank of ARRAY.

Result Type, Type Parameter and Shape. Default integer scalar.

**Result Value.** The result has a value equal to the extent of dimension DIM of the actual argument associated with the actual argument associated with ARRAY or, if DIM is absent, the total number of elements in the actual argument associated with the actual argument associated with ARRAY.

Examples. Assuming A is declared by the statement

INTEGER A(3:10, 10)

and is argument associated with B, the value of  $GLOBAL\_SIZE(B, 1)$  is 8. If B is argument associated with the section, A(5:10, 2:4), the value of  $GLOBAL\_SIZE(B)$  is 18.

# ABSTRACT\_TO\_PHYSICAL(ARRAY, INDEX, PROC)

**Description.** Returns processor identification for the physical processor associated with a specified abstract processor relative to a global actual argument array.

Class. Subroutine.

1
21
22
23
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# PHYSICAL\_TO\_ABSTRACT(ARRAY, PROC, INDEX)

**Description.** Returns coordinates for an abstract processor, relative to a global actual argument array, corresponding to a specified physical processor.

1 2 3	PROC	must be scalar and of type default integer. It is an INTENT(IN) argument. It contains an identifying value for a physical processor.
4 5 7 8 9 10 11 12 13	INDEX	must be a rank-1 integer array. It is an INTENT(OUT) ar- gument. The size of INDEX must equal the rank of the processor arrangement onto which the global HPF array is mapped. INDEX receives the coordinates within this processors arrangement of the abstract processor associ- ated with the physical processor specified by PROC. The value of the $i^{th}$ element will be in the range 1 to $e_i$ , where $e_i$ is the extent of the $i^{th}$ dimension of the processors ar- rangement.
14 15 16 17	between abstract processors an is one-to-many an equivalent, s	d only on systems where there is a one-to-one correspondence d physical processors. On systems where this correspondence system-dependent procedure should be provided.
18	LOCAL_TO_GLOBAL(A	RRAY, L_INDEX, G_INDEX)
19	<b>Description</b> . Converts	s a set of local coordinates within a local dummy array to an
20 21		oordinates within the associated global HPF actual argument
22	array.	
23 24	Class. Subroutine.	
25	Arguments.	
26 27 28 29	ARRAY	may be of any type; it must be a dummy array that is associated with a global HPF array actual argument. It is an INTENT(IN) argument.
30 31 32 33 34 35	L_INDEX	must be a rank-1 integer array whose size is equal to the rank of ARRAY. It is an INTENT(IN) argument. It contains the coordinates of an element within the local dummy array ARRAY. The value of the $i^{th}$ element must be in the range 1 to $e_i$ , where $e_i$ is the extent of the $i^{th}$ dimension of ARRAY.
36	<b>G_INDEX</b>	must be a rank-1 integer array whose size is equal to
37		the rank of ARRAY. It is an INTENT(OUT) argument. It re-
38 39		ceives the coordinates within the global HPF array actual
40		argument of the element identified within the local array by L_INDEX. The value of the $i^{th}$ element will be in the
41		by L_INDEX. The value of the $i^{th}$ element will be in the range 1 to $e_i$ , where $e_i$ is the extent of the $i^{th}$ dimension
42		of the global HPF actual argument array associated with
43		ARRAY.
44		
45 46		RRAY, G_INDEX, L_INDEX,
40	LOCAL, NCOPIES, PRO	DCS)
48	Optional arguments.	L_INDEX, LOCAL, NCOPIES, PROCS

**Description.** Converts a set of global coordinates within a global HPF actual argument array to an equivalent set of local coordinates within the associated local dummy array.

Class. Subroutine.

#### Arguments.

ARRAY	may be of any type; it must be a dummy array that is associated with a global HPF array actual argument. It is an INTENT(IN) argument.	8 9 10 11
G_INDEX	must be a rank-1 integer array whose size is equal to the rank of ARRAY. It is an INTENT(IN) argument. It contains the coordinates of an element within the global HPF array actual argument associated with the local dummy array ARRAY. The value of the $i^{th}$ element must be in the range 1 to $e_i$ , where $e_i$ is the extent of the $i^{th}$ dimension of the global HPF actual argument array associated with ARRAY.	12 13 14 15 16 17 18 19
L_INDEX (optional)	must be a rank-1 integer array whose size is equal to the rank of ARRAY. It is an INTENT(OUT) argument. It receives the coordinates within a local dummy array of the element identified within the global actual argument array by G_INDEX. (These coordinates are identical on any processor that holds a copy of the identified global array element.)	20 21 22 23 24 25 26 27
	The value of the $i^{th}$ element will be in the range 1 to $e_i$ , where $e_i$ is the extent of the $i^{th}$ dimension of ARRAY.	28 29
LOCAL (optional)	must be scalar and of type LOGICAL. It is an INTENT(OUT) argument. It is set to .TRUE. if the local array contains a copy of the global array element and to .FALSE. otherwise.	30 31 32 33
NCOPIES (optional)	must be scalar and of type integer. It is an INTENT(OUT) argument. It is set to the number of processors that hold a copy of the identified element of the global actual array.	34 35 36 37
PROCS (optional)	must be a rank-1 integer array whose size is at least the number of processors that hold copies of the identified element of the global actual array. The identifying num- bers of those processors are placed in <b>PROCS</b> . The order in which they appear is implementation dependent.	38 39 40 41 42 43

# MY\_PROCESSOR()

Description. Returns the identifying number of the calling physical processor.

**Class.** Pure function.

**Result Type, Type Parameter, and Shape.** The result is scalar and of type 1 default integer. 2 3 **Result Value.** Returns the identifying number of the physical processor from which 4 the call is made. This value is in the range  $0 \leq MY_PROCESSOR < n - 1$  where n is 5 the value returned by NUMBER\_OF\_PROCESSORS. 6 7 LOCAL\_BLKCNT(ARRAY, DIM, PROC) 8 9 Optional arguments. DIM, PROC. 10  $1\,1$ **Description**. Returns the number of blocks of elements in each dimension, or of a 12specific dimension of the array on a given processor. 13 14 **Class.** Pure function. 1516Arguments. 17ARRAY may be of any type; it must be a dummy array that is 18 associated with a global HPF array actual argument. 19 20 **DIM** (optional) must be scalar and of type integer with a value in the 21 range  $1 \leq \text{DIM} \leq n$ , where n is the rank of ARRAY. The 22 corresponding actual argument must not be an optional 23 dummy argument.  $^{24}$ **PROC** (optional) must be scalar and of type integer. It must be a valid 25processor number. 26 27 **Result Type, Type Parameter, and Shape.** The result is of type default integer. 28 It is scalar if DIM is present; otherwise the result is an array of rank one and size n, 29 where n is the rank of ARRAY. 30 31 Result Value. 32 33 The value of LOCAL\_BLKCNT(ARRAY, DIM, PROC) is the number of blocks Case (i): 34 of the ultimate align target of ARRAY in dimension DIM that are mapped 35 to processor PROC and which have at least one element of ARRAY aligned 36 to them. 37 Case (ii): LOCAL\_BLKCNT(ARRAY, DIM) returns the same value as 38 LOCAL\_BLKCNT(ARRAY, DIM, PROC=MY\_PROCESSOR()). 39 Case (iii): LOCAL\_BLKCNT(ARRAY) has a value whose  $i^{th}$  component is equal to 40 LOCAL\_BLKCNT(ARRAY, i), for i = 1, ..., n, where n is the rank of ARRAY. 41 42 **Examples.** Given the declarations 43 44 REAL A(20,20), B(10) 45 TEMPLATE T(100, 100)46 !HPF\$ !HPF\$ ALIGN B(J) WITH A(\*,J) 47ALIGN A(I,J) WITH T(3\*I, 2\*J) !HPF\$ 48

```
!HPF$ PROCESSORS PR(5,5)
!HPF$ DISTRIBUTE T(CYCLIC(3), CYCLIC(3)) ONTO PR
!HPF$ CALL LOCAL_COMPUTE(A, B)
...
EXTRINSIC(HPF_LOCAL) SUBROUTINE LOCAL_COMPUTE(X, Y)
USE HPF_LOCAL_LIBRARY
REAL X(:,:), Y(:)
INTEGER NBY(1), NBX(2)
NBX = LOCAL_BLKCNT(X)
NBY = LOCAL_BLKCNT(Y)
```

the values returned on the physical processor corresponding to PR(2,4) in NBX is  $\begin{bmatrix} 4 & 3 \end{bmatrix}$  and in NBY is  $\begin{bmatrix} 1 \end{bmatrix}$ .

# LOCAL\_LINDEX(ARRAY, DIM, PROC)

<b>Optional argument.</b> PI
------------------------------

**Description.** Returns the lowest local index of all blocks of an array dummy argument in a given dimension on a processor.

Class. Pure function.

#### Arguments.

		27
ARRAY	may be of any type; it must be a dummy array that is	28
	associated with a global HPF array actual argument.	29
DIM	must be scalar and of type integer with a value in the	30
	range $1 < \text{DIM} < n$ , where n is the rank of ARRAY.	31
		32
$\texttt{PROC} \ (\text{optional})$	must be scalar and of type integer. It must be a valid	33
	processor number.	34

**Result Type, Type Parameter, and Shape.** The result is a rank-one array of type default integer and size b, where b is the value returned by LOCAL\_BLKCNT(ARRAY, DIM [, PROC])

#### Result Value.

- Case (i): The value of LOCAL\_LINDEX(ARRAY, DIM, PROC) has a value whose  $i^{th}$  component is the local index of the first element of the  $i^{th}$  block in dimension DIM of ARRAY on processor PROC. The value of the  $i^{th}$  element will be in the range 1 to  $e_i$ , where  $e_i$  is the extent of the  $i^{th}$  dimension of ARRAY.
- Case (ii): LOCAL\_LINDEX(ARRAY, DIM) returns the same value as LOCAL\_LINDEX(ARRAY, DIM, PROC=MY\_PROCESSOR()).

1 2 3	<b>Examples.</b> With the same declarations as in the example under LOCAL_BLKCNT, on the physical processor corresponding to PR(2,4) the value returned by LOCAL_LINDEX(X, DIM=1) is $\begin{bmatrix} 1 & 2 & 3 & 4 \end{bmatrix}$ ;				
4 5	the value of LOCAL_LINDEX(X, DIM=2) is $\begin{bmatrix} 1 & 3 & 4 \end{bmatrix}$ .				
6 7	LOCAL_UINDEX(ARRAY, DIM, PROC)				
8 9	Optional argument. PROC.				
10 11	<b>Description.</b> Returns the highest local index of all blocks of an array dummy argument in a given dimension on a processor.				
12 13	Class. Pure function.				
14	Arguments.				
15 16 17	ARRAY may be of any type; it must be a dummy array that is associated with a global HPF array actual argument.				
18 19	DIM must be scalar and of type integer with a value in the range $1 \leq \text{DIM} \leq n$ , where n is the rank of ARRAY.				
20 21 22	PROC (optional) must be scalar and of type integer. It must be a valid processor number.				
23 24 25	<b>Result Type, Type Parameter, and Shape.</b> The result is a rank-one array of type default integer and size $b$ , where $b$ is the value returned by LOCAL_BLKCNT(ARRAY, DIM [, PROC])				
26 27	Result Value.				
28 29 30 31 32 33	Case (i): The value of LOCAL_UINDEX(ARRAY, DIM, PROC) has a value whose $i^{th}$ component is the local index of the last element of the $i^{th}$ block in dimension DIM of ARRAY on processor PROC. The value of the $i^{th}$ element will be in the range 1 to $e_i$ , where $e_i$ is the extent of the $i^{th}$ dimension of ARRAY.				
34 35	Case (ii): LOCAL_UINDEX(ARRAY, DIM) returns the same value as LOCAL_UINDEX(ARRAY, DIM, PROC=MY_PROCESSOR()).				
36 37 38 39	<b>Examples.</b> With the same declarations as in the example under LOCAL_BLKCNT, on the physical processor corresponding to PR(2,4) the value returned by LOCAL_UINDEX(X, DIM=1) is $\begin{bmatrix} 1 & 2 & 3 & 4 \end{bmatrix}$ ;				
40 41	the value of LOCAL_UINDEX(X, DIM=2) is $\begin{bmatrix} 2 & 3 & 4 \end{bmatrix}$ .				
42 43	11.7.2 Library Access from Serial Extrinsics				
44 45 46	A SERIAL subprogram may contain references to any HPF_LIBRARY procedure or HPF intrin- sic function, except HPF_ALIGNMENT, HPF_DISTRIBUTION or HPF_TEMPLATE. Within a SERIAL scope the HPF_LOCAL_LIBRARY module must not be used. References to the intrinsic functions NUMBER_OF_PROCESSORS and PROCESSORS_SHAPE				
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will return the same value as if the function reference appeared in global HPF.

### SECTION 11. APPROVED EXTENSIONS FOR HPF EXTRINSICS

# Section 12

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# Approved Extensions to the HPF Intrinsic and Library Procedures

This chapter describes intrinsic and library routines that have been approved as extensions to HPF Version 2.0.

The extended intrinsic procedures include a transpose function that generalizes the Fortran TRANSPOSE intrinsic function. Certain algorithms require access to multidimensional arrays along different axes. In modern machines, it will usually be best to make the array axis along which an inner loop runs the first axis, so that in local memory the elements will be contiguous. A generalized transpose is required to do this data rearrangement, which is not simply a data remapping. In many cases, the result of the transpose will be assigned to a variable whose first axis is distributed with a *dist-format* of (\*).

For this sort of operation, **FORALL** is adequate when the rank and the particular set of axes to be exchanged are known; for example:

```
    FORALL(I1 = 1:SIZE(ARRAY,1))
    FORALL(I2 = 1:SIZE(ARRAY,2))
    FORALL(I3 = 1:SIZE(ARRAY,3))
    RESULT(I3,I1,I2) = ARRAY(I1,I2,I3)
    ENDFORALL
    ENDFORALL
    ENDFORALL
    ENDFORALL
```

If, however, the relation between input and result axes is to be variable, FORALL is an inconvenient idiom. Thus we have generalized the TRANSPOSE intrinsic function, allowing as arguments an input array (which is to be transposed) of any nonzero rank, and an integer rank-one array (giving the axis permutation) whose size is the rank of the first input array.

The default value for the order argument makes this an extension of the existing Fortran one-argument **TRANSPOSE** function.

Two new intrinsic inquiry functions, ACTIVE\_NUM\_PROCS and ACTIVE\_PROCS\_SHAPE are useful for determining the size and the shape of the processor subset executing the program, as modified by ON constructs.

The extended library consists of mapping inquiry subroutines. Extended versions of HPF\_ALIGNMENT and HPF\_TEMPLATE allow an additional, optional, DYNAMIC output argument. This allows a program to determine whether an object, or its align ultimate target, has the DYNAMIC attribute. There is a new version of HPF\_DISTRIBUTION, and two new mapping inquiry subroutines that are especially useful for determining mappings produced by the general block and indirect distribution forms.

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### **12.1** Specifications of Extended Intrinsic Procedures

### ACTIVE\_NUM\_PROCS(DIM)

**Optional Argument.** DIM

**Description.** Returns the total number of processors currently executing the program or the number of processors currently executing the program along a specified dimension of the processor array, as determined by the innermost ON block.

Class. Processors inquiry function.

#### Arguments.

DIM (optional)	must be scalar and of type integer with a value in the
	range $1 \leq \text{DIM} \leq n$ where n is the rank of the processor
	array.

#### Result Type, Type Parameter, and Shape. Default integer scalar.

**Result Value.** The result has a value equal to the extent of dimension DIM of the processor array determined by the innermost containing ON block or, if DIM is absent, the total number of elements of this processor array. The result is always greater than zero. Outside of any ON block, the result is the same as that returned by NUMBER\_OF\_PROCESSORS().

**Examples.** The program fragment

INTEGER X(16, 3)
!hpf\$ TEMPLATE T(16, 8)
<pre>!hpf\$ PROCESSORS PROCS(4, 4)</pre>
!hpf\$ ALIGN X(I, J) WITH T(I, 3∗J−1)
<pre>!hpf\$ DISTRIBUTE T(CYCLIC(2), BLOCK) ONTO PROCS</pre>
<pre>!hpf\$ ON (PROCS(:,:)) BEGIN</pre>
<pre>!hpf\$ ON HOME(X(2:12:10, :)) BEGIN</pre>
PRINT *, ACTIVE_NUM_PROCS()
PRINT *, ACTIVE_NUM_PROCS(DIM=1)
PRINT *, ACTIVE_NUM_PROCS(DIM=2)
hpf\$ END ON
hpf\$ END ON

prints 6, 2 and 3 regardless of the size or shape of the hardware processor array on which the program is running,

1	ACTIVE_PROCS_SHAPE()		
2 3 4	<b>Description.</b> Returns the shape of the currently active processor array, as determined by the innermost $ON$ block.		
5 6	Class. Processors inquiry function.		
7 8	Arguments. None		
9 10 11 12	<b>Result Type, Type Parameter, and Shape.</b> The result is a default integer array of rank one whose size is equal to the rank of the processor array determined by the innermost containing <b>ON</b> block.		
13 14 15 16	<b>Result Value.</b> The value of the result is the shape of the processor array determined by the innermost containing ON block. Outside of any ON block, the result is the same as that returned by PROCESSORS_SHAPE().		
17 18	<b>Examples.</b> The program fragment		
19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 33	<pre>INTEGER X(16, 3) !hpf\$ TEMPLATE T(16, 8) !hpf\$ TEMPLATE T(16, 8) !hpf\$ PROCESSORS PROCS(4, 4) !hpf\$ ALIGN X(I, J) WITH T(I, 3*J-1) !hpf\$ DISTRIBUTE T(CYCLIC(2), BLOCK) ONTO PROCS !hpf\$ ON (PROCS(:,:)) BEGIN PRINT *, ACTIVE_PROCS_SHAPE() !hpf\$ ON HOME(X(2:12:10, :)) BEGIN PRINT *, ACTIVE_PROCS_SHAPE() !hpf\$ END ON !hpf\$ END ON prints 4, 4 and 2,3 regardless of the size or shape of the hardware processor array on which the program is running,</pre>		
35 36	TRANSPOSE(ARRAY,ORDER)		
37	Optional Argument. ORDER		
38 39	<b>Description.</b> Permute the axes (a generalized transpose) of an array.		
40 41	Class. Transformational function.		
42 43	Arguments.		
44 45	ARRAY may be of any type, and must be array valued.		
46 47 48	<b>ORDER</b> (optional) must be of type integer, rank one, and of size equal to the rank of <b>ARRAY</b> . Its elements must be a permutation of $(1, 2,, n)$ , where n is <b>RANK</b> ( <b>ARRAY</b> ).		

**Result Type, Type Parameters, and Shape.** The result is an array of the same rank, type, and type parameters as ARRAY. Its shape satisfies the relation RS(ORDER) == AS, where RS is the shape of the result and AS is SHAPE(ARRAY). If ORDER is absent, it defaults to (n, n - 1, ..., 1), where n is RANK(ARRAY).

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**Result value.** Element  $(j_1, j_2, \ldots, j_n)$  of the result is ARRAY $(j_{order(1)}, j_{order(2)}, \ldots, j_{order(n)})$ .

**Examples.** For an array of rank two, **TRANSPOSE(ARRAY)** is the usual matrix transpose.

If ARRAY has shape  $\begin{bmatrix} 1 & 2 & 3 \end{bmatrix}$  and ARRAY(1,:,:) is  $\begin{bmatrix} 111 & 112 & 113 \\ 121 & 122 & 123 \end{bmatrix}$ and ORDER is  $\begin{bmatrix} 3 & 1 & 2 \end{bmatrix}$  then the shape of the result is  $\begin{bmatrix} 2 & 3 & 1 \end{bmatrix}$ ; if R is the result then R(:,:,1) is equal to ARRAY(1,:,:). The rule is that axis *i* of ARRAY becomes axis ORDER(i) of the result.

### 12.2 Specifications of Extended Library Procedures

## HPF\_ALIGNMENT(ALIGNEE, LB, UB, STRIDE, AXIS\_MAP, IDEN-TITY\_MAP, DYNAMIC, NCOPIES)

**Optional Arguments.** LB, UB, STRIDE, AXIS\_MAP, IDENTITY\_MAP, DYNAMIC, NCOPIES

**Description.** Returns information regarding the correspondence of a variable and the *align-target* (array or template) to which it is ultimately aligned.

Class. Mapping inquiry subroutine.

#### Arguments.

ALIGNEE	may be of any type. It may be scalar or array valued. It must not be an assumed-size array. It must not be a pointer that is disassociated or an allocatable array that is not allocated. It is an INTENT (IN) argument.	32 33 34 35
	If ALIGNEE has the pointer attribute, information about the alignment of its target is returned. The target must not be an assumed-size dummy argument or a section of an assumed-size dummy argument.	36 37 38 39 40
LB (optional)	must be of type default integer and of rank one. Its size must be at least equal to the rank of ALIGNEE. It is an INTENT (OUT) argument. The first element of the $i^{th}$ axis of ALIGNEE is ultimately aligned to the LB(i) <sup>th</sup> align-target element along the axis of the align-target associated with the $i^{th}$ axis of ALIGNEE. If the $i^{th}$ axis of ALIGNEE is a collapsed axis, LB(i) is implementation dependent.	41 42 43 44 45 46 47 48

1 2 3 4 5 6 7 8	UB (optional)	must be of type default integer and of rank one. Its size must be at least equal to the rank of ALIGNEE. It is an INTENT (OUT) argument. The last element of the $i^{th}$ axis of ALIGNEE is ultimately aligned to the UB(i) <sup>th</sup> align-target element along the axis of the align-target associated with the $i^{th}$ axis of ALIGNEE. If the $i^{th}$ axis of ALIGNEE is a collapsed axis, UB(i) is implementation dependent.
9 10 11 12 13 14 15	STRIDE (optional)	must be of type default integer and of rank one. Its size must be at least equal to the rank of ALIGNEE. It is an INTENT (OUT) argument. The i <sup>th</sup> element of STRIDE is set to the stride used in aligning the elements of ALIGNEE along its i <sup>th</sup> axis. If the i <sup>th</sup> axis of ALIGNEE is a collapsed axis, STRIDE(i) is zero.
16 17 18 19 20 21	AXIS_MAP (optional)	must be of type default integer and of rank one. Its size must be at least equal to the rank of ALIGNEE. It is an INTENT (OUT) argument. The i <sup>th</sup> element of AXIS_MAP is set to the <i>align-target</i> axis associated with the i <sup>th</sup> axis of ALIGNEE. If the i <sup>th</sup> axis of ALIGNEE is a collapsed axis, AXIS_MAP(i) is 0.
22 23 24 25 26 27 28 29 30 31	IDENTITY_MAP (optional)	must be scalar and of type default logical. It is an INTENT (OUT) argument. It is set to true if the ultimate <i>align-target</i> associated with ALIGNEE has a shape identical to ALIGNEE, the axes are mapped using the identity per- mutation, and the strides are all positive (and therefore equal to 1, because of the shape constraint); otherwise it is set to false. If a variable has not appeared as an <i>alignee</i> in an ALIGN or REALIGN directive, and does not have the INHERIT attribute, then IDENTITY_MAP must be true; it can be true in other circumstances as well.
32 33 34 35	DYNAMIC (optional)	must be scalar and of type default logical. It is an INTENT (OUT) argument. It is set to true if ALIGNEE has the DYNAMIC attribute; otherwise it is set to false.
36 37 38 39	NCOPIES (optional)	must be scalar and of type default integer. It is an INTENT (OUT) argument. It is set to the number of copies of ALIGNEE that are ultimately aligned to <i>align-target</i> . For a non-replicated variable, it is set to one.
40 41 42		If ALIGNEE is scalar, then no elements of LB, UB, STRIDE, or AXIS_MAP are set.
43 44	Examples.	
45 46 47	REAL PI = 3.1415927 DIMENSION A(10,10), HPF\$ TEMPLATE T(40,20)	3(20,30),C(20,40,10),D(40)

HPF\$ DYNAMIC A 48

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!HPF\$ ALIGN A(I,:) WITH T(1+3\*I,2:20:2) !HPF\$ ALIGN C(I,\*,J) WITH T(J,21-I) !HPF\$ ALIGN D(I) WITH T(I,4) !HPF\$ PROCESSORS PROCS(4,2), SCALARPROC !HPF\$ DISTRIBUTE T(BLOCK,BLOCK) ONTO PROCS !HPF\$ DISTRIBUTE B(CYCLIC,BLOCK) ONTO PROCS !HPF\$ DISTRIBUTE ONTO SCALARPROC :: PI

Arguments.

assuming that the actual mappings are as the directives specify, the results of calling HPF\_ALIGNMENT are:

	А	В	С	D
LB	[4, 2]	[1, 1]	[20, N/A, 1]	[1]
UB	[31, 20]	[20,30]	[1, N/A, 10]	[40]
STRIDE	[3, 2]	[1, 1]	[-1,0,1]	[1]
AXIS_MAP	[1, 2]	[1, 2]	[2,0,1]	[1]
IDENTITY_MAP	false	true	false	false
DYNAMIC	true	false	false	false
NCOPIES	1	1	1	1

where "N/A" denotes a implementation-dependent result. To illustrate the use of NCOPIES, consider:

LOGICAL BOZO(20,20), RONALD_MCDONALD(20)	22
HPF\$ TEMPLATE EMMETT_KELLY(100,100)	23
HPF\$ ALIGN RONALD_MCDONALD(I) WITH BOZO(I,*)	24
HPF\$ ALIGN BOZO(J,K) WITH EMMETT_KELLY(J,5*K)	25
.mrt kita bolo(3,k) with indif_Abble(3,0+k)	26
Then CALL HPF_ALIGNMENT(RONALD_MCDONALD, NCOPIES = NC) sets NC to 20. Now con-	27
sider:	28
	29
LOGICAL BOZO(20,20), RONALD_MCDONALD(20)	30
!HPF\$ TEMPLATE WILLIE_WHISTLE(100)	31
!HPF\$ ALIGN RONALD_MCDONALD(I) WITH BOZO(I,*)	32
!HPF\$ ALIGN BOZO(J,*) WITH WILLIE_WHISTLE(5*J)	33
Then CALL HPF_ALIGNMENT(RONALD_MCDONALD, NCOPIES = NC) sets NC to one.	34
Then CALL HFF_ALIGNMENT (NONALD_NODONALD, NCOFILS - NC) sets NC to one.	35
HDE DIGEDIDIELON/DIGEDIDIEDE AVIG EVDE AVIG INFO	36
HPF_DISTRIBUTION(DISTRIBUTEE, AXIS_TYPE, AXIS_INFO,	
PROCESSORS_RANK, PROCESSORS_SHAPE, PLB, PUB, PSTRIDE,	
LOW_SHADOW, HIGH_SHADOW)	39
<b>Optional Arguments.</b> AXIS_TYPE, AXIS_INFO, PROCESSORS_RANK,	40
PROCESSORS_SHAPE, PLB, PUB, PSTRIDE, LOW_SHADOW, HIGH_SHADOW.	41
	42
<b>Description</b> . The HPF_DISTRIBUTION subroutine returns information regarding the	43
distribution of the ultimate <i>align-target</i> associated with a variable.	44
	45
Class. Mapping inquiry subroutine.	46
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1 2 3 4	DISTRIBUTEE	may be of any type. It may be scalar or array valued. It must not be sequential. It must not be a pointer that is disassociated or an allocatable array that is not allocated. It is an INTENT (IN) argument.
5 6 7 8 9 10 11 12 13 14 15 16 17	AXIS_TYPE (optional)	must be a rank one array of type default character. It may be of any length, although it must be of length at least 9 in order to contain the complete value. Its elements are set to the values below as if by a char- acter intrinsic assignment statement. Its size must be at least equal to the rank of the <i>align-target</i> to which DISTRIBUTEE is ultimately aligned; this is the value re- turned by HPF_TEMPLATE in TEMPLATE_RANK. It is an INTENT (OUT) argument. Its i <sup>th</sup> element contains infor- mation on the distribution of the i <sup>th</sup> axis of that <i>align- target</i> . The following values are defined by HPF (imple- mentations may define other values):
18 19 20 21 22		<ul> <li>'BLOCK' The axis is distributed BLOCK. The corresponding element of AXIS_INFO contains the block size.</li> <li>'GEN_BLOCK' The axis is distributed BLOCK(array). The value of the corresponding element of AXIS_INFO is implementation dependent.</li> </ul>
23 24 25 26 27 28 29		<ul> <li>'COLLAPSED' The axis is collapsed (distributed with the "*" specification). The value of the corresponding element of AXIS_INFO is implementation dependent.</li> <li>'CYCLIC' The axis is distributed CYCLIC. The corresponding element of AXIS_INFO contains the block size.</li> </ul>
30 31 32 33 34		'INDIRECT' The axis is distributed INDIRECT(map-array). The value of the corre- sponding element of AXIS_INFO is implementation dependent.
35 36 37 38 39 40 41 42 43	AXIS_INFO (optional)	must be a rank one array of type default integer, and size at least equal to the rank of the <i>align-target</i> to which DISTRIBUTEE is ultimately aligned (which is returned by HPF_TEMPLATE in TEMPLATE_RANK). It is an INTENT (OUT) argument. The i <sup>th</sup> element of AXIS_INFO contains the block size in the block or cyclic distribution of the i <sup>th</sup> axis of the ultimate <i>align-target</i> of DISTRIBUTEE; if that axis is a collapsed axis, then the value is implementation de- pendent.
44 45 46 47 48	PROCESSORS_RANK (option	al) must be scalar and of type default integer. It is set to the rank of the processor arrangement onto which DISTRIBUTEE is distributed. It is an INTENT (OUT) ar- gument.

PROCESSORS_SHAPE (optional) must be a rank one array of type default integer and
of size at least equal to the value, $m$ , returned in PROCES-
SORS_RANK. It is an INTENT (OUT) argument. Its first $m$
elements are set to the shape of the processor arrange-
ment onto which <b>DISTRIBUTEE</b> is mapped. (It may be
necessary to call HPF_DISTRIBUTION twice, the first time
to obtain the value of PROCESSORS_RANK in order to allo-
cate PROCESSORS_SHAPE.)

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PLB (optional) must be a rank one array of type default integer and of size at least equal to the rank of the ultimate *aligntarget* of DISTRIBUTEE. It is an INTENT (OUT) argument. The i<sup>th</sup> element is set to the smallest processor index ONTO which the i<sup>th</sup> axis of the ultimate *align-target* of DISTRIBUTEE is mapped; if that axis is collapsed, then the corresponding element of PLB is implementation dependent. The value returned is in the range 1 to  $e_i$  where  $e_i$  is the extent of processor arrangement axis onto which the selected axis of the ultimate *align-target* of DISTRIBUTEE is mapped.

- **PUB** (optional) must be a rank one array of type default integer and  $^{21}$ of size at least equal to the rank of the ultimate align-22 target of DISTRIBUTEE. It is an INTENT (OUT) argument. 23 The i<sup>th</sup> element is set to the largest processor index ONTO  $^{24}$ which the i<sup>th</sup> axis of the ultimate *align-target* of DISTRI- $^{25}$ BUTEE is mapped; if that axis is collapsed, then the cor-26 responding element of PUB is implementation dependent.  $^{27}$ The value returned is in the range 1 to  $e_i$  where  $e_i$  is the 28 extent of processor arrangement axis onto which the se-29 lected axis of the ultimate *align-target* of **DISTRIBUTEE** is 30 mapped. 31
- PSTRIDE (optional) must be a rank one array of type default integer and of size at least equal to the rank of DISTRIBUTEE. It is an INTENT (OUT) argument. The i<sup>th</sup> element is set to the interprocessor stride in the ONTO clause with which the i<sup>th</sup> axis of DISTRIBUTEE is mapped; if that axis is collapsed, then the corresponding element of PSTRIDE is set to zero.

LOW\_SHADOW (optional) must be a rank one array of type default integer, and size 40 at least equal to the rank of the *align-target* to which 41 **DISTRIBUTEE** is ultimately aligned (which is returned 42 by HPF\_TEMPLATE in TEMPLATE\_RANK). It is an INTENT 43 (OUT) argument. The i<sup>th</sup> element of LOW\_SHADOW con-44 tains the low-side shadow width in the block or cyclic 45distribution of the i<sup>th</sup> axis of the ultimate *align-target* of  $^{46}$ DISTRIBUTEE; if that axis is a collapsed axis, then the 47 value is implementation dependent. 48

1 2 3 4 5 6 7 8 9 10 11	HIGH_SHADOW (optional) Example. Given the declassuming that the actual	at least equal to t DISTRIBUTEE is ut by HPF_TEMPLATE (OUT) argument. tains the high-side distribution of the DISTRIBUTEE; if th value is implement.	ample illustrating HPF_A	arget to which ch is returned is an INTENT H_SHADOW con- block or cyclic align-target of axis, then the LIGNMENT and	
12 13	HPF_DISTRIBUTION are:	mappings are as	the difectives speeny,	the results of	
13		А	В	PI	
15	AXIS_TYPE ['H	BLOCK', 'BLOCK']	['CYCLIC', 'BLOCK']		
16	AXIS_INFO	[10, 10]	[1, 15]	[]	
17	PROCESSORS_SHAPE	[4, 2]	[2,2]	[ ]	
18 19	PROCESSORS_RANK	2	2	0	
20	PLB PUB	[4, 1] [1, 2]	[2, 1] [3, 2]		
21	PSTRIDE	[-1, 1]	[1, 1]		
22				LJ	
23	HPF_TEMPLATE(ALIGN	EE, TEMPLAT	E_RANK, LB, UB,		
24	AXIS_TYPE, AXIS_INFO,	NUMBER_ALI	GNED, DYNAMI	C)	
25 26	<b>Optional Arguments.</b> L	B, UB, AXIS_TYPE, A	XIS_INFO, NUMBER_ALIG	NED,	
27	TEMPLATE_RANK, DYNAMIC	, , , , ,	,	,	
28	<b>Description</b> . The HPF_TE	MPLATE subroutine	returns information re-	parding the ul-	
29	timate <i>align-target</i> associa				
30	concerning the variable fr		4		
31 32	HPF_ALIGNMENT returns information from the variable's point of view.				
33	Class. Mapping inquiry subroutine.				
34					
35	Arguments.				
36	ALIGNEE	may be of any typ	be. It may be scalar or	array valued.	
37 38			assumed-size array. It		
39			associated or an allocata		
40			is an INTENT (IN) arg		
41			e pointer attribute, info		
42			s target is returned. The		
43		an assumed-size du	-size dummy argument	or a section of	
44				Te i sværsvæ	
45	TEMPLATE_RANK (optional)		of type default integer.		
46 47		-	It is set to the rank o can be different from t		
47			ollapsing and replicating		
-			mapping and replicating	· ·	

LB (optional)	must be of type default integer and of rank one. Its size	1
	must be at least equal to the rank of the <i>align-target</i> to	2
	which ALIGNEE is ultimately aligned; this is the value	3
	returned in TEMPLATE_RANK. It is an INTENT (OUT) argu-	4
	ment. The $i^{th}$ element of LB contains the declared <i>align</i> -	5
	$target$ lower bound for the $\mathrm{i}^{\mathrm{th}}$ template axis.	6

**UB** (optional) must be of type default integer and of rank one. Its size must be at least equal to the rank of the *align-target* to which ALIGNEE is ultimately aligned; this is the value returned in TEMPLATE\_RANK. It is an INTENT (OUT) argument. The i<sup>th</sup> element of UB contains the declared *aligntarget* upper bound for the i<sup>th</sup> template axis.

- must be a rank one array of type default character. It AXIS\_TYPE (optional) 14may be of any length, although it must be of length 15at least 10 in order to contain the complete value. Its 16elements are set to the values below as if by a char-17acter intrinsic assignment statement. Its size must be 18 at least equal to the rank of the *align-target* to which 19ALIGNEE is ultimately aligned; this is the value returned 20 in TEMPLATE\_RANK. It is an INTENT (OUT) argument. The  $^{21}$ i<sup>th</sup> element of AXIS\_TYPE contains information about the 22 i<sup>th</sup> axis of the *align-target*. The following values are de-23 fined by HPF (implementations may define other values):  $^{24}$ 
  - $^{25}$ 'NORMAL' The *align-target* axis has an axis of ALIGNEE 26 aligned to it. For elements of AXIS\_TYPE assigned  $^{27}$ this value, the corresponding element of AXIS\_INFO 28 is set to the number of the axis of ALIGNEE aligned 29 to this *align-target* axis. 30
  - 'REPLICATED' ALIGNEE is replicated along this align-tar-31 get axis. For elements of AXIS\_TYPE assigned this 32 value, the corresponding element of AXIS\_INFO is set 33 to the number of copies of ALIGNEE along this align-34 target axis. 35
  - 'SINGLE' ALIGNEE is aligned with one coordinate of the 36 align-target axis. For elements of AXIS\_TYPE assigned 37 this value, the corresponding element of AXIS\_INFO 38 is set to the *align-target* coordinate to which ALIGNEE 39 is aligned. 40

AXIS\_INFO (optional) must be of type default integer and of rank one. Its size must be at least equal to the rank of the *align-target* to which ALIGNEE is ultimately aligned; this is the value returned in TEMPLATE\_RANK. It is an INTENT (OUT) argument. See the description of AXIS\_TYPE above.

NUMBER\_ALIGNED (optional) must be scalar and of type default integer. It is an INTENT (OUT) argument. It is set to the total number 48

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1 2 3			variables tha	-	<i>-target.</i> This is the <i>align-target</i>
4 5 6 7	DYNAMIC (optional)	(OUT) argume	ent. It is set t	-	It is an INTENT <i>ilign-target</i> has vise.
8 9 10 11	<b>Example.</b> Given the dec assuming that the actual HPF_TEMPLATE are:		-	-	
12		А	С	D	
13	LB	[1, 1]		[1, 1]	
14	UB		[40, 20]	[40, 20]	
15	AXIS_TYPE		['NORMAL',		
16		-	-	'SINGLE']	
17	AXIS_INFO	[1, 2]	[3, 1]	[1, 4]	
18 19	NUMBER_ALIGNEI		3	3	
20	TEMPLATE_RANK	2 false	2 false	2 false	
21	DYNAMIC	laise	Talse	laise	
<sup>22</sup> <b>HDE</b>	-MAP_ARRAY(ARR	Αν τένισι	ATE DIM		
23 1117 I'				, MAF_ART	LAI)
24	<b>Description.</b> Returns t				bution of axis
25	TEMPLATE_DIM of the ultim	nate <i>align-targe</i>	t associated v	with ARRAY.	
26	Class Manning inquiny a	wheneuting			
27	<b>Class.</b> Mapping inquiry s	ubroutille.			
28 29	Arguments.				
30					
31	ARRAY				It must not be
32					is disassociated
33			v	at is not alloc	ated. It is an
34		INTENT(IN) a	rgument.		
35	TEMPLATE_DIM	must be scalar	and of type of	lefault integer.	Its value must
36					ate <i>align-target</i>
37		of ARRAY. It is	an INTENT()	[N) argument.	
38	MAP_ARRAY	must be of	type default	integer and	of rank one.
39		Its size must	be no sma	ller than the	extent of the
40 41		PROCESSORS_D	$IM^{th}$ axis of	the processor	s arrangement
41					ign-target asso-
43		ciated with AF	RAY. It is an	INTENT(OUT) a	argument.
44					e processor in-
45					ate <i>align-target</i>
46			0		apped. If axis
47					of ARRAY is col-
48		lapsed, then a	ll elements of	the result have	e the value one.

Example. Given the declarations

```
DIMENSION A(2)

!HPF$ TEMPLATE T(4,8)

!HPF$ ALIGN A(I,*) WITH T(2*I,5)

!HPF$ PROCESSORS PROCS(2,2)

!HPF$ DISTRIBUTE T(INDIRECT( (/1,2,2,1/) ), BLOCK( (/3,5/) )) ONTO PROCS
```

assuming that the actual mappings are as the directives specify, Then after calling HPF\_MAP\_ARRAY(A,TEMPLATE\_DIM=1, MAP\_ARRAY=M), M has the value  $\begin{bmatrix} 1 & 2 & 2 & 1 \end{bmatrix}$ . After calling HPF\_MAP\_ARRAY(A,TEMPLATE\_DIM=2, MAP\_ARRAY=M), M has the value  $\begin{bmatrix} 1 & 1 & 1 & 2 & 2 & 2 & 2 \end{bmatrix}$ .

# HPF\_NUMBER\_MAPPED(ARRAY, PROCESSORS\_DIM, NUMBER\_MAPPED)

**Description.** Returns the number of elements of the ultimate *align-target* of ARRAY mapped to the each element of axis PROCESSORS\_DIM of the processors arrangement onto which the ultimate *align-target* of ARRAY is distributed.

Class. Mapping inquiry subroutine.

Arguments.		23
ARRAY	may be of any type. It must not be scalar. It must not be sequential. It must not be a pointer that is disassociated or an allocatable array that is not allocated.	24 25 26 27
PROCESSORS_DIM	must be scalar and of type default integer. Its value must be between one and the rank of the processors arrange- ment onto which the ultimate <i>align-target</i> of <b>ARRAY</b> is distributed.	28 29 30 31 32
NUMBER_MAPPED	must be of type default integer and of rank one. Its size must be no smaller than the extent of axis PROCESSORS_DIM of the processors arrangement onto which the ultimate <i>align-target</i> of ARRAY is distributed. The i <sup>th</sup> element of NUMBER_MAPPED is set to the num- ber of elements of an axis of the ultimate <i>align-target</i> of ARRAY that are mapped to the i <sup>th</sup> processor of axis PROCESSORS_DIM of the processors arrangement onto which the ultimate <i>align-target</i> of ARRAY is distributed. If axis PROCESSORS_DIM of the processors arrangement onto which the ultimate <i>align-target</i> of ARRAY is distributed is associated with a BLOCK distributed axis, then MAP_ARRAY is set to the array of block sizes used to distribute that axis.	33 34 35 36 37 38 39 40 41 42 43 44 45 46

Example. Given the declarations

1 2

8

9

10

11

12 13 14

15

16 17

18

19

20 21

```
DIMENSION A(2,40)
1
          !HPF$ TEMPLATE T(4,8,4,16)
^{2}
          !HPF$ ALIGN A(I,*) WITH T(2*I, 5, *, *)
3
          !HPF$ PROCESSORS PROCS(2,2,3)
4
          !HPF$ DISTRIBUTE T(INDIRECT((/2,2,1,2/)), BLOCK((/3,5/)), *, BLOCK) &
5
          !HPF$
                          ONTO PROCS
6
7
       assuming that the actual mappings are as the directives specify, after calling
8
       HPF_NUMBER_MAPPED (A, PROCESSORS_DIM=1, NUMBER_MAPPED = M) M has the value 1 3 ;
9
      after calling HPF_NUMBER_MAPPED(A, PROCESSORS_DIM=2, NUMBER_MAPPED = M) M has the value \begin{bmatrix} 3 & 5 \end{bmatrix}; after calling HPF_NUMBER_MAPPED(A, PROCESSORS_DIM=3, NUMBER_MAPPED =
10
1\,1
12
      M) M has the value \begin{bmatrix} 6 & 6 & 4 \end{bmatrix}.
13
14
15
16
17
18
19
20
21
22
23
^{24}
^{25}
26
^{27}
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
^{46}
47
48
```

# Part IV

# Annexes

This major section organizes descriptions of the syntax and semantics of features of the High Performance Fortran language, version 2.0 (described n Parts I and II) and its approved extensions (described in Part III) for reference use. It is not a part of the HPF language specification proper.

# Annex A

15 16 17

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# Syntax Rules

This Appendix collects the formal syntax definitions of this High Performance Fortran Language Specification.

- A.2 Notati
  - Notation and Syntax
- A.2.2 Syntax of Directives

20 21	H201 hpf-a	lirective-line	is	directive-origin hpf-directive
22 23 24 25	H202 direc	ctive-origin	is or or	!HPF\$ CHPF\$ *HPF\$
26 27 28	H203 hpf-a	lirective	is or	specification- $directiveexecutable$ - $directive$
29 30 31 32 33 34 35	H204 spect	ification- $directive$	is or or or or or	processors-directive align-directive distribute-directive inherit-directive template-directive combined-directive sequence-directive
36 37	H205 exect	utable-directive	is	independent- $directive$
38 39	Constraint:	An <i>hpf-directive-line</i> can the same line.	not	be commentary following another statement on
40 41 42	Constraint:	A <i>specification-directive</i> appear.	nay	appear only where a <i>declaration-construct</i> may
43 44 45	Constraint:	An <i>executable-directive</i> n appear.	nay :	appear only where an <i>executable-construct</i> may
46 47	Constraint:			he rules of either Fortran free form (F95:3.3.1.1) nment lines, depending on the source form of the

surrounding Fortran source form in that program unit. (F95:3.3)

H206	$specification\hbox{-}directive\hbox{-}extended$	$\mathbf{is}$	processors- $directive$	1
		or	subset-directive	2
		or	align-directive	3
		or	distribute- $directive$	4
		or	inherit- $directive$	5
		or	template- $directive$	6
		or	combined- $directive$	7
		or	sequence- $directive$	8
		or	dynamic- $directive$	9
		or	range-directive	10
		or	shadow- $directive$	11
				12
H207	executable-directive-extended	is	independent- $directive$	13
		or	realign-directive	14
		or	redistribute-directive	15
		or	on-directive	16
		or	resident-directive	17
H208	executable-construct-extended	is	action-stmt	18
11208	executable-construct-extended		case-construct	19
		or or	do-construct	20
		or	if-construct	21
		or	where-construct	22
		or	on-construct	23
		or	resident-construct	24
		or	task-region-construct	25 26
		01		20
A.3	Data Mapping			28
11.0	Data mapping			29
A.3.2	2 Syntax of Data Alignme	$\mathbf{ent}$	and Distribution Directives	30
H301	combined- $directive$	is	$combined\-attribute\-list\::\ combined\-decl\-list$	31 32
H302	combined- $attribute$	is	ALIGN align-attribute-stuff	33
		or	DISTRIBUTE dist-attribute-stuff	34
		or	INHERIT	35
		or	TEMPLATE	36
		or	PROCESSORS	37
		or	DIMENSION ( $explicit$ -shape-spec-list )	38
H303	combined- $decl$	is	hpf-entity [ ( explicit-shape-spec-list ) ]	39
11000		or	object-name	40
TTOO (			·	41
H304	hpf- $entity$	is	processors-name	42
		or	template-name	43
Const	raint: The same kind of <i>combir</i>	ned-a	attribute must not appear more than once in a	44
	given <i>combined-directive</i> .		•••	45
a i	_		· · · · · · · · · · · · · · · · · · ·	46
Consti			pears in a <i>combined-directive</i> , any entity to which th the HPF <b>TEMPLATE</b> or <b>PROCESSORS</b> type spec-	47 48

1		ifier.		
2 3	A.3.3 Tł	ne DISTRIBUTE Dire	ecti	ve
4 5	H305 distri	bute-directive	is	DISTRIBUTE distributee dist-directive-stuff
6	H306 dist-a	lirective-stuff	is	dist-format-clause [ dist-onto-clause ]
7 8 9	H307 dist-a	uttribute-stuff	is or	dist-directive-stuff dist-onto-clause
10 11	H308 distri	butee	is or	object-name template-name
12 13 14	H309 dist-f	ormat-clause	is or or	( dist-format-list ) * ( dist-format-list ) *
15 16 17 18	H310 dist-f	ormat	is or or	BLOCK [ ( scalar-int-expr ) ] CYCLIC [ ( scalar-int-expr ) ] *
19	H311 dist-a	onto-clause	is	ONTO dist-target
20 21 22 23	H312 dist-t	arget	is or or	processors-name * processors-name *
24 25	Constraint:	An <i>object-name</i> mentione subobject designator or a		s a <i>distributee</i> must be a simple name and not a <i>nponent-name</i> .
26 27	Constraint:	An object-name mentione	ed as	a <i>distributee</i> may not appear as an <i>alignee</i> .
28 29	Constraint:	An <i>object-name</i> mention tribute.	ed a	as a $distributee \mbox{ may not have the POINTER}$ at-
30 31	Constraint:	An object-name mentione	d as	a <i>distributee</i> may not have the <b>TARGET</b> attribute.
32 33 34	Constraint:		n th	e dist-format-list (and its surrounding parenthe- tis case, the statement form of the directive is clause of "*" is present.
35 36 37	Constraint:	If a <i>dist-format-list</i> is spectrum to which it applies.	cified	l, its length must equal the rank of each <i>distribu</i> -
38 39 40	Constraint:			d a <i>dist-target</i> appear, the number of elements re not "*" must equal the rank of the specified
41 42 43	Constraint:			ot a <i>dist-format-list</i> , the rank of each <i>distributee</i> ecified processor arrangement.
44 45 46	Constraint:			se or the <i>dist-target</i> in a <b>DISTRIBUTE</b> directive <i>istributee</i> must be a dummy argument.
40 47 48	Constraint:	Any <i>scalar-int-expr</i> appear be a <i>specification-expr</i> .	aring	g in a $dist$ -format of a DISTRIBUTE directive must

A.3.4	The ALIGN Directive			1
H313	a lign-directive	is	ALIGN alignee align-directive-stuff	2 3
H314	a lign-directive-stuff	is	( $align-source-list$ ) $align-with-clause$	4
H315	align-attribute-stuff	is	[ ( align-source-list ) ] align-with-clause	5
H316	alignee	is	object-name	6 7
H317	align-source	is	, ,	8
11011		or	*	9
		or	align-dummy	10
H318	align- $dummy$	is	scalar- $int$ - $variable$	11 12
Const	raint: An <i>object-name</i> mention subobject designator or a		s an <i>alignee</i> must be a simple name and not a <i>nponent-name</i> .	13 14 15
Const	raint: An object-name mentione	d as	an <i>alignee</i> may not appear as a <i>distributee</i> .	16 17
Const	raint: An object-name mentione	d as	an <i>alignee</i> may not have the POINTER attribute.	18
Const	raint: An object-name mentione	d as	an <i>alignee</i> may not have the <b>TARGET</b> attribute.	19 20
Const	8		<i>ign-source-list</i> (and its surrounding parentheses) the statement form of the directive is not allowed.	21 22 23
Const	raint: If the <i>align-source-list</i> is p to which it applies.	orese	nt, its length must equal the rank of each alignee	24 25
Const	raint: An <i>align-dummy</i> must be	a n	amed variable.	26 27
Const	raint: An object may not have b	ooth	the INHERIT attribute and the ALIGN attribute.	28 29
H319	align-with-clause	is	WITH align-spec	30 31
H320	align-spec	is or	align-target [ ( align-subscript-list ) ] * align-target [ ( align-subscript-list ) ]	32 33
H321	align-target	is or	object-name template-name	34 35
H322	align-subscript	is or or or	int-expr align-subscript-use subscript-triplet *	36 37 38 39 40
H323	a lign-subscript-use	is or	[[ int-level-two-expr ] add-op ] align-add-operand align-subscript-use add-op int-add-operand	41 42 43
H324	align-add-operand	is or	[ int-add-operand * ] align-primary align-add-operand * int-mult-operand	44 45
H325	align-primary	is or	align-dummy ( align-subscript-use )	46 47 48

## A 2 4 The ALICN Direction

### A.3. DATA MAPPING

H326 int-a	dd-operand	is	add-operand
H327 int-m	ult-operand	$\mathbf{is}$	mult-operand
H328 int-le	vel-two-expr	is	level-2-expr
Constraint:	An <i>object-name</i> mention a subobject designator o		s an <i>align-target</i> must be a simple name and not <i>omponent-name</i> .
Constraint:	An align-target may not	have	the OPTIONAL attribute.
Constraint:	If the <i>align-spec</i> in an AL be a dummy argument.	IGN (	lirective begins with " $*$ " then every <i>alignee</i> must
Constraint:	In an <i>align-directive</i> any <i>mult-operand</i> must be a		<i>expr</i> , <i>int-level-two-expr</i> , <i>int-add-operand</i> or <i>int-</i> fication expression.
Constraint:	Any <i>subscript</i> or <i>stride</i> <i>align-directive</i> must be a		subscript-triplet that is an align-subscript in an cification expression.
Constraint:	Each align-dummy may	appe	ar at most once in an <i>align-subscript-list</i> .
Constraint:	An <i>align-subscript-use</i> e <i>align-dummy</i> .	xpres	ssion may contain at most one occurrence of an
Constraint:	where in the <i>align-spec</i> e of the grammar shown <i>subscript-use</i> only by sta	excep abov arting	used as an <i>align-dummy</i> may not appear any- t where explicitly permitted to appear by virtue e. Paraphrased, one may construct an <i>align- g</i> with an <i>align-dummy</i> and then doing additive t with integer specification expressions that con-
Constraint:	A subscript within an alignature dummy.	gn-sı	ubscript may not contain occurrences of any align-
Constraint:	An <i>int-add-operand</i> , <i>int</i> -integer.	-mult	<i>e-operand</i> , or <i>int-level-two-expr</i> must be of type
A.3.6 Tł	ne PROCESSORS Di	irect	ive
H329 proce	ssors- $directive$	is	PROCESSORS processors-decl-list
H330 proce	ssors-decl	is	<pre>processors-name [ ( explicit-shape-spec-list ) ]</pre>
A.3.7 Tł	ne TEMPLATE Dire	ctiv	e
H331 temp	late-directive	is	TEMPLATE template-decl-list
H332 temp	late-decl	is	template-name [ ( explicit-shape-spec-list ) ]

A.3.8 St	orage and Sequence A	Asso	ociation	1	
H333 seque	ence-directive	is or	SEQUENCE [ [ :: ] association-name-list ] NO SEQUENCE [ [ :: ] association-name-list ]	2 3 4	
H334 assoc	iation-name	is or	object-name / [ common-block-name ] /	4 5 6	
Constraint:	An object name or COMMO directive within any scop		ock name may appear at most once in a <i>sequence</i> - 1nit.	7 8 9	
Constraint:	Only one sequence direct same scoping unit.	ive v	with no association-name-list is permitted in the	10 11	
A.4 Dat	a Mapping in Subpro	gra	m Interfaces	12 13 14	
A.4.4 Al	ignment			15	
H401 inher	it-directive	is	INHERIT inheritee-list	16 17	
H402 inher		is	object-name	18	
11402 1111101	<i>ucc</i>	15	oojeer-name	19	
Constraint:	An <i>inheritee</i> must be a d	umn	ny argument.	20	
Constraint:	An <i>inheritee</i> must not be	an	alignee.	21 22	
	An <i>inheritee</i> must not be			23	
Constraint:	All <i>undernee</i> must not be	e a u	istributee.	24	
A.5 IND	EPENDENT and Re	elate	ed Directives	25 26	
A.5.1 Tł	ne INDEPENDENT I	Dire	ective	26 27 28	
A.5.1 Tł				26 27	
A.5.1 Tł	ne INDEPENDENT I vendent-directive	Dire	ective INDEPENDENT [ , <i>new-clause</i> ]	26 27 28 29 30 31	
<b>A.5.1 Th</b> H501 indep H502 new-e	ne INDEPENDENT I vendent-directive clause	Dire is is	ective INDEPENDENT [ , new-clause ] [ , reduction-clause ] NEW ( variable-name-list )	26 27 28 29 30 31 32	
A.5.1         Th           H501         indep           H502         new-e           H503         reduct	ne INDEPENDENT I vendent-directive clause ction-clause	Dire is is is	ective INDEPENDENT [ , new-clause ] [ , reduction-clause ] NEW ( variable-name-list ) REDUCTION ( reduction-variable-list )	26 27 28 29 30 31	
A.5.1         Th           H501         indep           H502         new-e           H503         reduct	ne INDEPENDENT I vendent-directive clause	Dire is is is is	ective INDEPENDENT [ , new-clause ] [ , reduction-clause ] NEW ( variable-name-list ) REDUCTION ( reduction-variable-list ) array-variable-name	26 27 28 29 30 31 32 33	
A.5.1         Th           H501         indep           H502         new-e           H503         reduct	ne INDEPENDENT I vendent-directive clause ction-clause	Dire is is is	ective INDEPENDENT [ , new-clause ] [ , reduction-clause ] NEW ( variable-name-list ) REDUCTION ( reduction-variable-list ) array-variable-name scalar-variable-name	26 27 28 29 30 31 32 33 34	
A.5.1     TI       H501     indep       H502     new-q       H503     reduct       H504     reduct	ne INDEPENDENT I tendent-directive clause etion-clause etion-variable	Dire is is is is or or	ective INDEPENDENT [ , new-clause ] [ , reduction-clause ] NEW ( variable-name-list ) REDUCTION ( reduction-variable-list ) array-variable-name scalar-variable-name structure-component	26 27 28 29 30 31 32 33 34 35	
A.5.1     TI       H501     indep       H502     new-q       H503     reduct       H504     reduct	ne INDEPENDENT I eendent-directive clause etion-clause etion-variable	Dire is is is or or ne fe	ective INDEPENDENT [ , new-clause ] [ , reduction-clause ] NEW ( variable-name-list ) REDUCTION ( reduction-variable-list ) array-variable-name scalar-variable-name structure-component ollowing an independent-directive must be a do-	26 27 28 29 30 31 32 33 34 35 36	
<b>A.5.1 Tl</b> H501 <i>indep</i> H502 <i>new-0</i> H503 <i>reduc</i> H504 <i>reduc</i> Constraint:	ne INDEPENDENT I bendent-directive clause ction-clause ction-variable The first non-comment li stmt, forall-stmt, or a for	Dire is is is or or ne f- all-c	Pective INDEPENDENT [ , new-clause ] [ , reduction-clause ] NEW ( variable-name-list ) REDUCTION ( reduction-variable-list ) array-variable-name scalar-variable-name structure-component ollowing an independent-directive must be a do- onstruct.	26 27 28 29 30 31 32 33 34 35 36 37 38 39 40	
<b>A.5.1 Tl</b> H501 <i>indep</i> H502 <i>new-0</i> H503 <i>reduc</i> H504 <i>reduc</i> Constraint:	<b>ne INDEPENDENT I</b> bendent-directive clause ction-clause ction-variable The first non-comment li <i>stmt</i> , <i>forall-stmt</i> , or a <i>for</i> If the first non-comment	Dire is is is or or all-c line	<pre>ective INDEPENDENT [ , new-clause ]   [ , reduction-clause ] NEW ( variable-name-list ) REDUCTION ( reduction-variable-list ) array-variable-name scalar-variable-name structure-component ollowing an independent-directive must be a do- onstruct. following an independent-directive is a do-stmt,</pre>	26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41	
<b>A.5.1 Tl</b> H501 <i>indep</i> H502 <i>new-0</i> H503 <i>reduc</i> H504 <i>reduc</i> Constraint:	<b>ne INDEPENDENT I</b> bendent-directive clause ction-clause ction-variable The first non-comment li <i>stmt</i> , <i>forall-stmt</i> , or a <i>for</i> If the first non-comment	Dire is is is or or all-c line	Pective INDEPENDENT [ , new-clause ] [ , reduction-clause ] NEW ( variable-name-list ) REDUCTION ( reduction-variable-list ) array-variable-name scalar-variable-name structure-component ollowing an independent-directive must be a do- onstruct.	26 27 28 29 30 31 32 33 34 35 36 37 38 39 40	
A.5.1     Tl       H501     indep       H502     new-o       H503     reduct       H504     reduct       Constraint:     Constraint:	ne INDEPENDENT I bendent-directive clause ction-clause ction-variable The first non-comment li stmt, forall-stmt, or a for If the first non-comment then that statement mus able.	Dire is is is or or all-c line t con	<pre>ective INDEPENDENT [ , new-clause ]   [ , reduction-clause ] NEW ( variable-name-list ) REDUCTION ( reduction-variable-list ) array-variable-name scalar-variable-name structure-component ollowing an independent-directive must be a do- onstruct. following an independent-directive is a do-stmt, ntain a loop-control option containing a do-vari-</pre>	26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42	
A.5.1     Tl       H501     indep       H502     new-o       H503     reduct       H504     reduct       Constraint:     Constraint:	ne INDEPENDENT I bendent-directive clause stion-clause stion-variable The first non-comment li stmt, forall-stmt, or a for If the first non-comment then that statement mus able. If either the NEW clause on	Dire is is is or or nef <i>all-c</i> line t con	<pre>ective INDEPENDENT [ , new-clause ]   [ , reduction-clause ] NEW ( variable-name-list ) REDUCTION ( reduction-variable-list ) array-variable-name scalar-variable-name structure-component ollowing an independent-directive must be a do- onstruct. following an independent-directive is a do-stmt, ntain a loop-control option containing a do-vari- e REDUCTION clause is present, then the first non- </pre>	26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42 43	
A.5.1 Tl H501 indep H502 new-0 H503 reduc H504 reduc Constraint: Constraint:	ne INDEPENDENT I bendent-directive clause ction-clause ction-variable The first non-comment li stmt, forall-stmt, or a for If the first non-comment then that statement mus able. If either the NEW clause of comment line following the	Dire is is is or or nef <i>all-c</i> line t con	<pre>ective INDEPENDENT [ , new-clause ]   [ , reduction-clause ] NEW ( variable-name-list ) REDUCTION ( reduction-variable-list ) array-variable-name scalar-variable-name structure-component ollowing an independent-directive must be a do- onstruct. following an independent-directive is a do-stmt, ntain a loop-control option containing a do-vari-</pre>	26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42 43 44	

1		• Be a dummy arg	gument;	
2		• Have the SAVE o	r TARGET	f attribute;
3 4		• Occur in a COMM	ON block	;
4 5 6		• Be storage assoc EQUIVALENCE sta		th another object as a result of appearing in an
7		• Be use associate	d;	
8		• Be host associat	,	
9 10			,	coping unit via host association.
11 12 13 14 15	Constraint:	in the same <i>independe</i> a <i>reduction-clause</i> in	e <i>nt-direc</i> the rang	duction-variable may not appear in a new-clause tive, nor may it appear in either a new-clause or e (i.e., the lexical body) of the following do-stmt, to which the independent-directive applies.
16 17 18	Constraint:	A structure-component section-list.	nt in a	reduction-variable may not contain a subscript-
19 20 21	Constraint:	A variable that occurs be of type CHARACTER		<i>luction-var</i> must be of intrinsic type. It may not
22 23 24 25 26 27 28 29 30 31 32 33 33 34	H505 reduc	ction-stmt	is or or or or or or or or or or	<pre>variable = variable mult-op mult-operand variable = add-operand * variable variable = variable add-op add-operand variable = level-2-expr + variable variable = variable and-op and-operand variable = and-operand and-op variable variable = variable or-op or-operand variable = or-operand or-op variable variable = variable equiv-op equiv-operand variable = equiv-operand equiv-op variable variable = reduction-function ( variable , expr ) variable = reduction-function ( expr , variable ) MAX</pre>
35 36 37 38 39	n 500 reauc	enon-junction	or or or or	MAX MIN IAND IOR IEOR
40 41	Constraint:	The two occurances o	f <i>variabl</i>	e in a <i>reduction-stmt</i> must be textually identical.
42 43	A.6 Ext	rinsic Program Un	its	
44 45	A.6.2 D	eclaration of Extri	nsic Pr	ogram Units
46 47 48	H601 funct	ion-stmt	is	<pre>[ prefix ] FUNCTION function-name  ( [ dummy-arg-name-list ] )  [ RESULT ( result-name ) ]</pre>

H602 subre	outine-stmt	is	<pre>[ prefix ] SUBROUTINE subroutine-name [ ( [ dummy-arg-list ] ) ]</pre>	1 2
				2
H603 prefi	x	is	prefix-spec [ prefix-spec ]	4
H604 prefi	x-spec	is	type-spec	5
1 0	•	or	RECURSIVE	6
		or	PURE	7
		$\mathbf{or}$	ELEMENTAL	8
		or	extrinsic- $prefix$	9
Constraint	Within any UDE ontonn	al au	honogram every internal subprogram must be of	10
Constraint:	-		<i>bprogram</i> , every <i>internal-subprogram</i> must be of host and any <i>internal-subprogram</i> whose extrinsic	11
			assumed to be of that extrinsic kind.	12
	kind is not given explicit.	19 16	assumed to be of that extrinsic kind.	13
H605 progr	ram-stmt	is	[ extrinsic-prefix ] PROGRAM program-name	14
H606 mode	ıle-stmt	is	[ extrinsic-prefix ] MODULE module-name	15 16
H607 block	-data-stmt	is	[ extrinsic-prefix ] BLOCK DATA	17
11007 01004	-uutu-stinit	15	[ block-data-name ]	18
				19
Constraint:	Every module-subprogram	n of	any HPF module must be of the same extrinsic	20
			<i>lule-subprogram</i> whose extrinsic kind is not given	21
	explicitly is assumed to h	be of	that extrinsic kind.	22
Constraint:	Every <i>internal-subprogra</i>	m of	f any HPF main-program or module-subprogram	23
			c kind as its host, and any internal-subprogram	24 25
			iven explicitly is assumed to be of that extrinsic	25
	kind.			27
IICOO		•	EVEDINGIA ( setaineir second)	28
H608 extri	nsic-prefix	is	EXTRINSIC ( <i>extrinsic-spec</i> )	29
H609 extri	nsic- $spec$	$\mathbf{is}$	extrinsic-spec-arg-list	30
		or	extrinsic-kind-keyword	31
H610 extri	nsic-spec-arg	is	language	32
	1 5	or	model	33
		or	external-name	34
H611 langu	10.00	is	[LANGUAGE = ]	35
11011 lange	iuge	15	scalar-char-initialization-expr	36
		_	-	37
H612 mode	el	is	[ MODEL = ]	38
			scalar- $char$ - $initialization$ - $expr$	39 40
H613 exter	nal-name	is	[ EXTERNAL_NAME = ]	40
			scalar- $char$ - $initialization$ - $expr$	42
Constantat	In on onthin-is such	int	t logat one of language we del an external	43
Constraint:			at least one of <i>language</i> , <i>model</i> , or <i>external-name</i>	44
	must be specified and no	пеп	nay be specified more than once.	45
Constraint:	If <i>language</i> is specified w	ithou	ut LANGUAGE=, <i>language</i> must be the first item in	46

Constraint: If *language* is specified without LANGUAGE=, *language* must be the first item in the *extrinsic-spec-arg-list*. If *model* is specified without MODEL=, *language* without LANGUAGE= must be the first item and *model* must be the second item in the the second

1 2 3		<i>extrinsic-spec-arg-list.</i> If <i>external-name</i> is specified without EXTERNAL_NAME=, <i>language</i> without LANGUAGE= must be the first item and <i>model</i> without MODEL= must be the second item in the <i>extrinsic-spec-arg-list</i> .
4 5 6	Constraint:	The forms with LANGUAGE=, MODEL=, and EXTERNAL_NAME= may appear in any order except as prohibited above.
7 8 9		Note that these rules for <i>extrinsic-spec-arg-list</i> are as if EXTRINSIC were a procedure with an explicit interface with a <i>dummy-arg-list</i> of LANGUAGE, MODEL, EXTERNAL_NAME, each of which were OPTIONAL.
10 11	Constraint:	In language, values of scalar-char-initialization-expr may be:
12 13		• 'HPF', referring to the HPF language; if a <i>model</i> is not explicitly specified, the <i>model</i> is implied to be 'GLOBAL';
14 15 16		• 'FORTRAN', referring to the ANSI/ISO standard Fortran language; if a <i>model</i> is not explicitly specified, the <i>model</i> is implied to be 'SERIAL';
17 18 19		• 'F77', referring to the former ANSI/ISO standard FORTRAN 77 lan- guage; if a <i>model</i> is not explicitly specified, the <i>model</i> is implied to be 'SERIAL';
20 21		• 'C', referring to the ANSI standard C programming language; if a <i>model</i> is not explicitly specified, the <i>model</i> is implied to be 'SERIAL'; or
22 23 24		• an implementation-dependent value with an implementation-dependent implied <i>model</i> .
25 26		Note that, for most implementations, 'C' will only be allowed for <i>function-stmts</i> and <i>subroutine-stmts</i> occurring in an <i>interface-body</i> .
27 28	Constraint:	If language is not specified it is the same as that of the host scoping unit.
29 30	Constraint:	In model, values of scalar-char-initialization-expr may be:
31 32 33 34 35		<ul> <li>'GLOBAL', referring to the global model,</li> <li>'LOCAL', referring to the local model,</li> <li>'SERIAL', referring to the serial model, or</li> <li>an implementation-dependent value.</li> </ul>
36 37 38	Constraint:	If <i>model</i> is not specified or implied by the specification of a language, it is the same as that of the host scoping unit.
39 40 41	Constraint:	All <i>languages</i> and <i>models</i> whose names begin with the three letters HPF are reserved for present or future definition by this specification and its successors.
42 43 44 45 46 47 48	Constraint:	In <i>external-name</i> , the value of <i>scalar-char-initialization-expr</i> is a character string whose use is determined by the extrinsic kind. For example, an extrinsic kind may use the <i>external-name</i> to specify the name by which the procedure would be known if it were referenced by a C procedure. In such an implementation, a user would expect the compiler to perform any transformations of that name that the C compiler would perform. If <i>external-name</i> is not specified, its value is implementation-dependent.

H614 extrir	nsic-kind-keyword	is or or	HPF HPF_LOCAL HPF_SERIAL	1 2 3
Constraint:	sence of an <i>extrinsic-pre</i> as if it were of extrinsic EXTRINSIC(HPF) or EXTR	<i>fix</i> a kin INSI	t to EXTRINSIC('HPF', 'GLOBAL'). In the ab- an HPF compiler interprets a compilation unit d HPF. Thus, for an HPF compiler, specifying C('HPF', 'GLOBAL') is redundant. Such explicit e required for use with a compiler that supports	4 5 7 8 9 10
Constraint:	<i>main-program</i> whose extroutine of extrinsic kind	insio HPF <u></u>	quivalent to EXTRINSIC('HPF', 'LOCAL'). A c kind is HPF_LOCAL behaves as if it were a sub- LOCAL that is called with no arguments from a nd HPF whose executable part consists solely of	11 12 13 14 15 16
Constraint:	nstraint: EXTRINSIC(HPF_SERIAL) is equivalent to EXTRINSIC('HPF', 'SERIAL'). main-program whose extrinsic kind is HPF_SERIAL behaves as if it were a su routine of extrinsic kind HPF_SERIAL that is called with no arguments from main program of extrinsic kind HPF whose executable part consists solely that call.			
Constraint:	0		hose names begin with the three letters HPF are definition by this specification and its successors.	22 23 24 25
A.8 App	roved Extensions for	Dat	ta Mapping	26 27
A.8.2 Sy	ntax of Attributed Fo	rm	s of Extended Data Mapping Directives	28
H801 comb	ined-attribute-extended	or or or or or or	ALIGN align-attribute-stuff DISTRIBUTE dist-attribute-stuff INHERIT TEMPLATE PROCESSORS DIMENSION ( explicit-shape-spec-list ) DYNAMIC RANGE range-attr-stuff SHADOW shadow-attr-stuff SUBSET	29 30 31 32 33 34 35 36 37 38 39 40 41
Constraint:	The SUBSET attribute ma	y be	applied only to a processors arrangement.	42
A.8.3 Th	ne REDISTRIBUTE	Dire	ective	43 44
H802 redist	tribute-directive	is or	<b>REDISTRIBUTE</b> distributee dist-directive-stuff <b>REDISTRIBUTE</b> dist-attribute-stuff :: distributee-list	45 46 47 48

1 2	Constraint:	A distributee that appears in a REDISTRIBUTE directive must have the DYNAMIC attribute (see Section $8.5$ ).						
3 4 5	Constraint:	A <i>distributee</i> in a REDISTRIBUTE directive may not appear as an <i>alignee</i> in an ALIGN or REALIGN directive.						
6 7 8	Constraint:	Neither the <i>dist-format-clause</i> nor the <i>dist-target</i> in a REDISTRIBUTE directive may begin with "*".						
9 10	A.8.4 T	ne REALIGN Direct	ive					
11 12 13	H803 realig	ŋn-directive	is or	REALIGN alignee align-directive-stuff REALIGN align-attribute-stuff :: alignee-list				
14 15 16	Constraint:	Any <i>alignee</i> that appears in a <b>REALIGN</b> directive must have the <b>DYNAMIC</b> attribute (see Section 8.5).						
17 18	Constraint:	If the <i>align-target</i> specified in the <i>align-with-clause</i> has the DYNAMIC attribute, then each <i>alignee</i> must also have the DYNAMIC attribute.						
19 20 21	Constraint:	An <i>alignee</i> in a <b>REALIGN</b> directive may not appear as a <i>distributee</i> in a <b>DISTRIBUTE</b> or <b>REDISTRIBUTE</b> directive.						
22 23	A.8.5 Th	ne DYNAMIC Direc	tive					
24 25	H804 dyna	mic- $directive$	is	DYNAMIC alignee-or-distributee-list				
26 27	H805 align	ee-or-distributee	is or	alignee distributee				
28 29 30 31	Constraint:	An object in COMMON may not be declared DYNAMIC and may not be aligned to an object (or template) that is DYNAMIC. (To get this kind of effect, modules must be used instead of COMMON blocks.)						
32 33 34	Constraint:	A component of a derived type may have the DYNAMIC attribute only if it also has the POINTER attribute. (See Section 8.9 for further discussion.)						
35 36 37	Constraint:	An object with the SAVE attribute may not be declared DYNAMIC and may not be aligned to an object (or template) that is DYNAMIC.						
38 39	A.8.7 M	apping to Processor	Sub	sets				
40 41 42 43	H806 exten	ded-dist-target	is or or	<pre>processors-name [ ( section-subscript-list ) ] * processors-name [ ( section-subscript-list ) ] *</pre>				
44 45	Constraint:	-		section-subscript-list may not be vector-subscripts r subscripts or subscript-triplets.				
46 47 48	Constraint:	: In the <i>section-subscript-list</i> , the number of <i>section-subscripts</i> must equal the rank of the <i>processor-name</i> .						

Constraint:	Within a DISTRIBUTE directive, each $section$ -subscript must be a $specification$ -expr.					
Constraint:	straint: Within a DISTRIBUTE or a REDISTRIBUTE directive, if both a <i>dist-format-lis</i> and a <i>dist-target</i> appear, the number of elements of the <i>dist-format-list</i> the are not "*" must equal the number of <i>subscript-triplets</i> in the named processe arrangement.					
Constraint:	nstraint: Within a DISTRIBUTE or a REDISTRIBUTE directive, if a <i>dist-target</i> appears I not a <i>dist-format-list</i> , the rank of each <i>distributee</i> must equal the number <i>subscript-triplets</i> in the named processor arrangement.					
Constraint:	onstraint: If either the <i>dist-format-clause</i> or the <i>dist-target</i> in a <b>DISTRIBUTE</b> dire begins with "*" then every <i>distributee</i> must be a dummy argument, <i>exce</i> the distributee has the <b>POINTER</b> attribute.					
Constraint: If the <i>align-spec</i> in an ALIGN directive begins with "*" then every <i>alignee</i> m be a dummy argument, <i>except if the</i> alignee <i>has the</i> <b>POINTER</b> attribute.				15 16 17		
Constraint: An <i>inheritee</i> must be a dummy argument, <i>except if the</i> alignee <i>has the</i> <b>POINT</b> <i>attribute.</i>						
A.8.9 M	apping of Derived Ty	pe (	Components	21 22 23		
H807 distri	butee-extended	is or or or	object-name template-name component-name structure-component	24 25 26 27		
Constraint:	A component of a derived of the component is not a		be may be explicitly distributed only if the type plicitly mapped type.	28 29 30		
Constraint: An object of a derived type may be explicitly distributed only if the deriv type is not an explicitly mapped type.			nay be explicitly distributed only if the derived	31 32 33		
Constraint:	raint: A <i>distributee</i> in a <b>DISTRIBUTE</b> directive may not be a <i>structure-component</i> .			34 35		
Constraint:	int: A <i>distributee</i> in a <b>DISTRIBUTE</b> directive which occurs in a <i>derived-type-def</i> must be the <i>component-name</i> of a component of the derived type.			36 37		
Constraint:	traint: A <i>component-name</i> may occur as a <i>distributee</i> in a <b>DISTRIBUTE</b> directive o curing within the derived type definition only.			38 39 40		
Constraint: A <i>distributee</i> that is a <i>structure-component</i> may occur only in a <b>REDISTRIE</b> directive and every <i>part-ref</i> except the rightmost must be scalar (rank ze The rightmost <i>part-name</i> in the <i>structure-component</i> must have the <b>DYNA</b> attribute.			xcept the rightmost must be scalar (rank zero).	41 42 43 44		
H808 align	ee-extended	is or or	object-name component-name structure-component	45 46 47 48		

1 2	Constraint:	t: A component of a derived type may be explicitly aligned only if the type of the component is not an explicitly mapped type.					
3 4 5	Constraint:	An object of a derived type may be explicitly aligned only if the derived type is not an explicitly mapped type.					
6 7	Constraint:	An <i>alignee</i> in an ALIGN dire	ective may not be a <i>structure-component</i> .				
8 9	Constraint:	An <i>alignee</i> in an <b>ALIGN</b> dire <i>component-name</i> of a comp	ctive that occurs in a <i>derived-type-def</i> must be the onent of the derived type.				
10 11 12	Constraint:	A <i>component-name</i> may occ within the derived type def	ur as an <i>alignee</i> only in an ALIGN directive occuring nition.				
13 14 15 16 17	Constraint:	rective and every part-ref	An <i>alignee</i> that is a <i>structure-component</i> may occur only in a <b>REALIGN</b> di- ective and every <i>part-ref</i> except the rightmost must be scalar (rank zero). The rightmost <i>part-name</i> in the <i>structure-component</i> must have the <b>DYNAMIC</b> .ttribute.				
18 19 20 21 22	H809 align	C	s object-name r template-name r component-name r structure-component				
23 24 25	Constraint:	A <i>component-name</i> may appear as an align target only in an ALIGN directive occuring within the derived type definition that defines that component.					
26 27 28	Constraint:	In an <i>align-target</i> that is a <i>structure-component</i> , every <i>part-ref</i> except the rightmost must be scalar (rank zero).					
29	A.8.10 N	New Distribution Form	ats				
30 31 32 33 34 35	H810 exten	c c	s BLOCK [ ( <i>int-expr</i> ) ] r CYCLIC [ ( <i>int-expr</i> ) ] r GEN_BLOCK ( <i>int-array</i> ) r INDIRECT ( <i>int-array</i> ) r *				
36 37 38	Constraint:	An <i>int-array</i> appearing in a <i>extended-dist-format</i> of a DISTRIBUTE directive or REDISTRIBUTE directive must be an integer array of rank 1.					
39 40 41	Constraint:	An <i>int-array</i> appearing in a <i>extended-dist-format</i> of a <b>DISTRIBUTE</b> directive must be a <i>restricted-expr</i> .					
42 43 44	Constraint:	The size of any <i>int-array</i> appearing with a GEN_BLOCK distribution must be equal to the extent of the corresponding dimension of the target processor arrangement.					
45 46 47 48	Constraint:	The size of any <i>int-array</i> appearing with an <b>INDIRECT</b> distribution must be equal to the extent of the corresponding dimension of the <i>distributee</i> to which the distribution is to be applied.					

A.8.11 The RANGE Directive					
H811 range-directive	is RANGE ranger range-attr-stuff	2 3			
H812 ranger	is object-name	4			
	or template-name	5			
H813 range-attr-stuff	is range-distribution-list	6 7			
H814 range-distribution	is ( range-attr-list )	8			
H815 range-attr	is range-dist-format	9			
-	or ALL	10 11			
H816 range-dist-format		12			
		13			
	OF TNDIDECT	14			
		15 16			
		17			
		18			
Constraint: At least one of the follow	ing must be true:	19			
	-	20 21			
• The <i>ranger</i> has the	DYNAMIC attribute.	22			
• The <i>ranger</i> has the	INHERIT attribute.	23			
• The <i>ranger</i> is specifi	ed with a <i>dist-format-clause</i> of <b>*</b> in a <b>DISTRIBUTE</b> or	24			
combined directive.		25 26			
Constanting. The longth of each parage	attentist must be equal to the papts of the papage	27			
Constraint. The length of each <i>range</i>		28			
Constraint: The ranger must not app	ear as an alignee in an ALIGN or REALIGN directive.	29 30			
		31			
A.8.12 The SHADOW Direct	ive	32			
H817 shadow-directive		33 34			
		35			
H818 shadow-target		36			
H819 shadow-attr-stuff	is ( shadow ence list )	37			
H820 shadow-spec		38 39			
11820 shuuow-spec		40			
H821 width	is internr	41			
	2	42 43			
H822 low-width	<b>15</b> <i>inv-cupt</i>	43			
H823 high-width	is int-expr	45			
Constraint. The interne concepting	a width low width on high width must be a constant	46			
Constraint: The <i>int-expr</i> representing a <i>width</i> , <i>low-width</i> , or <i>high-width</i> must be a constant $_{43}$					

## A.8.11 The RANGE Directive

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Constraint: The *int-expr* representing a *width*, *low-width*, or *high-width* must be a constant specification-expr with value greater than or equal to 0.

## A.9. APPROVED EXTENSIONS FOR DATA AND TASK PARALLELISM

A.9	Approved Extensions fo	r Da	ta and Task Parallelism
A.9.1	Active Processor Sets		
H901	subset- $directive$	is	SUBSET processors-name
A.9.2	2 The ON Directive		
H902	on-directive	is	ON on-stuff
H903	on-stuff	is	home $[$ , resident-clause $]$ $[$ , new-clause
H904	on-construct	is	directive-origin block-on-directive block directive-origin end-on-directive
H905	block- $on$ - $directive$	is	ON on-stuff BEGIN
H906	$end\-on\-directive$	is	END ON
H907	home	is or or	HOME ( variable ) HOME ( template-elmt ) ( processors-elmt )
H908	template-elmt	is	template-name [ ( section-subscript-list )
H909	processors- $elmt$	is	processors-name [ ( section-subscript-list ]
A.9.3	B The RESIDENT Clau	se, D	irective, and Construct
H910	resident-clause	is	RESIDENT resident-stuff
H911	resident-stuff	is	[ ( res-object-list ) ]
H912	resident- $directive$	is	RESIDENT resident-stuff
H913	resident-construct	is	directive-origin block-resident-directive block directive-origin end-resident-directive
H914	block-resident-directive	is	RESIDENT resident-stuff BEGIN
H915	$end\-resident\-directive$	is	END RESIDENT
H916	res-object	is	object
A.9.4	The TASK_REGION	Cons	truct
H917	$task\-region\-construct$	is	directive-origin block-task-region-directive-block
			directive- $origin$ $end$ - $task$ - $region$ - $directive$
H918	block-task-region-directive	is	directive-origin end-task-region-directive

A.10 Approved Extension for	Asynchronous I/O	1	
	or ASYNCHRONOUS	2	
		4	
	<b>or</b> ID = scalar-default-int-variable	5	
	or ASYNCHRONOUS	6	
		7	
Constraint: If either an ASYNCHBONOUS	S or an ID= specifier is present, then both shall be	8	
present.	o or an 12 specifici is present, then soon shan se	9	
Preserve		10	
		11	
Constraint: If an ASYNCHRONOUS specifier is present, the REC= specifier shall appear, a			
format shall not appear, a	nd a <i>namelist-group-name</i> shall not appear.	13	
		14	
Constraint: If an ASYNCHRONOUS specifi	er is present, then no function reference may appear	15	
in an expression anywhere	in the data transfer statement.	16	
		17	
		18	
	or ID = scalar-default-int-variable	19	
	<b>OF PENDING</b> = scalar-aejauli-logical-variable	20	
		21	
Constraint: The ID= and PENDING= sp	or PENDING = scalar-default-logical-variable traint: The ID= and PENDING= specifiers shall not appear in an INQUIRE statement if the FILE= specifier is present. traint: If either an ID= specifier or a PENDING= specifier is present, then both shall be		
-		23	
		24	
Constraint: If either an ID= specifier o	r a PENDING= specifier is present then both shall be	25	
present.	a <b>Habina</b> specifici is present; then both shari be	26	
present.		27	
		28	
A.10.1 The WAIT Statement		29	
A.IO.I The WAII Statement		30	
		31	
H1001 wait-stmt	is WAIT ( $wait$ -spec-list )	32	
	•	33	
1	is UNIT = <i>io-unit</i>	34	
	or ID = scalar-default-int-expr	35	
	or ERR = $label$	36	
	or IOSTAT = label	37	
		38 39	
Constraint: A wait-spec-list shall conta	in exactly one UNIT= specifier, exactly one ID= spec-	40	
ifier, and at most one of e		41	
	-	42	
		43	
A.11 Approved Extensions for	HPF Extrinsics	44	
		45	
A.11.2 Extrinsic Language Bir	ndings	46	
47			
${\rm H1101}\ type-declaration-stmt-extended$	is type-spec [[, attr-spec-extended]:] e	nt#sty-decl-list	

1	H1102 attr-s	spec- $extended$	is	PARAMETER		
2			or	access-spec		
3			$\mathbf{or}$	ALLOCATABLE		
4			or	DIMENSION ( $array$ - $spec$ )		
5			$\mathbf{or}$	EXTERNAL		
6			$\mathbf{or}$	INTENT ( <i>intent-spec</i> )		
7			or	INTRINSIC		
8			or	OPTIONAL		
9			or	POINTER		
10			or	SAVE		
11			or	TARGET		
12			or	MAP_TO ( map-to-spec )		
13				LAYOUT ( <i>layout-spec</i> ) PASS_BY ( <i>pass-by-spec</i> )		
14			or	PASS_BI ( pass-by-spec )		
15 16	H1103 map-	-	is	scalar-char-initialization-expr		
17	H1104 layou	-	is	$scalar\-char\-initialization\-expr$		
18 19	H1105 pass-	by-spec	is	scalar- $char$ - $initialization$ - $expr$		
20						
21	<b>a</b>	<b>T</b> 1 (/	, ,			
22 23	Constraint:	l ne same <i>attr-spec-exten</i> <i>declaration-stmt</i> .	aea	shall not appear more than once in a given $type$ -		
24 25 26	Constraint:	An entity shall not be explicitly given any attribute more than once in a scoping unit.				
27 28 29 30	Constraint:	The attributes MAP_TO, LAYOUT, and PASS_BY may be specified only for dummy arguments within a scoping unit of an extrinsic type for which these attributes have been explicitly defined.				
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44 45						
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48						

# Annex B

# Syntax Cross-reference

This Appendix cross-references symbols used in the formal syntax rules. Rule identifiers beginning with "H" refer to syntax rules of this High Performance Fortran Language Specification; the full rule may be found in Appendix A. Rule identifiers beginning with "R" refer to syntax rules of the Fortran Language Standard ("Fortran 95").

### **B.1** Nonterminal Symbols That Are Defined

Symbol	$\mathbf{Defined}$	Referenced	22
action- $stmt$	R216	H208	23
add-op	R710	H323 H505	24
add-operand	$\mathbf{R706}$	H326 H505	25
align-add-operand	H324	H323 H324	26
a lign-attribute-stuff	H315	H302 H801 H803	27
a lign-directive	H313	H204 H206	28
a lign-directive-stuff	H314	H313 H803	29
align-dummy	H318	H317 H325	30
align-primary	H325	H324	31
align-source	H317	H314 H315	32
align-spec	H320	H319	33
a lign-subscript	H322	H320	34
a lign-subscript-use	H323	H322 H323 H325	35
align-target	H321	H320	36
a lign-target-extended	H809		37
a lign-with-clause	H319	H314 H315	38
alignee	H316	H313 H803 H805	39
alignee-extended	H808		40
alignee- $or$ - $distributee$	H805	H804	41
allocate- $object$	R625		42
allocate-stmt	R622		43
and-op	m R720	H505	44
and-operand	R715	H505	45
array-constructor	R432		46
array-spec	R513	H1102	47
assignment- $stmt$	R735		48

1	association-name	H334	H333
2	attr-spec	R503	11555
3	attr-spec-extended	H1102	H1101
4	block	R801	H904 H913 H917
5	block-data-stmt	H607	
6	block-on-directive	H905	H904
7	block-resident-directive	H914	H913
8	block-task-region-directive	H918	H917
9	call-stmt	R1211	
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12	combined- $attribute$ - $extended$	H801	
13	combined- $decl$	H303	H301
14	combined- $directive$	H301	H204 H206
15	data- $stmt$	R532	
16	deallocate-stmt	R631	
17	directive-origin	H202	H201 H904 H913 H917
18	dist-attribute-stuff	H307	H302 H801 H802
19	dist-directive-stuff	H306	H305 H307 H802
20	dist-format	H310	H309
21	dist-format-clause	H309	H306
22	dist-onto-clause	H311	H306 H307
23	dist-target	H312	H311
24	distribute-directive	H305	H204 H206
25	distributee	H308	H305 H802 H805
26	distributee- $extended$	H807	
27	do-construct	R816	H208
28	dummy-arg	R1223	H602
29	dynamic-directive	H804	H206
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31	end- $on$ - $directive$	H906	H904
32	$end\-resident\-directive$	H915	H913
33	end- $subroutine$ - $stmt$	R1224	
34	$end\-task\-region\-directive$	H919	H917
35	entity- $decl$	R504	H1101
36	equiv- $op$	R722	H505
37	equiv- $operand$	R717	H505
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*       processors-directive       H330       H329         *       processors-clire       H329       H204       H206         *       processors-clire       H909       H907         *       processors-clire       H909       H907         *       processors-clire       H909       H907         *       range-attr       H815       H814         *       range-distribution       H811       H206         *       range-distribution       H814       H813         *       range-distribution       H812       H811         *       range-distribution       H810       H812         *       reduction-clirective       H803       H207         *       reduction-stint       H503       H501         *       reduction-stint       H506       H503         *       resident-clause       H910       H908         *       resident-directive					H602		
4       processors-directive       H329       H204       H206         5       processors-elmt       H909       H907         6       program-stmt       H605         7       range-altr       H815       H814         8       range-directive       H813       H801       H811         9       range-dist-format       H816       H815       H814         10       range-dist-format       H816       H815       H813         11       range-dist-format       H816       H815       H814         12       range-dist-format       H816       H815       H817         13       read-stmt       R909       H74       Ralign-directive       H802       H207         14       redign-directive       H802       H207       H916       H911         15       reduction-function       H506       H505       H504       H503         16       reduction-stante       H910       H912       H914       H916         16       resident-clause       H910       H912       H914       H916         17       reduction-stuff       H911       H910       H912       H914         17       residen	2						
1       processors-elmt       H909       H907         6       program-stmt       H605         7       range-attr       H815       H814         6       range-attr-stuff       H815       H814         7       range-dist-format       H816       H813         10       range-dist-format       H816       H813         11       range-dist-format       H816       H813         12       range-dist-format       H816       H813         12       range-distribution       H814       H813         12       range-distribution       H814       H813         13       redistribute-directive       H803       H207         14       redistribute-directive       H802       H207         15       reduction-clause       H505       H505         16       reduction-stmt       H505       H503         17       resident-clause       H910       H903         28       resident-clause       H910       H903         29       resident-clause       H911       H910         20       resident-clause       H911       H910         21       resident-clause       H913       H208<	3	-			TTOOR		
a       program-stmt       H605         7       range-attr       H815       H814         8       range-attr-stuff       H813       H801       H811         9       range-directive       H811       H206         10       range-dist-format       H816       H815         11       range-dist-format       H816       H815         12       range-dist-format       H816       H813         12       range-dist-format       H810       H813         12       range-dist-format       H814       H813         13       read-stm1       R009       1         14       realign-directive       H803       H207         15       reduction-function       H506       H501         16       reduction-stmt       H506       H503         17       reduction-stmt       H504       H503         18       resident-clause       H910       H911         19       resident-clause       H910       H903         20       resident-stuff       H911       H207         21       resident-stuff       H911       H910       H912       H914         22       section-stuce	4	-			H206		
$\tau$ range-attr       H815       H814 $\epsilon$ range-attr-stuff       H813       H801       H811 $\epsilon$ range-dist-formal       H813       H801       H815 $range-dist-formal$ H816       H815       H813 $range-dist-formal$ H814       H813       H814 $range-dist-formal$ H814       H813       H814 $range-dist-formal$ H809       H207       H814 $read-stml$ R009       H207       H815 $read-stml$ H802       H207       H816 $read-ction-function$ H505       H501       H505 $reduction-function$ H506       H503       H503 $reduction-stmth$ H505       H803       H207 $resident-clause$ H916       H911       H903       H208 $resident-clause$ H910       H903       H208       H206 $resident-stuff$ H912       H207       H206       H206 $resident-stuff$ H911       H910       H912       H914 $section-subscript       R618       H806       H098       H909     $	5	-		H907			
*         range-attr-stuff         H813         H801         H811           *         range-dist-format         H816         H815           10         range-dist-format         H816         H815           11         range-dist-format         H814         H813           12         ranger         H814         H813           12         ranger         H814         H813           12         ranger         H814         H813           13         read-stmt         R909           14         real-stmt         R909           15         redistribute-directive         H803         H207           15         redistribute-directive         H802         H207           15         reduction-function         H505         H503           16         reduction-variable         H504         H503           17         reduction-variable         H504         H901           18         reduction-variable         H504         H903           19         resident-clause         H910         H903           10         resident-directive         H912         H204           10         section-subscript         R618         H	6						
*       range-directive       H811       H206         *       range-dist-format       H816       H815         *       range-dist-format       H816       H813         *       range-distribution       H814       H813         *       ranger       H812       H811         *       range-distribute-directive       H803       H207         *       redistribute-directive       H802       H207         *       reduction-function       H505       H501         *       reduction-function       H505       H503         *       reduction-stmt       H505       H503         *       reduction-turvariable       H504       H503         *       resident-clause       H910       H903         *       resident-clause       H911       H910       H912         *       resident-stuff       H911       H910       H912       H914         *       section-subscript       R618       H806       H817         *       secution-directive       H333       H204       H206         *       shadow-aterset       H818       H817       H819         *       shadow-aterset <t< td=""><td>7</td><td>-</td><td></td><td></td><td></td><td></td><td></td></t<>	7	-					
10       range-dist-format       H816       H815         11       range-distribution       H814       H813         12       ranger       H812       H811         13       read-stmt       R909         14       realign-directive       H803       H207         15       redistribute-directive       H802       H207         16       reduction-clause       H503       H501         17       reduction-function       H505       H802         18       reduction-variable       H504       H503         20       res-object       H916       H911         21       resident-clause       H910       H903         22       resident-clause       H913       H208         23       resident-directive       H913       H208         24       resident-directive       H911       H910       H912         25       section-subscript       R618       H806       H909         26       sequence-directive       H333       H204       H206         27       shadow-directive       H817       H206       H204         28       shadow-target       H819       H811       H817 <td>8</td> <td>0 00</td> <td></td> <td></td> <td>H811</td> <td></td> <td></td>	8	0 00			H811		
11       range-distribution       H814       H813         12       ranger       H812       H811         13       read-stmt       R909         14       realign-directive       H803       H207         15       redistribute-directive       H802       H207         16       reduction-clause       H503       H501         17       reduction-function       H506       H503         18       reduction-stmt       H505       H503         20       res-object       H916       H911         21       resident-clause       H910       H903         22       resident-clause       H910       H903         23       resident-struct       H913       H208         24       resident-stuff       H911       H910       H912         25       section-subscript       R618       H806       H909         26       sequence-directive       H333       H204       H206         27       shadow-directive       H819       H817       H206         28       shadow-directive       H819       H817       H206         29       shadow-directive       H204       H203       H204 <td>9</td> <td>5</td> <td></td> <td></td> <td></td> <td></td> <td></td>	9	5					
12       ranger       H812       H811         13       read-stmt       R909         14       realign-directive       H803       H207         15       redistribute-directive       H802       H207         16       reduction-clause       H503       H501         17       reduction-function       H506       H505         18       reduction-stmt       H506       H503         20       res-object       H910       H903         21       resident-clause       H910       H903         22       resident-clause       H912       H207         24       resident-clause       H911       H910       H912         25       resident-stuff       H911       H910       H912       H914         26       sequence-directive       H333       H204       H206         27       shadow-attr-stuff       H819       H801       H817         28       shadow-directive       H817       H206         29       shadow-target       H818       H817         30       shadow-target       H818       H817         31       specification-directive-extended       H206       H203	10	range-dist-format					
13       real-stmt       R909         14       realign-directive       H803       H207         15       redistribute-directive       H802       H207         16       reduction-clause       H503       H501         17       reduction-function       H506       H505         18       reduction-stmt       H505       H503         19       reduction-variable       H504       H503         20       res-object       H916       H911         21       resident-clause       H910       H903         22       resident-construct       H913       H208         23       resident-stuff       H911       H910       H912         24       resident-stuff       H911       H910       H912       H914         25       section-subscript       R618       H806       H908       H909         26       sequence-directive       H817       H204       H206       H204         29       shadow-directive       H817       H206       H204       H206       H204         29       shadow-target       H818       H817       H203       H2       H204       H206       H204       H206 <t< td=""><td>11</td><td>range-distribution</td><td></td><td></td><td></td><td></td><td></td></t<>	11	range-distribution					
14       realign-directive       H803       H207         15       reduction-clause       H503       H501         16       reduction-clause       H503       H501         17       reduction-function       H506       H503         18       reduction-stmt       H505       H503         19       reduction-variable       H504       H503         20       res-object       H916       H911         21       resident-clause       H910       H903         22       resident-construct       H913       H208         23       resident-stuff       H911       H910       H912         24       resident-stuff       H911       H910       H912       H914         25       section-subscript       R618       H806       H909       H909         26       sequence-directive       H333       H204       H206       H206         27       shadow-directive       H817       H801       H817         28       shadow-directive       H817       H206       H206         29       shadow-directive       H204       H203       H204         20       specification-directive extended       H204 <td>12</td> <td>ranger</td> <td>H812</td> <td>H811</td> <td></td> <td></td> <td></td>	12	ranger	H812	H811			
15       redistribute-directive       H802       H207         16       reduction-clause       H503       H501         17       reduction-function       H506       H503         18       reduction-stmt       H505       1         19       reduction-variable       H504       H503         20       res-object       H916       H911         21       resident-clause       H910       H903         22       resident-construct       H913       H208         23       resident-directive       H912       H207         24       resident-directive       H911       H910       H912 H914         25       section-subscript       R618       H806       H909         26       sequence-directive       H333       H204       H206         27       shadow-attr-stuff       H817       H801       H817         28       shadow-directive       H817       H206       H204         29       shadow-farget       H818       H817         31       specification-directive-extended       H204       H203         32       specification-expr       R734       H204         34       specification-e	13	read- $stmt$	$\mathbf{R909}$				
16reduction-clauseH503H50117reduction-functionH506H50518reduction-stmtH506H50319reduction-variableH504H50320res-objectH916H91121resident-clauseH910H90322resident-clauseH910H90323resident-stuffH911H91024resident-stuffH911H91025section-subscriptR618H80626sequence-directiveH819H80127shadow-attr-stuffH819H80128shadow-directiveH817H20629shadow-attr-stuffH817H20629shadow-directiveH817H20629shadow-targetH818H81730shadow-targetH204H20331specification-directive-extendedH20433specification-partR20434specification-partR61435stat-variableR62038structure-componentR61741subscriptR61742subscript-tripletR61943targetR73744task-region-constructH91145template-declH33146template-declH33147template-elimtH90848template-elimtH90849H907	14	realign- $directive$	H803	H207			
17reduction-functionH506H50518reduction-stmtH50519reduction-variableH504H50320res-objectH916H91121resident-clauseH910H90322resident-constructH912H20823resident-irectiveH912H20724resident-stuffH911H910H91225section-subscriptR618H806H90826sequence-directiveH333H204H20627shadow-attr-stuffH819H801H81728shadow-directiveH817H206H81929shadow-directiveH818H817H81920shadow-targetH818H817H81921specification-directiveH204H203H20422specification-directiveH204H203H20423specification-directiveH204H203H20424specification-directive-extendedH206H807H80825stal-variableR623H809H80926structure-componentR614H504H807H80826subscriptR617H808H80927subscriptR617H808H80928subscript-tripletR619H322H31140subscript-tripletR619H32241targetR737H40842subscript-triplet<	15	redistribute- $directive$	H802	H207			
18reduction-stmtH50519reduction-variableH504H50320res-objectH916H91121resident-clauseH910H90322resident-constructH913H20823resident-directiveH911H910H91224resident-stuffH911H910H91225section-subscriptR618H806H90826sequence-directiveH333H204H20627shadow-attr-stuffH819H801H81728shadow-directiveH817H20629shadow-atgetH818H81729shadow-directiveH204H20329specification-directiveH204H20329specification-directiveH204H20320specification-directiveH204H20320specification-directiveH204H20320specification-partR204H20320subcitation-texprR73424specification-partR62325strideR614H50426subscriptR61727subscriptR61728structure-componentR61429subscript-tripletR61720subscriptR61721subscript-tripletR61922subscript-tripletR191724targetR73725template-declH331 <td< td=""><td>16</td><td>reduction-<math>clause</math></td><td>H503</td><td>H501</td><td></td><td></td><td></td></td<>	16	reduction- $clause$	H503	H501			
19reduction-variableH504H50320res-objectH916H91121resident-clauseH910H90322resident-constructH913H20823resident-directiveH912H20724resident-stuffH911H910H91225section-subscriptR618H806H90826sequence-directiveH333H204H20627shadow-attr-stuffH819H801H81728shadow-directiveH817H20629shadow-targetH818H81730shadow-targetH818H81731specification-directive-extendedH20632specification-directive-extendedH20633specification-partR62334specification-partR62435stat-variableR62336structure-componentR614H50437strideR61740subscriptR61741subscript-tripletR619H32242subscript-tripletR619H32243targetR737444task-region-constructH917H20845template-declH332H33146template-declH332H33146template-declH332H331	17	reduction-function	H506	H505			
20       res-object       H916       H911         21       resident-clause       H910       H903         22       resident-construct       H913       H208         23       resident-directive       H912       H207         24       resident-stuff       H911       H910       H912       H914         25       section-subscript       R618       H806       H909       H909         26       sequence-directive       H333       H204       H206       H206         27       shadow-attr-stuff       H819       H801       H817         28       shadow-directive       H817       H206         29       shadow-spec       H817       H206         29       shadow-target       H818       H817         31       specification-directive       H204       H203         32       specification-airective-extended       H206       H204         33       specification-part       R623       H809         34       specification-part       R614       H504       H807       H808         35       stride       R614       H504       H807       H808       H809       H809       H809       H	18	reduction-stmt	H505				
21resident-clauseH910H90322resident-clauseH913H20823resident-directiveH912H20724resident-stuffH911H910H91225section-subscriptR618H806H90826sequence-directiveH333H204H20627shadow-attr-stuffH819H801H81728shadow-directiveH817H206129shadow-directiveH818H81730shadow-targetH818H81731specification-directive-extendedH206132specification-aperR73434specification-partR20435stat-variableR62336stop-stmtR614H50437strideR61740subscriptR61741subscript-tripletR619H32242subscript-tripletR619H32243targetR737444task-region-constructH917H20845template-declH332H33146template-decliveH331H20447template-elmtH908H907	19	reduction- $variable$	H504	H503			
22       resident-construct       H913       H208         23       resident-directive       H912       H207         24       resident-stuff       H911       H910       H912       H914         25       section-subscript       R618       H806       H908       H909         26       sequence-directive       H333       H204       H206         27       shadow-attr-stuff       H819       H801       H817         28       shadow-directive       H817       H206         29       shadow-directive       H817       H206         29       shadow-target       H818       H817         30       shadow-target       H818       H817         31       specification-directive-extended       H204       H203         32       specification-apart       R204       H203         33       specification-expr       R734       H818       H807         34       specification-part       R204       H204       H203         35       stat-variable       R623       H807       H808       H809         36       stop-stmt       R614       H504       H807       H808       H809 <t< td=""><td>20</td><td>res-object</td><td>H916</td><td>H911</td><td></td><td></td><td></td></t<>	20	res-object	H916	H911			
23resident-directiveH912H20724resident-stuffH911H910H912H91425section-subscriptR618H806H908H90926sequence-directiveH333H204H20627shadow-attr-stuffH819H801H81728shadow-directiveH819H801H81729shadow-directiveH817H20620shadow-targetH818H81730specification-directiveH204H20331specification-directive-extendedH206I32specification-airective-extendedH206I33specification-gartR734I34specification-partR613I35stat-variableR623I36stop-stmtR640I37strideR617I38structure-componentR614H50440subscriptR61741subscript-tripletR619H32242subscript-tripletR619H32243targetR73744task-region-constructH917H20845template-declH332H33146template-directiveH331H20447template-declH331H20447template-directiveH331H20448template-directiveH331H204	21	resident-clause	H910	H903			
24       resident-stuff       H911       H910       H912       H914         25       section-subscript       R618       H806       H908       H909         26       sequence-directive       H333       H204       H206         27       shadow-attr-stuff       H819       H801       H817         28       shadow-directive       H817       H206       H206         29       shadow-spec       H820       H819       H817         30       shadow-target       H818       H817       H206         31       specification-directive       H204       H203       H204         32       specification-directive-extended       H204       H203       H204         33       specification-gart       R734       H204       H203         34       specification-part       R623       H204       H204         35       stat-variable       R620       H204       H807       H808         36       stop-stmt       R614       H504       H807       H808         37       stride       subscript       R617       H206       H206         38       stucture-component       R617       H206       H206	22	resident- $construct$	H913	H208			
25section-subscriptR618H806H908H90926sequence-directiveH333H204H20627shadow-attr-stuffH819H811H81728shadow-directiveH817H20629shadow-specH820H81930shadow-targetH818H81731specification-directiveH204H20332specification-directive-extendedH20633specification-partR73434specification-partR20435stat-variableR62336stop-stmtR614H50437strideR61740subscriptR61741subscript-tripletR619H32242subscript-tripletR619H32244targetR73744task-region-constructH917H20845template-declH331H20447template-elmtH908H907	23	resident- $directive$	H912	H207			
25section-subscriptR618H806H908H90926sequence-directiveH333H204H20627shadow-attr-stuffH819H811H81728shadow-directiveH817H20629shadow-specH820H81930shadow-targetH818H81731specification-directiveH204H20332specification-directive-extendedH20633specification-partR73434specification-partR20435stat-variableR62336stop-stmtR614H50437strideR61740subscriptR61741subscript-tripletR619H32242subscript-tripletR619H32244targetR73744task-region-constructH917H20845template-declH331H20447template-elmtH908H907	24	resident-stuff	H911	H910	H912	H914	
226sequence-directiveH333H204H20627shadow-attr-stuffH819H801H81728shadow-directiveH817H20629shadow-specH820H81930shadow-targetH818H81731specification-directiveH204H20332specification-directive-extendedH20633specification-partR73434specification-partR62335stat-variableR62336stop-stmtR62037strideR614H50438structure-componentR61440subscriptR61741subscript-tripletR61942subscript-tripletR61943targetR73744task-region-constructH91745template-declH33246template-declH33147template-elmtH908490H907	25		R618	H806	H908	H909	
27       shadow-attr-stuff       H819       H801       H817         28       shadow-directive       H817       H206         29       shadow-spec       H820       H819         30       shadow-target       H818       H817         31       specification-directive       H204       H203         32       specification-directive-extended       H206       Image: Comparison of the system of	26	-					
28shadow-directiveH817H20629shadow-specH820H81930shadow-targetH818H81731specification-directiveH204H20332specification-directive-extendedH20633specification-directive-extendedH20634specification-partR73434specification-partR62336stop-stmtR62037strideR62038structure-componentR61440subscriptR61741subscript-tripletR61942subset-directiveH90143targetR73744task-region-constructH917H20845template-declH331H20446template-directiveH331H20447template-elmtH908H907	27	-					
29shadow-specH820H81930shadow-targetH818H81731specification-directiveH204H20332specification-directive-extendedH20633specification-airective-extendedH20634specification-partR73434specification-partR20435stat-variableR62336stop-stmtR62038structure-componentR614H50440subscriptR61741subscript-tripletR619H32242subset-directiveH901H20643targetR73744task-region-constructH917H20845template-declH331H20446template-directiveH331H20447template-elmtH908H907	28						
soshadow-targetH818H81731specification-directiveH204H20332specification-directive-extendedH20633specification-exprR73434specification-partR20435stat-variableR62336stop-stmtR84037strideR62038structure-componentR61440subscriptR61741subscriptR61742subscript-tripletR61943targetR73744task-region-constructH91745template-declH33146template-demtH90847template-elmtH908							
sinspecification-directiveH204H203specification-directive-extendedH206specification-directive-extendedH206specification-exprR734specification-partR204stat-variableR623stat-variableR623strideR620strideR614structure-componentR614subscriptR617subscriptR617subscriptR619subscript-tripletR619subscript-directiveH901subscript-directiveH901subscript-directiveH917subscript-directiveH332stat-directiveH331stat-directiveH331stat-directiveH331stat-directiveH308subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directiveH331subscript-directi		_					
32specification-directive-extendedH20633specification-exprR73434specification-partR20435stat-variableR62336stop-stmtR84037strideR62038structure-componentR614H50439subroutine-stmtH60240subscriptR61741subscript-tripletR619H32242subset-directiveH901H20643targetR73744task-region-constructH917H20845template-declH331H20447template-elmtH908H907		-					
33specification-exprR73434specification-partR20435stat-variableR62336stop-stmtR84037strideR62038structure-componentR614H50439subroutine-stmtH60240subscriptR61741subscript-tripletR619H32242subset-directiveH901H20643targetR73744task-region-constructH917H20845template-declH331H20447template-elmtH908H907				11200			
34specification-partR20435stat-variableR62336stop-stmtR84037strideR62038structure-componentR614H50439subroutine-stmtH60240subscriptR61741subscript-tripletR619H32242subset-directiveH901H20643targetR73744task-region-constructH917H20845template-declH331H20447template-elmtH908H907							
35stat-variableR62336stop-stmtR84037strideR62038structure-componentR614H50439subroutine-stmtH60240subscriptR61741subscript-tripletR61942subset-directiveH90143targetR73744task-region-constructH917H20845template-declH332H33146template-directiveH908H907		1 0 1					
36         stop-stmt         R840           37         stride         R620           38         structure-component         R614         H504         H807         H808         H809           39         subroutine-stmt         H602							
37       stride       R620         38       structure-component       R614       H504       H807       H808       H809         39       subroutine-stmt       H602							
38structure-componentR614H504H807H808H80939subroutine-stmtH60240subscriptR61741subscript-tripletR619H32242subset-directiveH901H20643targetR73744task-region-constructH917H20845template-declH332H33146template-directiveH908H907		-					
39subroutine-stmt $H602$ 40subscript $R617$ 41subscript-triplet $R619$ $H322$ 42subset-directive $H901$ $H206$ 43target $R737$ 44task-region-construct $H917$ $H208$ 45template-decl $H332$ $H331$ 46template-directive $H331$ $H204$ 47template-elmt $H908$ $H907$				H504	H807	H808	H809
40subscriptR61741subscript-tripletR619H32242subset-directiveH901H20643targetR73744task-region-constructH917H20845template-declH332H33146template-directiveH331H20447template-elmtH908H907		-		11004	11001	11000	11005
41subscript-tripletR619H32242subset-directiveH901H20643targetR73744task-region-constructH917H20845template-declH332H33146template-directiveH331H20447template-elmtH908H907							
42subset-directiveH901H20643targetR73744task-region-constructH917H20845template-declH332H33146template-directiveH331H20447template-elmtH908H907		-		Цзоо			
43     target     R737       44     task-region-construct     H917     H208       45     template-decl     H332     H331       46     template-directive     H331     H204       47     template-elmt     H908     H907							
44task-region-constructH917H20845template-declH332H33146template-directiveH331H204H20647template-elmtH908H907				11200			
45template-declH332H33146template-directiveH331H204H20647template-elmtH908H907		-		<u>По</u> 00			
46template-directiveH331H204H20647template-elmtH908H907		_					
47 template-elmt H908 H907		-			Hooe		
•		-			П <i>2</i> 00		
48 <i>iype-aeciaration-simi</i> K501		-		H907			
	48	type-aectaration-stmt	K901				

type - $declaration$ - $stmt$ - $extended$	H1101		1
type-spec	R502	H604 H1101	2
variable	R601	H505 H907	3
wait-spec	H1002	H1001	4
wait-stmt	H1001		5
where-construct	R739	H208	6
where-stmt	R738	11200	7
width	H821	H820	8
write-stmt	R910	11020	
write-simi	N910		9
			10
	1 . A T.T.		11
B.2 Nonterminal Symbols T	hat Are Not	Defined	12
Symbol		Referenced	13 14
access-spec		H1102	15
array-variable-name		H1102 H504	16
block-data-name		H607	
char-initialization-expr		H611 H612 H613 H1103 H1104 H1104	17
common-block-name		H334	
component-name		H807 H808 H809 H818	19
*		H1002	20
default-int-expr		H601	21
dummy-arg-name		H601	22
function-name			23
int-array		H810	24
intent-spec		H1102	25
module-name		H606	26
object		H916	27
object- $name$		H303 H308 H316 H321 H334 H402	28
		H807 H808 H809 H812 H818	29
processors-name		H304 H312 H330 H806 H901 H909	30
program-name		H605	31
result-name		H601	32
subroutine-name		H602	33
template-name		H304 H308 H321 H332 H807 H809	34
		H812 H908	35
variable- $name$		H502 H504	36
			37
			38
B.3 Terminal Symbols			39
			40
Symbol		Referenced	41
!HPF\$		H202	42
(		H302 H303 H309 H310 H314 H315	43
		H320 H325 H330 H332 H502 H503	44
		H505 H601 H602 H608 H801 H806	45
		H810 H814 H816 H819 H907 H908	46
		H909 H911 H1001H1102	47
)		H302 H303 H309 H310 H314 H315	48

1	
2	
3	
4	
5	*
6	
7	*HPF\$
8	+
9	3
10	1
11	:
12	::
13	=
14	ALIGN
15	ALL
16	ALLOCATABLE
17	BEGIN
18	BLOCK
19	CHPF\$
20	CYCLIC
21	DATA
22	DIMENSION
23	DISTRIBUTE
24	DYNAMIC
25	ELEMENTAL
26	END
27	ERR
28	EXTERNAL
29	EXTERNAL_NAME
30	EXTRINSIC
31	FUNCTION
32	GEN_BLOCK
33	HOME
34	HPF
35	HPF_LOCAL
36	HPF_SERIAL IAND
37	ID
38 39	IEOR
	INDEPENDENT
40 41	INDEPENDENT
41	INHERIT
	INTENT
43 44	INTRINSIC
44 45	IOR
45	IOSTAT
46	LANGUAGE
47	LAYOUT
40	

H320 H325 H330 H332 H502 H503 H505 H601 H602 H608 H801 H806 H810 H814 H816 H819 H907 H908 H909 H911 H1001 H1102 H309 H310 H312 H317 H320 H322 H324 H505 H806 H810 H816 H202 H505H501 H505 H903 H1101 H334 H317 H820 H301 H333 H802 H803 H1101 H505 H611 H612 H613 H1002 H302 H313 H801 H815H1102 H905 H914 H310 H607 H810 H816 H202 H310 H810 H816 H607 H302 H801 H1102 H302 H305 H801 H801 H804 H604 H906 H915 H919 H1002 H1102 H613H608H601H810 H816 H907 H614 H614 H614 H506H1002 H506H501H810 H816 H302 H401 H801 H1102 H1102 H506H1002 H611

H1102

MAP_TO
MAX
MIN
MODEL
MODULE
NEW
NO
ON
ONTO
OPTIONAL
PARAMETER
PASS_BY
POINTER
PROCESSORS
PROGRAM
PURE
RANGE
REALIGN
RECURSIVE
REDISTRIBUTE
REDUCTION
RESIDENT
RESULT
SAVE
SEQUENCE
SHADOW
SUBROUTINE
SUBSET
TARGET
TASK_REGION
TEMPLATE
UNIT
WAIT
WITH

H1102			1
H506			2
H506			3
H612			4
H606			5
H502			6
H333			7
H902 H905	H906		8
H311			9
H1102			10
H1102			11
H1102			12
H1102			13
H302 H329	H801		14
H605			15
H604			16
H801 H811			17
H803			18
H604			19
H802			20
H503			21
H910 H912	H914	H915	22
H601			23
H1102			24
H333			25
${ m H801}$ ${ m H817}$			26
H602			27
H801 H901			28
H1102			29
H918 H919			30
H302 H331	H801		31
H1002			32
H1001			33
H319			34
			35
			36
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			39
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## Annex C

## HPF 1.1 Subset

As part of the definition of the previous version of the High Performance Fortran language, HPF 1.1, a subset language was formally defined, based on the Fortran 77 language. The goal was to permit more rapid implementations of a useful subset of HPF that did not require full implementation of the new ANSI/ISO standard Fortran ("Fortran 90").

No subset language is defined as part of the current version, HPF 2.0. This Annex is included in the HPF 2.0 language document as a convenient summary of the HPF 1.1 Subset, which has served as a minimum requirement for HPF implementations.

 $^{25}$ 

#### C.1 Fortran 90 Features in the HPF 1.1 Subset

The features of the HPF 1.1 subset languages are listed below. For reference, the section numbers from the Fortran 90 standard are given along with the related syntax rule numbers:

- All FORTRAN 77 standard conforming features, except for storage and sequence association.
- The Fortran 90 definitions of MIL-STD-1753 features:
  - DO WHILE statement (8.1.4.1.1 / R821)
  - END DO statement (8.1.4.1.1 / R825)
  - IMPLICIT NONE statement (5.3 / R540)
  - INCLUDE line (3.4)
    - scalar bit manipulation intrinsic procedures: IOR, IAND, NOT, IEOR, ISHFT, ISHFTC, BTEST, IBSET, IBCLR, IBITS, MVBITS (13.13)
    - binary, octal and hexadecimal constants for use in DATA statements (4.3.1.1 / R407 and 5.2.9 / R533)
- Arithmetic and logical array features:
  - array sections (6.2.2.3 / R618-621)
    - \* subscript triplet notation (6.2.2.3.1)
    - \* vector-valued subscripts (6.2.2.3.2)
- array constructors limited to one level of implied DO (4.5 / R431)

- arithmetic and logical operations on whole arrays and array sections $(2.4.3, 2.4.5, and 7.1)$	1 2
- array assignment (2.4.5, 7.5, 7.5.1.4, and 7.5.1.5)	3
- masked array assignment (7.5.3)	4
* WHERE statement $(7.5.3 / R738)$	5 6
* block WHERE ELSEWHERE construct (7.5.3 / R739)	7
- array-valued external functions (12.5.2.2)	8
- automatic arrays (5.1.2.4.1)	9
<ul> <li>ALLOCATABLE arrays and the ALLOCATE and DEALLOCATE statements (5.1.2.4.3, 6.3.1 / R622, and 6.3.3 / R631)</li> </ul>	10 11 12
- assumed-shape arrays (5.1.2.4.2 / R516)	13
	14
• Intrinsic procedures:	15
The list of intrinsic functions and subroutines below is a combination of (a) routines	16 17
which are entirely new to Fortran and (b) routines that have always been part of Fortran but have been extended here to new encument and result types. The new	18
Fortran, but have been extended here to new argument and result types. The new or extended definitions of these routines are part of the subset. If a FORTRAN 77	19
routine is not included in this list, then only the original FORTRAN 77 definition is	20
part of the subset.	21 22
For all of the intrinsics that have an optional argument $\texttt{DIM}$ , only actual argument	23
expressions for DIM that are initialization expressions are part of the subset. The	24
intrinsics with this constraint are marked with †in the list below.	25
- the argument presence inquiry function: <b>PRESENT</b> (13.10.1)	26 27
<ul> <li>all the numeric elemental functions: ABS, AIMAG, AINT, ANINT, CEILING, CMPLX, CONJG, DBLE, DIM, DPROD, FLOOR, INT, MAX, MIN, MOD, MODULO, NINT, REAL, SIGN (13.10.2)</li> </ul>	28 29 30
<ul> <li>all mathematical elemental functions: ACOS, ASIN, ATAN, ATAN2, COS, COSH, EXP, LOG, LOG10, SIN, SINH, SQRT, TAN, TANH (13.10.3)</li> </ul>	31 32
<ul> <li>all the bit manipulation elemental functions : BTEST, IAND, IBCLR, IBITS, IBSET, IEOR, IOR, ISHFT, ISHFTC, NOT (13.10.10)</li> </ul>	33 34
$-$ all the vector and matrix multiply functions: DOT_PRODUCT, MATMUL (13.10.13)	35 36
- all the array reduction functions: ALL <sup>†</sup> , ANY <sup>†</sup> , COUNT <sup>†</sup> , MAXVAL <sup>†</sup> , MINVAL <sup>†</sup> , PRODUCT <sup>†</sup> , SUM <sup>†</sup> (13.10.14)	37 38
- all the array inquiry functions: ALLOCATED, LBOUND <sup><math>\dagger</math></sup> , SHAPE, SIZE <sup><math>\dagger</math></sup> , UBOUND <sup><math>\dagger</math></sup> (13.10.15)	39 40
- all the array construction functions: MERGE, PACK, SPREAD†, UNPACK (13.10.16)	41 42
- the array reshape function: <b>RESHAPE</b> $(13.10.17)$	43
- all the array manipulation functions: CSHIFT <sup>†</sup> , EOSHIFT <sup>†</sup> , TRANSPOSE (13.10.18)	44
- all array location functions: MAXLOC <sup>†</sup> , MINLOC <sup>†</sup> (13.10.19)	45
- all intrinsic subroutines: DATE_AND_TIME, MVBITS, RANDOM_NUMBER, RANDOM_SEED,	46
SYSTEM_CLOCK (3.11)	47 48
	10

1	• Declarations:
2 3 4 5	- Type declaration statements, with all forms of <i>type-spec</i> except <i>kind-selector</i> and TYPE(type-name), and all forms of <i>attr-spec</i> except <i>access-spec</i> , TARGET, and POINTER. (5.1 / R501-503, R510)
6 7	<ul> <li>attribute specification statements: ALLOCATABLE, INTENT, OPTIONAL, PARAMETER, SAVE (5.2)</li> </ul>
8 9	• Procedure features:
10 11	- INTERFACE blocks with no generic-spec or module-procedure-stmt $(12.3.2.1)$
12	- optional arguments $(5.2.2)$
13 14	- keyword argument passing (12.4.1 /R1212)
15 16	• Syntax improvements:
17	- long (31 character) names (3.2.2)
18 19	- lower case letters $(3.1.7)$
20	$-$ use of "_" in names (3.1.3)
21 22	– "!" initiated comments, both full line and trailing $(3.3.2.1)$
23 24	C.2 HPF 1.1 Directives and Language Extensions in the HPF 1.1 Subset
25	The following HPF 1.1 directives and language extensions to Fortran 90 were included in
26	the HPF 1.1 Subset:
26 27 28 29	
27 28	<ul><li>the HPF 1.1 Subset:</li><li>The basic data distribution and alignment directives: ALIGN, DISTRIBUTE,</li></ul>
27 28 29 30	<ul> <li>the HPF 1.1 Subset:</li> <li>The basic data distribution and alignment directives: ALIGN, DISTRIBUTE, PROCESSORS. and TEMPLATE.</li> </ul>
27 28 29 30 31 32 33 34	<ul> <li>the HPF 1.1 Subset:</li> <li>The basic data distribution and alignment directives: ALIGN, DISTRIBUTE, PROCESSORS. and TEMPLATE.</li> <li>The forall-statement (but not the forall-construct).</li> </ul>
27 28 29 30 31 32 33	<ul> <li>the HPF 1.1 Subset:</li> <li>The basic data distribution and alignment directives: ALIGN, DISTRIBUTE, PROCESSORS. and TEMPLATE.</li> <li>The forall-statement (but not the forall-construct).</li> <li>The INDEPENDENT directive.</li> </ul>
27 28 29 30 31 32 33 34 35 36	<ul> <li>the HPF 1.1 Subset:</li> <li>The basic data distribution and alignment directives: ALIGN, DISTRIBUTE, PROCESSORS. and TEMPLATE.</li> <li>The forall-statement (but not the forall-construct).</li> <li>The INDEPENDENT directive.</li> <li>The SEQUENCE and NO SEQUENCE directives.</li> <li>The system inquiry intrinsic functions NUMBER_OF_PROCESSORS and</li> </ul>
27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42 43 44	<ul> <li>the HPF 1.1 Subset:</li> <li>The basic data distribution and alignment directives: ALIGN, DISTRIBUTE, PROCESSORS. and TEMPLATE.</li> <li>The forall-statement (but not the forall-construct).</li> <li>The INDEPENDENT directive.</li> <li>The SEQUENCE and NO SEQUENCE directives.</li> <li>The system inquiry intrinsic functions NUMBER_OF_PROCESSORS and PROCESSORS_SHAPE.</li> <li>The computational intrinsic functions ILEN, and the HPF extended Fortran intrinsics MAXLOC and MINLOC, with the restriction that any actual argument expression</li> </ul>
27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42 43	<ul> <li>the HPF 1.1 Subset:</li> <li>The basic data distribution and alignment directives: ALIGN, DISTRIBUTE, PROCESSORS. and TEMPLATE.</li> <li>The forall-statement (but not the forall-construct).</li> <li>The INDEPENDENT directive.</li> <li>The SEQUENCE and NO SEQUENCE directives.</li> <li>The system inquiry intrinsic functions NUMBER_OF_PROCESSORS and PROCESSORS_SHAPE.</li> <li>The computational intrinsic functions ILEN, and the HPF extended Fortran intrinsics MAXLOC and MINLOC, with the restriction that any actual argument expression corresponding to an optional DIM argument must be an initialization expression.</li> <li>For a discussion of the rationale by which features were chosen for the HPF 1.1 Subset,</li> </ul>
27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42 43 44 45	<ul> <li>the HPF 1.1 Subset:</li> <li>The basic data distribution and alignment directives: ALIGN, DISTRIBUTE, PROCESSORS. and TEMPLATE.</li> <li>The forall-statement (but not the forall-construct).</li> <li>The INDEPENDENT directive.</li> <li>The SEQUENCE and NO SEQUENCE directives.</li> <li>The system inquiry intrinsic functions NUMBER_OF_PROCESSORS and PROCESSORS_SHAPE.</li> <li>The computational intrinsic functions ILEN, and the HPF extended Fortran intrinsics MAXLOC and MINLOC, with the restriction that any actual argument expression corresponding to an optional DIM argument must be an initialization expression.</li> <li>For a discussion of the rationale by which features were chosen for the HPF 1.1 Subset,</li> </ul>

## Annex D

# **Previous HPFF Acknowledgments**

The current HPF 2.0 document would not have been possible without the contributions of the previous series of HPFF meetings. Following are the acknowledgements for those efforts.

#### D.1 HPFF Acknowledgments

Technical development for HPF 1.0 was carried out by subgroups, and was reviewed by the full committee. Many people served in positions of responsibility:

- Ken Kennedy, Convener and Meeting Chair;
- Charles Koelbel, Executive Director and Head of the FORALL Subgroup;
- Mary Zosel, Head of the Fortran 90 and Storage Association Subgroup;
- Guy Steele, Head of the Data Distribution Subgroup;
- Rob Schreiber, Head of the Intrinsics Subgroup;
- Bob Knighten, Head of the Parallel I/O Subgroup;
- Marc Snir, Head of the Extrinsics Subgroup;
- Joel Williamson and Marina Chen, Heads of the Subroutine Interface Subgroup; and
- David Loveman, Editor.

Geoffrey Fox convened the first HPFF meeting with Ken Kennedy and later led a group to develop benchmarks for HPF. Clemens-August Thole organized a group in Europe and was instrumental in making this an international effort. Charles Koelbel produced detailed meeting minutes that were invaluable to subgroup heads in preparing successive revisions to the draft proposal. Guy Steele developed LATEX macros for a variety of tasks, including formatting BNF grammar, Fortran code and pseudocode, and commentary material; the document would have been much less aesthetically pleasing without his efforts.

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University of Vienna	37
Yale University	38

Many people contributed sections to the final language specification and HPF Journal of Development, including Alok Choudhary, Geoffrey Fox, Tom Haupt, Maureen Hoffert, Ken Kennedy, Robert Knighten, Charles Koelbel, David Loveman, Piyush Mehrotra, John Merlin, Tin-Fook Ngai, Rex Page, Sanjay Ranka, Robert Schreiber, Richard Shapiro, Marc Snir, Matt Snyder, Guy Steele, Richard Swift, Min-You Wu, and Mary Zosel. Many others contributed shorter passages and examples and corrected errors.

39

Because public input was encouraged on electronic mailing lists, it is impossible to 46 identify all who contributed to discussions; the entire mailing list was over 500 names long. 47 Following are some of the active participants in the HPFF process not mentioned above: 48

1		N. Arunasalam	Werner Assmann	Marc Baber
2		Babak Bagheri	Vasanth Bala	Jason Behm
3		Peter Belmont	Mike Bernhardt	Keith Bierman
4		Christian Bishof	John Bolstad	William Camp
5		Duane Carbon	Richard Carpenter	Brice Cassenti
6		Doreen Cheng	Mark Christon	Fabien Coelho
7		Robert Corbett	Bill Crutchfield	J. C. Diaz
8		James Demmel	Alan Egolf	Bo Einarsson
9		Pablo Elustondo	Robert Ferrell	Rhys Francis
10		Hans-Hermann Frese	Steve Goldhaber	Brent Gorda
11		Rick Gorton	Robert Halstead	Reinhard von Hanxleden
12		Hiroki Honda	Carol Hoover	Steven Huss-Lederman
13		Ken Jacobsen	Elaine Jacobson	Behm Jason
14		Alan Karp	Ronan Keryell	Anthony Kimball
15		Ross Knippe	Bruce Knobe	David Kotz
16		Ed Krall	Tom Lake	Peter Lawrence
17		Bryan Lawver	Bruce Leasure	Stewart Levin
18		David Levine	Theodore Lewis	Woody Lichtenstein
19		Ruth Lovely	Doug MacDonald	Raymond Man
20		Stephen Mark	Philippe Marquet	Jeanne Martin
21		Oliver McBryan	Charlie McDowell	Michael Metcalf
22		Charles Mosher	Len Moss	Lenore Mullin
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29		Paul St.Pierre	Nick Stanford	Mia Stephens
30		Jaspal Subhlok	Xiaobai Sun	Hanna Szoke
31		Bernard Tourancheau	Anna Tsao	Alex Vasilevsky
32		Stephen Vavasis	Arthur Veen	Brian Wake
33		Ji Wang	Karen Warren	D.C.B. Watson
34		Matthijs van Waveren	Robert Weaver	Fred Webb
35		Stephen Whitley	Michael Wolfe	Fujio Yamamoto
36		Marco Zagha		
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The following organizations made the language draft available by anonymous FTP access and/or mail servers: AT&T Bell Laboratories, Cornell Theory Center, GMD-I1.T (Sankt Augustin), Oak Ridge National Laboratory, Rice University, Syracuse University, and Thinking Machines Corporation. These outlets were instrumental in distributing the document.

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• Alok Choudhary, Head of Parallel I/O Subgroup;
• Chuck Koelbel, Head of Irregular Distributions Subgroup;
• Rob Schreiber, Head of Implementation Subgroup;
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1	Paula Vaughan, Donna Reese	
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3	P. Sadayappan, Chua-Huang H	uangOhio State University
4	Andrew Johnson	OSF Research Institute
5	Chip Rodden, Jeff Vanderlip .	Pacific Sierra Research
6	Larry Meadows, Doug Miles .	The Portland Group, Inc.
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9	Ira Baxter	Schlumberger
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12	-	Thinking Machines Corp., Silicon Graphics Inc.
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14		University of Denver
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31	8	WWW home-page for HPFF, and to the University of
32	Maryland for establishing a benchm	· · · · ·
33		Rice University were responsible for meeting planning
34		I continued to handle financial details of the project.
35		t for research and administrative activities from grants
36	from ARPA, DOE, and NSF.	0
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## Annex E

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# Policy and Mechanism for Recognized Extrinsic Interfaces

HPF defines certain extrinsics such as HPF\_LOCAL, and HPF\_SERIAL as interfaces that HPFF believes are useful to the HPF community. But there are many more such extrinsic interfaces beyond those maintained by HPFF. HPFF has a adopted a policy of formally recognizing certain extrinsic interface definitions, where the interface, and its addition to the HPF document is considered to be a service to the HPF community. Examples are language bindings to HPF or library packages.

#### E.1 Extrinsic Policy

To be considered for HPFF recognition, a proposed extrinsic must demonstrate the following things. It should be noted, however, that meeting these criteria does not guarentee acceptance of a proposed interface by HPFF.

- conformance to HPF rules for calling extrinsics,
- significant new functionality,
- existing practice such as users, implementations, etc.,
- institutional backing with evidence of ongoing support,
- coherent documentation,
- non-proprietary interface definition, and
- copyright goes to HPFF for interface, with permission to use (royalty free).

If a proposed extrinsic is accepted by HPFF, then:

- HPFF will recogize the interface and reference it in documentation, but HPFF does not assume responsibility for the extrinsic or its interface.
- The sponsor of the extrinsic must continue to conform to the HPF interface rules for extrinsics. The interface HPFF approves must not change without HPFF approval.
- The sponsor must assume responsibility for any CCI requests concernting the extrinsic.

A list of recognized extrinsic interfaces will be included in HPF documentation, with the following guidelines:	1 2
• There should be a single page introduction to the extrinsic which contains:	3 4
	5
- the name of the extrinsic,	6
- a brief abstract of functionality,	7
- a brief and informal description of the interface,	8 9
- information about platform and system availability, and	10
<ul> <li>reference and contacts for formal documentation, continued responsibility, and additional information (e.g. compiler availability).</li> </ul>	11 12
• There should be about two pages with short examples of usage.	13 14
• A short paper with the formal definition of the interface and an informal description of the functionality of the extrinsic.	15 16
E.2 Extrinsic Interface Mechanism	17 18 19
The HPF www-home page will have instructions for submission of an extrinsic interface. For HPFF consideration, the sponsor prepares a proposal that includes:	20 21 22
ullet a statement of what significant new functionality is provided,	23 24
• a description of existing practice,	25
$\bullet$ a statement of institutional backing with evidence of ongoing support,	26 27
• a copy of the complete documentation or a reference to an online version of the documentation,	28 29
• a draft of the text (described above) that would be included in the HPFF documen- tation, and	30 31 32
• a statement justifying the claim that the interface follows HPF conventions for calling extrinsics.	33 34 35
If the proposed extrinsic interface is approved by HPFF, the sponsor then submits:	36 37
• a formal statement for HPFF records that the interface definition is non-proprietary and that the copyright of the interface belongs HPFF,	38 39
ullet the formal contact for CCI and continued maintenance of the interface, and	40 41
ullet a copy of the interface documentation formatted for HPFF use, including a copy in	42
the current document and web mark-up languages.	43 44
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## Annex F

## HPF\_CRAFT

HPF\_CRAFT is a hybrid language, combining an SPMD execution model with high performing HPF features. The model combines the multi-threaded execution of HPF\_LOCAL and the HPF syntax. The goal of HPF\_CRAFT is to attain the potential performance of an SPMD programming model with access to HPF features and a well-defined extrinsic interface to HPF.

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### F.1 Introduction

HPF\_CRAFT is a hybrid language, combining an SPMD execution model with high performing and portable HPF features. The model combines the multi-threaded execution of HPF\_LOCAL and the HPF syntax and features. The goal of HPF\_CRAFT is to attain the potential performance of an SPMD programming model with access to HPF features and a well-defined extrinsic interface to HPF. It is built on top of the HPF\_LOCAL extrinsic environment.

SPMD features and a multi-threaded model allow the user to take advantage of the performance and opportunity for low level access of a more general purpose programming model. Including HPF data distribution features gives the programmer access to high performing aspects of both models, but with the added responsibility of working with a more low-level execution model. HPF\_CRAFT is best suited for platforms that support one way communication features, but is consistent with HPF and easily targeted for platforms that have HPF and can support SPMD programming styles.

The HPF features included in HPF\_CRAFT are a subset of the full HPF language chosen for their performance and their broad portability and ease of use. HPF\_CRAFT contains additional features to support SPMD programming styles. There are some differences from HPF, however. For example, I/O causes differences; in HPF\_CRAFT different processors are allowed to read from different files at the same time, in HPF the processors must all read from the same file. The differences in the models are principally caused by the multi-threaded execution model and the introduction of HPF\_LOCAL data rules.

HPF\_CRAFT allows for the notion of *private data*. Data defaults to a mapping in
which data items are allocated so that each processor has a unique copy. The values of
the individual data items and the flow of control may vary from processor to processor
within HPF\_CRAFT. This behavior is consistent with the behavior of HPF\_LOCAL. In
HPF\_CRAFT a processor may be individually named and code executed based upon which
processor it is executing on. HPF\_CRAFT also allows for the notion of *private loops*. A

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private loop is executed in entirety by each processor. The rules governing the interface to HPF\_CRAFT subprograms are similar to those for the HPF\_LOCAL interface. Dummy arguments use a hybrid of the interfaces between HPF and itself and that of HPF and HPF\_LOCAL. Explicitly mapped dummy arguments behave just as they do in HPF, while default (private) dummy arguments use the HPF\_LOCAL calling convention. HPF\_CRAFT will be initially made available on Cray MPP systems and may also be available on Cray vector architectures. Future versions of HPF\_CRAFT are possible on other vendor's architectures as well. HPF\_CRAFT is being implemented for Cray Research by The Portland Group, Inc. For Cray systems, HPF\_CRAFT may be obtained through the Cray Research Inc. Orderdesk, Cray Research Inc. orderdsk@cray.com

Additional formal documentation, requests, and suggestions can be made to

The Portland Group 9150 SW Pioneer Ct., Suite H Wilsonville, OR 97070 (503) 682-2806 trs@pgroup.com

#### F.2 Examples of Use

(612) 683-5907

HPF\_CRAFT is intended for use in circumstances where greater control and performance are desired for MIMD style architectures. Since data may be declared to be private, local control is made more available and since processor information is available message passing and direct memory access programming styles can be seamlessly integrated with explicitly mapped data.

The following examples show some of the capabilities of HPF\_CRAFT that are different from those of HPF. Others, such as integrated message passing and synchronization primitives are not shown. Much of HPF can also be used within HPF\_CRAFT.

Example 1 illustrates the difference between the default distribution for data and the distribution of mapped data.

```
! Example 1
```

```
INTEGER PRIVATE_A(100, 20), PRIVATE_B(12, 256), PRIVATE_C
INTEGER MAPPED_A(100, 20), MAPPED_B(12, 256), MAPPED_C
!HPF$ DISTRIBUTE MAPPED_A(BLOCK, BLOCK), MAPPED_B(BLOCK, *), MAPPED_C
```

In the above example, given 8 processors, there would be 8 \* 100 \* 20 (or 16,000) elements 46 in the array PRIVATE\_A. Each processor contains an entire array named PRIVATE\_A. The 47 elements of PRIVATE\_A on processor 1 cannot be referenced using implicit syntax by any 48 other processor. There are only 100 \* 20 (or 2000) elements of array MAPPED\_A, however,
 and these elements are distributed about the machine in a (BLOCK, BLOCK) fashion.

The difference between the PRIVATE\_A declaration in HPF\_CRAFT and that in HPF is the most instructive. In HPF\_CRAFT each processor contains one copy of the array, and the values of the elements of the array may vary from processor to processor. HPF implementations are permitted to make one copy of the array per processor the default, but the values of these copies must remain coherent across all processors. In HPF there is no way to write a conforming program in which different processors have different values for the same array.

Example 2 shows the usefulness of the ON clause for the INDEPENDENT loop as well as giving an example of how private data may be used.

```
12
     !
        Example 2
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14
            PRIVATE_C = 0
15
     !HPF$ INDEPENDENT (I, J) ON MAPPED_B(I, J)
16
            DO J=1,256
17
               DO I=1,12
18
                  MAPPED_B(I, J) = MAPPED_B(I, J) + 5
19
                  PRIVATE_C = PRIVATE_C + MAPPED_B(I, J)
20
               ENDDO
21
            ENDDO
22
```

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In this example, each iteration is executed on the processor containing the data that is mapped to it. The user was allowed to specify this.

In addition, the private variable PRIVATE\_C is used to compute a total for each processor. At the end of execution of the loop, the values of PRIVATE\_C may be different on each processor depending upon the values in the elements of the array on each processor. This data may be used as is, or it can be quickly summed using a barrier or an ATOMIC UPDATE.

Example 3 shows the final total value being combined into the variable MAPPED\_C whose value is available to all processors.

```
! Example 3
```

```
MAPPED_C = 0
!HPF$ ATOMIC UPDATE
MAPPED_C = MAPPED_C + PRIVATE_C
```

Example 4 shows how the language allows private data to vary from processor to processor.

```
43 ! Example 4
44
45 IF (MY_PE() .EQ. 5) THEN
46 PRIVATE_C = some-big-expression
47 ENDIF
48
```

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In this example, PRIVATE\_C on processor 5 will have the result of *some-big-expression*. Each processor can do distinctly different work and communicate through mapped data.

The code fragment in Example 5 is from an application and shows a few features of the language.

```
Example 5
!
!HPF$ GEOMETRY G(*, CYCLIC)
      REAL FX(100,100), FY(100,100), FZ(100,100)
!HPF$ DISTRIBUTE (G) :: FX,FY,FZ
      REAL FXP(100,16,100), FYP(100,16,100)
!HPF$ DISTRIBUTE FXP(*,*, BLOCK) FYP(*,*, BLOCK)
      INTEGER CELL, ATOM, MAP(1000), NACELL(1000)
!HPF$ INDEPENDENT (CELL) ON FX(1,CELL)
      DO CELL=1,100
         JCELLO = 16*(CELL-1)
         DO NABOR = 1, 13
           JCELL = MAP(JCELLO+NABOR)
           DO ATOM=1, NACELL(CELL)
              FX(ATOM, CELL) = FX(ATOM, CELL) + FXP(ATOM, NABOR, JCELL)
              FY(ATOM, CELL) = FY(ATOM, CELL) + FYP(ATOM, NABOR, JCELL)
           ENDDO
         ENDDO
      ENDDO
```

The GEOMETRY directive allows the user to generically specify a mapping and use it to apply to many arrays (they need not have the same extents.)

Example 5 has a single INDEPENDENT loop which is the outer loop. It executes 100 iterations total. Within this loop the private value of JCELLO is set for each processor (ensuring that it is a local computation everywhere.) Nested inside the INDEPENDENT loop is a private loop; this loop executes 13 times *per* processor. Inside this loop JCELL is computed locally on each processor, minimizing unnecessary communication. Finally the innermost loop is also private.

#### F.3 External Interface

This section describes the behavior when an HPF\_CRAFT routine is called from HPF.

The calling convention and argument passing rules for HPF\_CRAFT are a hybrid of those for HPF calling HPF\_LOCAL and HPF calling HPF. Explicit interfaces are required. Where dummy arguments are private (default) storage, the HPF calling HPF\_LOCAL conventions are used. Where dummy arguments are explicitly mapped, the calling convention matches HPF calling HPF.

There are a number of constraints on HPF\_CRAFT routines that are called from HPF. The following is a list of restrictions placed on HPF\_CRAFT routines called from HPF:

• Recursive HPF\_CRAFT routines cannot be called from HPF.

1 2	• HPF_CRAFT routines called from HPF may only enter the routine at a single place (no alternate entries).
3 4 5 6	• An HPF_CRAFT supprogram may not be invoked directly or indirectly from within the body of a FORALL construct or within the body of an INDEPENDENT DO loop that is inside an HPF program.
7 8 9 10	• The attributes (type, kind, rank, optional, intent) of the dummy arguments in a supprogram called by HPF must match the attributes of the corresponding dummy arguments in the explicit interface.
11	• A dummy argument of an HPF_CRAFT supprogram called by HPF
12	must not he a presedure norma
13	- must not be a procedure name.
14	<ul> <li>must not have the POINTER attribute.</li> </ul>
15	<ul> <li>must not be sequential, unless it is also PE_PRIVATE.</li> </ul>
16	– must have assumed shape even when it is explicit shape in the interface.
17 18	- if scalar, it must be mapped so that each processor has a copy of the argument.
19 20	• The default mapping of scalar dummy arguments and of scalar function results when an HPF program calls an HPF_CRAFT routine is that it is replicated on each pro-
21	cessor.
22	
23	If a dummy argument of an EXTRINSIC('HPF_CRAFT') routine interface block is an
24	array and the dummy argument of the HPF_CRAFT supprogram has the default private
25	mapping, then the corresponding dummy argument in the specification of the $\mathrm{HPF\_CRAFT}$
26	procedure must be an array of the same rank, type, and type parameters. When the extrinsic
27	procedure is invoked, the dummy argument is associated with the local array that consists
28	of the subgrid of the global array that is stored locally.
0.0	If the dummy argument of the HPE CPAET suppression is explicitly mapped, it must

If the dummy argument of the HPF\_CRAFT supprogram is explicitly mapped, it must have the same mapping as the dummy argument of the EXTRINSIC('HPF\_CRAFT') supprogram. Note that this restriction does not require actual and dummy arguments to match and is no more stringent than saying that mappings of dummy arguments in interface blocks must match those in the actual routine.

F.4 Execution Model

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HPF\_CRAFT is built upon the fundamental execution model of HPF\_LOCAL, augmented
 with data mapping and work distribution features from HPF. It is also augmented with
 explicit low-level control features, many taken from Cray Research's CRAFT language.

In HPF\_CRAFT there is a single task on each processor and all tasks begin executing in parallel, with data defaulting to a private distribution, the same default distribution used in HPF\_LOCAL. Each processor gets a copy of the data storage unless specified otherwise by the user. Consequently I/O works identically to I/O in HPF\_LOCAL and message passing libraries are easily integrated.

<sup>45</sup> Simply stated, the execution model is that of HPF\_LOCAL.

To provide correct behavior when explicitly mapped data is involved, this model defines implicit barrier points at which the execution model requires that all processors must stop and wait for the execution of all other processors before continuing. These barriers add

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additional semantics to the HPF\_LOCAL behavior. An implementation may remove any 1 of these barriers that are deemed unnecessary, but every processor must participate in the 2 barriers at each one of these points.

The points where there are implicit barriers are conceptually after those instances in which the processors in the HPF\_CRAFT program are executing cooperatively, as if in an HPF program (e.g., after an INDEPENDENT loop). An HPF\_CRAFT program treats operations on explicitly mapped objects as if they were operations in an HPF program and it treates operations on private data as if they were executed within the HPF\_LOCAL framework. It is occasionally useful for an advanced programmer to indicate to the compilation system where barriers are not needed; HPF\_CRAFT has syntax to allow this capability.

#### F.5**HPF\_CRAFT** Functional Summary

HPF\_CRAFT contains a number of features not available in HPF, and restricts the usage of many of the features currently available. The following is a concise list of the differences.

- INDEPENDENT has been extended to better support an ON clause.
- There are new rules defining the interaction of explicitly mapped and private data.
- Parallel inquiry intrinsics IN\_PARALLEL() and IN\_INDEPENDENT() have been added.
- Serial regions (MASTER / END MASTER) have been added.
- Explicit synchronization primitives are provided.
- The ATOMIC UPDATE, SYMMETRIC, and GEOMETRY directives have been added.
- Many other compiler information directives have been added to assist the compiler in producing good quality code.

#### F.5.1 **Data Mapping Features**

Data mapping features provided are those that have been found useful most often. When data is explicitly mapped, only one copy of the data storage is created unless the explicit mapping directs otherwise. The value of explicitly mapped replicated data items must be consistent between processors as is the case in HPF. Storage and sequence association for explicitly mapped arrays is not guaranteed in HPF\_CRAFT. For private data, storage and sequence association follows the Fortran 90 rules.

A new directive is included for completeness: **PE\_PRIVATE**, which specifies that the data should conform to the default behavior. The values of private varables may vary on different processors.

#### F.5.2 Subprogram Interfaces

The behavior and requirements of an HPF\_CRAFT program at subprogram interfaces may 43 be divided into three cases. Each case is also available using some combination of HPF and 44 HPF\_LOCAL. For dummy arguments that are explicitly mapped, the behavior is identical 45to that of HPF. All processors must cooperate in a subprogram invocation that remaps or 46 explicitly maps data. In other words, if an explicit interface is required (by the HPF rules) 47or the subprogram declares explicitly mapped data, the subprogram must be called on all 48 processors. Processors need not cooperate if there are only reads to non-local data. The
 INHERIT attribute may only be applied to explicitly mapped data.

<sup>3</sup> Data that has the default private mapping (case two) the behavior of an HPF\_CRAFT <sup>4</sup> subprogram at subprogram interfaces is identical to that of HPF\_LOCAL. Data is passed <sup>5</sup> individually on every processor and the processors need not interact in any way.

6 When a subprogram is passed actual arguments that are a combination of both explic-7 itly mapped data and private data, the explicitly mapped data follows the HPF rules and 8 the private data follows the HPF\_LOCAL rules.

In case three, the user has the option of passing data with explicitly mapped actual 9 arguments to dummy arguments that are not explicitly mapped (i.e., private.) The mapping 10  $1\,1$ rules for this data are identical to the mapping rules when HPF calls an HPF\_LOCAL subprogram. The data remains "in-place." All HPF arrays are logically carved up into 12pieces; the HPF\_CRAFT procedure executing on a particular physical processor sees an 13 array containing just those elements of the global array that are mapped to that physical 14processor. There is implicit barrier synchronization after an INDEPENDENT loop. Transfer 1516of control into or out of an INDEPENDENT loop is prohibited.

Finally, it is undefined behavior when an actual argument is private and the dummy argument is explicitly mapped. A definition could be supplied for this interaction, but it is the same solution that one might propose for a calling sequence when HPF\_LOCAL subprograms call HPF subprograms.

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### F.5.3 The INDEPENDENT Directive

The INDEPENDENT directive is part of HPF\_CRAFT with the same semantics as in HPF. However, within INDEPENDENT loops the values of private data may vary from processor to processor. INDEPENDENT applied to FORALL has identical syntax and semantics as in HPF.

An HPF independent loop optionally may have a NEW clause. The NEW clause is not required by HPF\_CRAFT for default (not explicitly mapped) data. In HPF\_CRAFT data defaults to private so values may differ from processor to processor.

Private data has slightly different behavior than data specified in the NEW clause. The value of a private datum on each processor can be used beyond a single iteration of the loop. Private data may be used to compute local sums, for example. The values of data items named in a NEW clause may not be used beyond a single iteration. The NEW clause asserts that the INDEPENDENT directive would be valid if new objects were created for the variables named in the clause for each iteration of the loop. The semantics of the NEW clause are identical in HPF\_CRAFT and HPF.

The semantics of an INDEPENDENT applied to loops containing private data references changes with respect to the private data. The change can be summarized to say that instead of indicating that iterations have no dependencies upon one-another, with respect to the private data, iterations on different processors have no dependencies upon one-another.

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#### F.5.4 The ON Clause

In addition to the version of INDEPENDENT available from HPF, a new version of INDEPENDENT
 is included that incorporates the ON clause. There are a number of differences between the
 versions of INDEPENDENT with and without the ON clause.

The new version of the INDEPENDENT directive may be applied to the first of a group of tightly nested loops and may apply to more than one of them. This more easily facilitates

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the use of the ON clause. The current INDEPENDENT directive applies only to a single loop nest. The INDEPENDENT directive is extended so that multiple loop nests can be named. The general syntax for these new independent loops is as follows:

```
\begin{array}{l} \texttt{!HPF\$ INDEPENDENT} \; (I_1,I_2,\ldots,I_n) \; \texttt{ON} \; array\text{-}name(h_1(I_1),h_2(I_2),\ldots,h_n(I_n)) \\ \texttt{DO} \; I_1 = L_1, \; U_1, \; S_1 \\ \texttt{DO} \; I_2 = L_2, \; U_2, \; S_2 \\ & \cdots \\ \texttt{DO} \; I_n = L_n, \; U_n, \; S_n \\ & \cdots \\ \texttt{END} \; \texttt{DO} \\ & \cdots \\ \texttt{END} \; \texttt{DO} \\ \texttt{END} \; \texttt{DO} \\ \texttt{END} \; \texttt{DO} \\ \texttt{END} \; \texttt{DO} \end{array}
```

The syntax and semantics of INDEPENDENT with the ON clause are different from its syntax and semantics without the ON clause. With the ON clause the directive states that there are no cross-processor dependencies, but there may be dependencies between iterations on a processor. There is an implicit barrier synchronization after an INDEPENDENT loop. Transfer of control into or out of an INDEPENDENT loop is prohibited.

The iteration space of an INDEPENDENT nest must be rectangular. That is, the lower loop bound, the upper loop bound, and the step expression for each loop indicated by the INDEPENDENT induction list must be invariant with regard to the INDEPENDENT nest. Each index expression of *array-name* in the ON clause (the functions  $h_i$  above,) must be one of the following two forms:

```
[ a * loop_control_variable + ] b
[ a * loop_control_variable - ] b
```

where a and b must be integer values; they can be expressions, constants, or variables. The values of a and b must be invariant with regard to the INDEPENDENT loop nest. For example, specifying A(I,J,K) is valid. Specifying A(3,I+J,K) is not valid. Specifying A(I,I,K) is not valid because I appears twice. Division is prohibited in any index expression of the ON clause.

### F.5.5 Array Syntax

Array syntax is treated identically in HPF\_CRAFT as in HPF for explicitly mapped objects. For private objects the behavior is identical to that of HPF\_LOCAL. When private objects and explicitly mapped objects are combined the rules are as follows:

```
result = rhs_1 \text{ op}_1 rhs_2 \text{ op}_2 \dots \text{ op}_m rhs_n
```

- If *result* is explicitly mapped and all *rhs* arrays are explicitly mapped, the work is distributed as in HPF.
- If *result* is private and all *rhs* arrays are private the computation is done on all processors as an HPF\_LOCAL program would do it.
- If *result* is private and all *rhs* arrays are explicitly mapped, the work is distributed as
   <sup>47</sup> in HPF and the values of the results are broadcast to the *result* on each processor.

- If *result* is explicitly mapped and *not* all *rhs* arrays are explicitly mapped, the results of the operation are undefined, unless all corresponding elements of all private *rhs* arrays have the same values.
- If *result* is private and some, but not all *rhs* arrays are explicitly mapped, the value is computed on each processor and saved to the local *result*.

All processors must participate in any array syntax statement in which the value of an explicitly mapped array is modified, and there is implicit barrier synchronization after the statement executes.

F.5.6 Treatment of FORALL and WHERE Statements

13 The FORALL and WHERE statements are treated exactly as in HPF when data is explicitly 14mapped. When private data is modified, the statement is executed separately on each pro-15cessor. Finally, when data in a FORALL or WHERE are mixed, the rules for array syntax apply. 16If any explicitly mapped data item is modified in a *forall-stmt* or *where-stmt* then arrays in 17the *forall-header* or *where-header* must be explicitly mapped. In a **FORALL** construct, if any 18 explicitly mapped array is modified, all modified arrays must be explicitly mapped. There 19is an implicit barrier synchronization after FORALL and WHERE statements if any arrays in 20 the forall-header or where-header are explicitly mapped.

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### F.5.7 Synchronization Primitives

A number of synchronization primitives are provided. These primitives include:

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Barriers (test, set, wait) Locks ( test, set, clear) Critical Sections

Events (test, set, wait, clear)

<sup>30</sup> Barriers provides an explicit mechanism for a task to indicate its arrival at a program <sup>31</sup> point and to wait there until all other tasks arrive. A task may test and optionally wait <sup>32</sup> at an explicit barrier point. In the following example, a barrier is used to make sure that <sup>33</sup> block3 is not entered by any task until all tasks have completed execution of block1.

```
<sup>34</sup>
<sup>35</sup> block1
<sup>36</sup> CALL SET_BARRIER()
<sup>37</sup> block2
<sup>38</sup> CALL WAIT_BARRIER()
<sup>39</sup> block3
```

The following example performs a similar function as above. However, while waiting for all tasks to arrive at the barrier, the early tasks perform work within a loop.

```
43 block1
44 CALL SET_BARRIER()
45 DO WHILE (.NOT. TEST_BARRIER())
46 block2
47 END DO
48 block3
```

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Locks are used to prevent the simultaneous access of data by multiple tasks.

The SET\_LOCK(*lock*) intrinsic sets the mapped integer variable *lock* atomically. If the lock is already set, the task that called SET\_LOCK is suspended until the lock is cleared by another task and then sets it. Individual locks may be tested or cleared using *result* = TEST\_LOCK(*lock*) and CLEAR\_LOCK(*lock*) respectively.

A *critical section* protects access to a section of code rather than to a data object. The CRITICAL directive marks the beginning of a code region in which only one task can enter at a time. The END CRITICAL directive marks the end of the critical section. Transfer of control into or out of a critical section is prohibited.

```
!HPF$ CRITICAL
    GLOBAL_SUM = GLOBAL_SUM + LOCAL_SUM
!HPF$ END CRITICAL
```

Events are typically used to record the state of a program's execution and to communicate that state to another task. Because they do not set locks, as do the lock routines described earlier, they cannot easily be used to enforce serial access of data. They are suited to work such as signalling other tasks when a certain value has been located in a search procedure. There are four routines needed to perform the event functions, and each requires a mapped argument.

The SET\_EVENT(*event*) routine sets or *posts* an event; it declares that an action has been accomplished or a certain point in the program has been reached. A task can post an event at any time, whether the state of the event is cleared or already posted. The CLEAR\_EVENT(*event*) routine clears an event, the WAIT\_EVENT(*event*) routine waits until a particualr event is posted, and the *result* = TEST\_EVENT(*event*) function returns a logical value indicating whether a particular event has been posted.

### F.5.8 Barrier Removal

You can explicitly remove an implicit barrier after any INDEPENDENT loop, or after any array syntax statement that modifies explicitly mapped arrays, by using the NO BARRIER directive.

```
!HPF$ NO BARRIER
```

### F.5.9 Serial Regions

It is often useful to enter a region where only one task is executing. This is particularly useful for certain types of I/O. To facilitate this, two directives are provided. In addition, one may optionally attach a COPY clause to the END MASTER directive which specifies the private data items whose values should be broadcast to all processors. The syntax of this directive is:

```
      !HPF$ MASTER
      45

      sequential region
      46

      ...
      47

      !HPF$ END MASTER [, COPY( var1 [, var2, ..., varn ])]
      48
```

where var is SYMMETRIC private data to be copied to the same named private data on other 1 processors. 2

3 If a routine is called within a serial region, the routine executes serially; there is no way to get back to parallel execution within the routine. All explicitly mapped data is 4 accessible from within routines called in a serial region, but a routine called from within 5 a serial region cannot allocate explicitly mapped data or remap data. All processors must 6 participate in the invocation of the serial region. Transfer of control into or out of a serial 7 region is not permitted. 8

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F.5.10 Libraries

 $1\,1$ The HPF Local Routine Library is available in HPF\_CRAFT. The HPF\_LOCAL extrinsic 12environment contains a number of libraries that are useful for local SPMD programming and 13 a number of libraries that allow the user to determine global (rather than local) state infor-14 mation. These library procedures take as input the name of a dummy argument and return 15information on the corresponding global HPF actual argument. They may only be invoked 16 by an HPF\_CRAFT procedure that was directly invoked by global HPF code. They may 17be called only for private data. The libraries reside in a module called HPF\_LOCAL\_LIBRARY. 18

The HPF Library is available to HPF\_CRAFT when called with data that is explicitly 19 mapped and all processors are participating in the call. In addition, as in HPF\_LOCAL, 20 the entire HPF Library is available for use with private data. Mixing private and explicitly 21 mapped data in calls to the HPF library produces undefined behavior.

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#### F.5.11 **Parallel Inquiry Intrinsics**

These intrinsic functions are provided as an extension to HPF. They return a logical value that provides information to the programmer about the state of execution in a program.

> IN\_PARALLEL() IN\_INDEPENDENT()

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#### F.5.12 Task Identity

MY\_PE() may be used to return the local processor number. The physical processors are identified by an integer in the range of 0 to n-1 where n is the value returned by the 34 global HPF\_LIBRARY function NUMBER\_OF\_PROCESSORS. Processor identifiers are returned by ABSTRACT\_TO\_PHYSICAL, which establishes the one-to-one correspondence between the abstract processors of an HPF processors arrangement and the physical processors. Also, the local library function MY\_PROCESSOR returns the identifier of the task executing the call.

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#### F.5.13 **Parallelism Specification Directives**

41These directives allow a user to assert that a routine will only be called from within a 42 parallel region, a serial region, or from within both regions. Without these directives an 43 implementation might be required to generate two versions of code for each routine, de-44 pending upon implementation strategies. The directives simply make the generated code 45size smaller and remove a test. 46

47!HPF\$ PARALLEL\_ONLY

!HPF\$ SERIAL\_ONLY 48

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The RESIDENT directive can be specified at the loop level and at the routine level. It is an assertion that the references to particular variables in the routine (or loop) are only references to data that are local to the task making the assertion. In the following loop, all references to arrays A, B, and C are local to the task executing each iteration.

### F.5.16 The ATOMIC UPDATE Directive

END DO

In HPF\_CRAFT, the ATOMIC UPDATE directive tells the compiler that a particular data item or the elements of a particular array for a specified operation must be updated atomically. This can be used within loops or in array syntax and applies to both the elements of an array with an assignment of a permutation and the elements of an array within a loop.

In the following example, all references to R(IX(I)) occur atomically, thus eliminating the possibility that different iterations might try to modify the same element concurrently.

```
REAL R(200), S(1000)
INTEGER IX(1000)
!HPF$ DISTRIBUTE R(BLOCK), S(BLOCK), IX(BLOCK)
!HPF$ INDEPENDENT (I) ON S(I)
DO I = 1, 1000
!HPF$ ATOMIC UPDATE
```

```
R(IX(I)) = R(IX(I)) + S(I)
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             END DO
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3
               The GEOMETRY Directive
      F.5.17
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      The GEOMETRY directive is similar to a typedef in C, only it is for data mapping. It allows
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      the user to conveniently change the mappings of many arrays at the same time. It is similar
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      in many ways to the TEMPLATE directive, but since it is bound to no particular extent it is
8
      sometimes easier to apply.
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      !HPF$ GEOMETRY geom(d_1 [, d_2, ..., d_n])
11
      !HPF$ DISTRIBUTE ( geom ) [::] var_1[, var_2, ..., var_m]
12
      Where d_i indicates one of the allowable distribution formats.
13
14
      !HPF$ GEOMETRY GBB(BLOCK, CYCLIC)
15
             REAL A(300,300), B(400,400)
16
      !HPF$ DISTRIBUTE (GBB) :: A, B
17
             if GBB changes then both {\tt A} and {\tt B} change
      !
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```

Annex G

# The FORTRAN 77 Local Library

14The HPF standard now describes an EXTRINSIC(LANGUAGE='F77', MODEL='LOCAL') interface, or EXTRINSIC (F77\_LOCAL) to use the keyword identification (see Section 11.6 for its 1516description), similar in characteristics to the EXTRINSIC(LANGUAGE='HPF', MODEL='LOCAL') 17and EXTRINSIC (LANGUAGE='FORTRAN', MODEL='LOCAL') interfaces. This section describes 18 a set of library routines to make it easier to make use of the F77\_LOCAL interface when pass-19 ing distributed array data. These library routines can facilitate, for example, a portable 20 blend of global data parallel code with preexisting FORTRAN 77-based code using explicit 21 message passing calls for interprocessor communication. The FORTRAN 77 Local Library 22 interface described in this section was originally developed as part of Thinking Machines 23 TMHPF and is now supported by Sun Microsystems Inc. For suggestions, requests, or corrections concerning this interface, please contact 24

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Sun Microsystems Inc.

- 27 High Performance Computing
- <sup>28</sup> M/S UCHLO5-104
- <sup>29</sup> 5 Omni Way
- <sup>30</sup> Chelmsford, MA 01824
- 31 f77-local-library@sun.com

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### G.1 Introduction

The basic constraints for the local model (Section 11.1) together with the F77\_LOCAL-specific 36 argument passing options (Section 11.6) define the nature of the F77\_LOCAL interface: how 37 control is to be transferred from a global HPF procedure to a set of local procedures de-38 scribed by an EXTRINSIC (F77\_LOCAL) procedure interface and how data can be passed be-39 tween these two types of procedures: by reference or by descriptor, and with or without tem-40 porary local reordering of data to satisfy FORTRAN 77 provisions for sequential, contiguous 41 storage of array data in Fortran array element order. These alternative methods of argument 42 passing can be obtained by use of the two special-purpose attributes for extrinsic dummy 43 arguments defined for LANGUAGE='F77' routines: LAYOUT('F77\_ARRAY') (the default) vs. 44 LAYOUT('HPF\_ARRAY'), and PASS\_BY('\*') (the default) vs. PASS\_BY('HPF\_HANDLE'). 45However, to take advantage of the option allowing one to pass global HPF array "handles" 46 to local FORTRAN 77 procedures and then obtain information locally about how the local 47portion of a given parallel array is actually distributed requires special inquiry routines 48

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comparable to the HPF Local Library of functions. Since this library is not only described 1 as a module, but uses many features such as array-valued functions and optional arguments 2 not available in FORTRAN 77 code, it is recommended that a modified FORTRAN 77 3 interface to this library be provided in the manner described below. Furthermore, there is 4 the problem of describing local portions of parallel arrays in the FORTRAN 77 code used 5 in each local routine called from a global HPF one. Since assumed-shape syntax may not 6 be used, explicit shape arrays are required. But it is common for global distribution of 7arbitrary sized arrays to result in local portions of arrays that do not have constant shapes 8 on all processors, and the actual extents in each processor cannot necessarily be predicted 9 in advance. In order to allow programmers to obtain axis extent information at run time 10 from the HPF global caller, a special HPF-callable subgrid inquiry subroutine is provided.  $1\,1$ A FORTRAN 77 callable version of the same routine is also described below, for flexibility 12in programming. 13

#### G.2 Summary

• One HPF-callable subgrid inquiry subroutine HPF_SUBGRID_INFO	
• A set of FORTRAN 77-callable inquiry subroutines F77_SUBGRID_INFO	
F77_GLOBAL_ALIGNMENT	
F77_GLOBAL_DISTRIBUTION	
F77_GLOBAL_TEMPLATE	
F77_ABSTRACT_TO_PHYSICAL	
F77_PHYSICAL_TO_ABSTRACT	
F77_LOCAL_TO_GLOBAL	
F77_GLOBAL_TO_LOCAL	
F77_LOCAL_BLKCNT	
F77LOCALLINDEX	

F77\_SIZE F77\_MY\_PROCESSOR

F77\_LOCAL\_UINDEX

F77\_GLOBAL\_SHAPE

F77\_GLOBAL\_SIZE

F77\_SHAPE

### G.3 Global HPF Subgrid Inquiry Routine

The F77\_LOCAL library interface includes only one global HPF subroutine, HPF\_SUBGRID\_INFO, whose implementation should be added as an extension to the standard HPF Library module. Its purpose is to provide per-processor information about the local subgrids of distributed arrays. This information is often critical when passing such arrays to local procedures written in FORTRAN 77, where array argument shapes must be stated explicitly 48 in the local procedure (except in the last dimension; there are "assumed size" but no "assumed shape" arrays), but may be expressed in terms of arguments passed at run time ("adjustable shape arrays"). Thus the subgrid parameters obtained from this subgrid inquiry routine can be passed as arguments to the local routines and used there to describe the extents of the locally visible portions of global HPF arrays, as the example in Section G.5 will demonstrates.

# HPF\_SUBGRID\_INFO (ARRAY, IERR, DIM, LB, UB, STRIDE, LB\_EMBED, UB\_EMBED, AXIS\_MAP)

- **Description.** Gives local information about local subgrid allocation onto each processor of a distributed array; callable from a global HPF routine.
  - **Class.** Inquiry subroutine.
- <sup>15</sup> Arguments.

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- 16ARRAY is a nonsequential array of any type, size, shape, or map-17ping. It is an INTENT (IN) argument. 18 19 IERR is a scalar integer of default kind. It is an INTENT (OUT) 20 argument. Its return value is zero upon successful return 21 and nonzero otherwise. Errors result if local subgrids 22 cannot be expressed as array sections of ARRAY. 23
  - If any of the optional arguments LB\_EMBED, UB\_EMBED, or AXIS\_MAP is present, then a nonzero value is also returned if the compiler does not organize the local data in serial memory by sequence associating a larger "embedding" array (see Section G.3.1 below for more explanation).
- 29DIM (optional)is a scalar integer of default kind. It is an INTENT (IN)30argument. DIM indicates the axis along which return val-31ues are desired. If DIM is not present, values are returned32for all axes.
- LB (optional) is an INTENT (OUT), default integer array. If this argument is present, and if the value returned in IERR is zero, the values returned in array LB are the lower bounds in global coordinates of each processor's subgrid, along one (if DIM is present) or each dimension of ARRAY.
- UB (optional) is an INTENT (OUT), default integer array. If this argument is present, and if the value returned in IERR is zero, the values returned in array UB are the upper bounds in global coordinates of each processor's subgrid, along one (if DIM is present) or each dimension of ARRAY.
- 44STRIDE (optional)is an INTENT (OUT), default integer array. If this argument is present, and if the value returned in IERR is zero,45ment is present, and if the value returned in IERR is zero,46the values returned in array STRIDE are the strides in lo-47cal memory between elements of each processor's subgrid,48along one (if DIM is present) or each dimension of ARRAY.

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LB_EMBED (optional)	is an INTENT (OUT), default integer array. If this ar- gument is present, and if the value returned in IERR is zero, the values returned in array LB_EMBED are the lower bounds in global coordinates of the actual global array elements allocated on each processor, possibly a superset of the user-visible subgrid, along one (if DIM is present) or each dimension of ARRAY.	1 2 3 4 5 6 7
UB_EMBED (optional)	is an INTENT (OUT), default integer array. If this ar- gument is present, and if the value returned in IERR is zero, the values returned in array UB_EMBED are the upper bounds in global coordinates of the actual global array el- ements allocated on each processor, possibly a superset of the user-visible subgrid, along one (if DIM is present) or each dimension of ARRAY.	8 9 10 11 12 13 14 15
AXIS_MAP (optional)	is a rank 2, INTENT (OUT), default integer array. If this argument is present, its shape must be at least [n,r], where n is the number of processors and r is the rank of ARRAY.	16 17 18 19
	If the value returned in IERR is zero, the values returned in $AXIS\_MAP(i,1:r)$ represent the numbers of the axes of the subgrid on processor i from fastest varying to slowest varying, and form a permutation of the sequence $1,2,\ldots,r$ .	20 21 22 23 24 25

For the last six arguments, LB, UB, STRIDE, LB\_EMBED, UB\_EMBED, and AXIS\_MAP, each array has a first axis of extent at least n, where n is the number of processors, and the first n indices of that axis of each array must be distributed (perhaps via an explicit CYCLIC or BLOCK distribution) one index per processor. If a second dimension is needed, it should be a collapsed axis of extent at least equal to the rank of ARRAY.

If HPF\_SUBGRID\_INFO is called, and the elements of ARRAY that are local to any particular processor are not representable as an array section of the global user array, then a nonzero value is returned for IERR. Otherwise, if any of the optional arguments LB, UB, or STRIDE is present, then the lower bounds, upper bounds, or strides, respectively, that describe the local array sections are returned in terms of one-based, global coordinates.

#### G.3.1 Subgrid Inquiries Involving Embedding Arrays

In the common case in which the elements of each local subgrid of the global array argument are distributed across processors, with no overlap, and allocated in local memory like a local FORTRAN 77 array, as a contiguous sequence of elements in Fortran array element order, these three last optional arguments would not be required.

However, some implementations may choose less common layouts in local memory, that involve "embedding" these elements in a larger array section of equal rank that *is* sequence-associated in serial memory. For example, alignment of axes of arrays in different orders may result in a permuting embedding of the subgrid. Or axes of subgrids map be padded with ghost cells, either for stencil optimizations or to achieve same-size subgrids on all nodes. In variations such as these, we may still view the subgrid as being "embedded" in a sequence associated array which may be accessible in F77\_LOCAL operations, if the permutation of axes, shape of any embedding array, and offsets into that array can be obtained at runtime. The last three arguments of HPF\_SUBGRID\_INFO are provided to allow programmers to obtain this information when it is appropriate, with the help of the IERR flag to signal when this is not the case.

In this mapping, local memory has been allocated for a larger array section, with coordinates (LB\_EMBED : UB\_EMBED : STRIDE). The coordinates of the *actual* computational elements are limited to the subset (LB : UB : STRIDE). The sequence association is generalized to an arbitrary mapping of axes. Here, AXIS\_MAP numbers the axes from fastest varying to slowest varying. If LB\_EMBED, UB\_EMBED, or AXIS\_MAP is specified in a call to HPF\_SUBGRID\_INFO but ARRAY does not satisfy the assumptions of this mapping model, then a nonzero value is returned for IERR.

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#### G.4 Local FORTRAN 77 Inquiry Routines

Here the F77-callable inquiry subroutines are described briefly. These provide essentially
 the same capability as the combination of the HPF intrinsic array inquiry functions such
 as SHAPE and SIZE, together with the HPF LOCAL LIBRARY inquiry routines. The subrou tine F77\_SUBGRID\_INFO serves as a local counterpart to the globally callable subroutine
 HPF\_SUBGRID\_INFO described above. In all of the following:

- ARRAY is a dummy argument passed in from a global HPF caller using the LAYOUT ('HPF\_ARRAY') attribute and declared within the FORTRAN 77 local subroutine as a scalar integer variable. It is an INTENT (IN) argument.
- DIM is a scalar integer of default kind. It is an INTENT (IN) argument. This argument specifies a particular axis of the global array associated with ARRAY or, if DIM = -1, inquiry is for all axes.
- An "inquiry result" is an INTENT (OUT) argument. If DIM = -1, it is a rank-one array of size equal to at least the rank of the global array associated with ARRAY, returning information associated with all axes. If DIM is positive, the "inquiry result" is a scalar, returning information only for the axis indicated by DIM.
- The arguments are defined in the same way as for the corresponding HPF or HPF\_LOCAL routines unless otherwise noted. See the description of HPF\_SUBGRID\_INFO above and Section 11.7.1 for full specifications of the similarly-named HPF\_LOCAL\_LIBRARY procedures.

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# F77\_SUBGRID\_INFO (ARRAY, IERR1, IERR2, DIM, LB, UB, STRIDE, LB\_EMBED, UB\_EMBED, AXIS\_MAP)

**Description.** This is a FORTRAN 77-callable version of the HPF subroutine HPF\_SUBGRID\_INFO.

Arguments.

46IERR1is a scalar integer of default kind. It is an INTENT (OUT)47argument. Its return value is zero if LB, UB, and STRIDE48were determined successfully and nonzero otherwise.

IERR2	is a scalar integer of default kind. It is an INTENT (OUT) argument. Its return value is zero if LB_EMBED and UB_EMBED were determined successfully and nonzero otherwise.	1 2 3 4	
LB, UB, STRIDE, LB_EMB	ED, UB_EMBED, AXIS_MAP are "inquiry results" of default integer type. They are the lower and upper bounds and strides of the array sections describing the local data (in terms of global indices), the lower and upper bounds of the embedding arrays (again, in terms of global indices), and the axes of the embedding arrays to which the axes of ARRAY are mapped.	5 6 7 8 9 10 11 12	
F77_GLOBAL_ALIGNME IDENTITY_MAP, DYNAR	NT (ALIGNEE, LB, UB, STRIDE, AXIS_MAP, MIC, NCOPIES)	13 14 15	
tine GLOBAL_ALIGNMENT. A	FORTRAN 77-callable version of the HPF_LOCAL subrou- ll but the first are INTENT (OUT) arguments whose return the corresponding HPF routine.	16 17 18 19	
Arguments.		20	
ALIGNEE	is a dummy argument passed in from global HPF. It is an INTENT (IN) argument.	21 22 23	
LB, UB, STRIDE, AXIS_M	AP are integer arrays of rank one. Their size must be at least equal to the rank of the global HPF array associated with ALIGNEE.	24 25 26	
IDENTITY_MAP, DYNAMIC	are scalar logicals.	27	
NCOPIES	is a scalar integer of default kind.	28 29	
	TION (DISTRIBUTEE, AXIS_TYPE, RS_RANK, PROCESSORS_SHAPE)	30 31 32	
GLOBAL_DISTRIBUTION. A	<b>Description.</b> This is a FORTRAN 77-callable version of the HPF_LOCAL subroutine GLOBAL_DISTRIBUTION. All but the first are INTENT (OUT) arguments whose return values are as specified by the corresponding HPF routine.		
Arguments.		37	
DISTRIBUTEE	is a dummy argument passed in from global HPF. It is an INTENT (IN) argument.	38 39 40	
AXIS_TYPE	is a CHARACTER*9 array of rank one. Its size must be at least equal to the rank of the global HPF array associated with DISTRIBUTEE.	41 42 43	
AXIS_INFO	is a default integer array of rank one. Its size must be at least equal to the rank of the global HPF array associated with <b>DISTRIBUTEE</b> .	44 45 46	
PROCESSORS_RANK	is a scalar of default integer type.	47 48	

1 2 3	PROCESSORS_SHAPE	is an integer array of rank one. Its size must be at least equal to the value returned by <b>PROCESSORS_RANK</b> .		
4		TE (ALIGNEE, TEMPLATE_RANK, LB, UB, O, NUMBER_ALIGNED, DYNAMIC)		
7 8 9	GLOBAL_TEMPLATE. All b	FORTRAN 77-callable version of the HPF_LOCAL subroutine ut the first are INTENT (OUT) arguments whose return values orresponding HPF routine.		
10	Arguments.			
12 13 14	ALIGNEE	is a dummy argument passed in from global HPF. It is an INTENT (IN) argument.		
15 16	TEMPLATE_RANK	is a scalar integer of default kind.		
17 18 19	LB, UB, AXIS_INFO	are integer arrays of rank one. Their size must be at least equal to the rank of the align-target to which the global HPF array associated with ALIGNEE is ultimately aligned.		
20 21 22 23 24	AXIS_TYPE	is a CHARACTER*10 array of rank one. Its size must be at least equal to the rank of the align-target to which the global HPF array associated with ALIGNEE is ultimately aligned.		
25	NUMBER_ALIGNED	is a scalar integer of default kind.		
26 27	DYNAMIC	is a scalar logical.		
	F77_ABSTRACT_TO_PE	IYSICAL(ARRAY, INDEX, PROC)		
30 31 32	<b>Description.</b> This is a ABSTRACT_TO_PHYSICAL.	FORTRAN 77-callable version of the HPF_LOCAL subroutine		
33 34	Arguments.			
35 36	INDEX	is a rank-one, INTENT (IN), integer array.		
37 38	PROC	is a scalar, INTENT (OUT), integer.		
39 40	F77_PHYSICAL_TO_AB	STRACT(ARRAY, PROC, INDEX)		
41 42 43	-	<b>Description.</b> This is a FORTRAN 77-callable version of the HPF_LOCAL subroutine PHYSICAL_TO_ABSTRACT.		
44 45	Arguments.			
46	PROC	is a scalar, INTENT (IN), integer.		
47 48	INDEX	is a rank-one, INTENT (OUT), integer array.		

$\mathbf{F77}_{-}$	LOCAL_TO_GLOBAL	(ARRAY, L_INDEX, G_INDEX)	1
	<b>Description.</b> This is a FORTRAN 77-callable version of the HPF_LOCAL subroutin LOCAL_TO_GLOBAL.		
	Arguments.		4 5
	L_INDEX	is a rank-one, INTENT (IN), integer array.	6
	G_INDEX	is a rank-one, INTENT (OUT), integer array.	7 8
	GLOBAL_TO_LOCAL PPIES, PROCS)	(ARRAY, G_INDEX, L_INDEX, LOCAL,	9 10 11
	<b>Description.</b> This is a FORTRAN 77-callable version of the HPF_LOCAL subrouger GLOBAL_TO_LOCAL.		
	Arguments.		15
	G_INDEX	is a rank-one, INTENT (IN), integer array.	16
	L_INDEX	is a rank-one, INTENT (OUT), integer array.	17 18
	LOCAL	is a scalar, INTENT (OUT), logical.	19
	NCOPIES	is a scalar, INTENT (OUT), integer.	20
	PROCS	is a rank-one, integer array whose size is at least the number of processors that hold copies of the identified element.	21 22 23 24
F77_LOCAL_BLKCNT(L_BLKCNT, ARRAY, DIM, PROC)			
	<b>Description.</b> This is a F LOCAL_BLKCNT.	ORTRAN 77-callable version of the HPF_LOCAL function	27 28 29
	Arguments.		30
	L_BLKCNT	is an "inquiry result" of type integer.	31
	PROC	is a scalar integer of default kind. It must be a valid	32 33
		processor number or, if $PROC = -1$ , the value returned	34
		by F77_MY_PROCESSOR() is implied.	35
$F77_$	LOCAL_LINDEX(L_L	INDEX, ARRAY, DIM, PROC)	36
		ORTRAN 77-callable version of the HPF_LOCAL function	37 38
	LOCAL_LINDEX.		39 40
	Arguments.		41
	L_LINDEX	is a rank-one, integer array of size equal to at least the value returned by F77_LOCAL_BLKCNT.	42 43
	DIM	may not be -1.	44 45
	PROC	is a scalar integer of default kind. It must be a valid processor number or, if $PROC = -1$ , the value returned	46 47
		by F77_MY_PROCESSOR() is implied.	48

<b>Description</b> . This	is a FORTRAN 77-callable version of the HPF_LOCAL function
LOCAL_UINDEX.	
Arguments.	
L_UINDEX	is a rank-one, integer array of size equal to at least the value returned by F77_LOCAL_BLKCNT.
DIM	may not be -1.
PROC	is a scalar integer of default kind. It must be a val processor number or, if PROC = -1, the value returne by F77_MY_PROCESSOR() is implied.
77_GLOBAL_SHAP	E(SHAPE, ARRAY)
<b>Description</b> . This GLOBAL_SHAPE.	is a FORTRAN 77-callable version of the HPF_LOCAL function
Arguments.	
SHAPE	is a rank-one, integer array of size equal to at least the rank of the global array associated with <b>ARRAY</b> . Its return value is the shape of that global array.
77_GLOBAL_SIZE(S	SIZE, ARRAY, DIM)
<b>Description</b> . This GLOBAL_SIZE.	is a FORTRAN 77-callable version of the HPF_LOCAL function
Arguments.	
SIZE	is a scalar integer equal to the extent of axis DIM of the global array associated with ARRAY or, if $DIM = -1$ , the total number of elements in that global array.
77_SHAPE(SHAPE,	ARRAY)
—	is a FORTRAN 77-callable version of the HPF intrinsic SHAF as called from HPF_LOCAL.
Arguments.	

F77_	SIZE(SIZE, ARRAY	, DIM)	1
	<b>Description</b> . This is a	FORTRAN 77-callable version of the HPF intrinsic SIZE,	2
	as it would behave as ca		3
			4 5
	Arguments.		6
	SIZE	is a scalar integer equal to the extent of axis DIM of the	7
		subgrid associated with ARRAY or, if $DIM = -1$ , the total	8
		number of elements in that subgrid.	9
			10
$\mathbf{F77}$	MY_PROCESSOR(M	IY_PROC)	11
Description. This is a FORTRAN 77-callable version of the HPF_LOCAL function		12 13	
	MY_PROCESSOR.		13 14
			15
	Arguments.		16
	MY_PROC	is a scalar, INTENT (OUT), integer. Its value is the iden-	17
		tifying number of the physical processor from which this	18
		call is made.	19
			20
G.5	Programming Exam	ple Using HPF_SUBGRID_INFO	21 22
C F	1 HPF Caller		23
G.5.	I HPF Caller		24
	PROGRAM EX	AMPLE	25
	! Declare the da	ta array and a verification copy	26
		ARAMETER :: NX = 100, NY = 100	27
		NSION(NX,NY) :: X, Y	28
		(BLOCK, BLOCK) :: X, Y	29
	0	will be computed	30
		tial sums on the processors	31
		AL_SUM(NUMBER_OF_PROCESSORS())	32
		PARTIAL_SUM(BLOCK) parameters are declared per processor	33 34
	! for a rank-two	· · ·	35
		IMENSION(NUMBER_OF_PROCESSORS(),2) ::	36
	& LB, UB, N		37
		(BLOCK,*) :: LB, UB, NUMBER	38
	! Define interfa		39
	INTERFACE		40
	EXTRINSI	C(F77_LOCAL) SUBROUTINE LOCAL1	41
	& ( LB1,	UB1, LB2, UB2, NX, X )	42
	•	1, LB2, UB2, and X are passed by default	43
		_ARRAY') and PASS_BY('*')	44
		DIMENSION(:) :: LB1, UB1, LB2, UB2	45
	INTEGER		46
	REAL X(:		47
	HPF\$ DISTRIBUTE	(BLOCK) :: LB1, UB1, LB2, UB2	48

```
!HPF$ DISTRIBUTE(BLOCK,BLOCK) :: X
1
                        END
2
3
                        EXTRINSIC(F77_LOCAL) SUBROUTINE LOCAL2(N,X,R)
                ! Arrays N, X, and R are passed by default
4
                ! as LAYOUT('F77_ARRAY') and PASS_BY('*')
5
                        INTEGER N(:)
6
                        REAL X(:,:), R(:)
7
                !HPF$ DISTRIBUTE N(BLOCK)
8
                !HPF$ DISTRIBUTE X(BLOCK, BLOCK)
9
                !HPF$ DISTRIBUTE R(BLOCK)
10
                        END
11
                      END INTERFACE
12
13
                ! Determine result using only global HPF
14
                      ! Initialize values
15
                      FORALL (I=1:NX, J=1:NY) X(I, J) = I + (J-1) * NX
16
                      ! Determine and report global sum
17
                      PRINT *, 'GLOBAL HPF RESULT: ',SUM(X)
18
                ! Determine result using local subroutines
19
                      ! Initialize values ( assume stride = 1 )
20
                      CALL HPF_SUBGRID_INFO( Y, IERR, LB=LB, UB=UB )
21
                      IF (IERR.NE.O) STOP 'ERROR!'
22
                      CALL LOCAL1( LB(:,1), UB(:,1), LB(:,2), UB(:,2), NX, Y )
23
                      ! Determine and report global sum
24
                      NUMBER = UB - LB + 1
25
                      CALL LOCAL2 ( NUMBER(:,1) * NUMBER(:,2) , Y , PARTIAL_SUM )
26
                      PRINT *, 'F77_LOCAL RESULT #1 : ',SUM(PARTIAL_SUM)
27
                      END
28
29
30
            FORTRAN 77 Callee
     G.5.2
31
                      SUBROUTINE LOCAL1( LB1, UB1, LB2, UB2, NX, X )
32
                ! The global actual arguments passed to LB1, UB1, LB2, and UB2
33
                ! have only one element apiece and so can be treated as scalars
34
                ! in the local Fortran 77 procedures
35
                      INTEGER LB1, UB1, LB2, UB2
36
                ! NX contains the global extent of the first dimension
37
                ! of the global array associated with local array X
38
                      INTEGER NX
39
                ! Note that X may have no local elements.
40
                      REAL X ( LB1 : UB1 , LB2 : UB2 )
41
                ! Initialize the elements of the array, if any
42
                      DO J = LB2, UB2
43
                        DO I = LB2, UB2
44
                          X(I,J) = I + (J-1) * NX
45
                        END DO
46
                      END DO
47
                      END
48
```

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13 14 15

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17 18

19

20 21

22

```
SUBROUTINE LOCAL2(N,X,R)
! Here, the rank of the original array is unimportant
! Only the total number of local elements is needed
! INTEGER N
    REAL X(N), R
! If N is zero, local array X has no elements, but R
! still computes the correct local sum
    R = 0.
    DO I = 1, N
        R = R + X(I)
        END DO
        END
```

#### G.6 Programming Example Using F77-Callable Inquiry Subroutines

This example performs only the initialization of the above example. It illustrates use of the F77-callable inquiry routines on descriptors passed from HPF, as well as the addressing of uncompressed local subgrid data in terms of "embedding arrays."

#### G.6.1 HPF Caller

```
PROGRAM EXAMPLE
                                                                        23
      INTEGER, PARAMETER :: NX = 100, NY = 100
                                                                        ^{24}
      REAL, DIMENSION(NX,NY) :: X
                                                                        ^{25}
!HPF$ DISTRIBUTE(BLOCK,BLOCK) :: X
                                                                        26
! Local subgrid parameters are declared per processor
                                                                        27
! for a rank-two array
                                                                        28
      INTEGER, DIMENSION(NUMBER_OF_PROCESSORS(),2) ::
                                                                        29
     & LB, UB, LB_EMBED, UB_EMBED
                                                                        30
!HPF$ DISTRIBUTE(BLOCK,*) :: LB, UB, LB_EMBED, UB_EMBED
                                                                        31
! Define interfaces
                                                                        32
      INTERFACE
                                                                        33
        EXTRINSIC(F77_LOCAL) SUBROUTINE LOCAL1(
                                                                        34
       LB1, UB1, LB_EMBED1, UB_EMBED1,
     &
                                                                        35
       LB2, UB2, LB_EMBED2, UB_EMBED2, X, X_DESC )
     ۶r.
                                                                        36
        INTEGER, DIMENSION(:) ::
                                                                        37
        LB1, UB1, LB_EMBED1, UB_EMBED1,
     &
                                                                        38
     &
         LB2, UB2, LB_EMBED2, UB_EMBED2
                                                                        39
! X is passed twice, both times without local reordering.
                                                                        40
! First, it is passed by reference for accessing array elements.
                                                                        41
        REAL, DIMENSION(:,:), LAYOUT('HPF_ARRAY'),
                                                                        42
                PASS_BY('*')
     &
                                                  :: X
                                                                        43
! It is also passed by descriptor for use in F77 LOCAL
                                                                        44
! LIBRARY subroutines only.
                                                                        45
        REAL, DIMENSION(:,:), LAYOUT('HPF_ARRAY'),
                                                                        46
                PASS_BY('HPF_HANDLE')
     &
                                                  :: X_DESC
                                                                        47
!HPF$ DISTRIBUTE(BLOCK) :: LB1, UB1, LB_EMBED1, UB_EMBED1
                                                                        48
```

```
!HPF$ DISTRIBUTE(BLOCK) :: LB2, UB2, LB_EMBED2, UB_EMBED2
1
                !HPF$ DISTRIBUTE(BLOCK,BLOCK) :: X
2
з
                        END
                      END INTERFACE
4
                ! Initialize values
5
                ! ( Assume stride = 1 and no axis permutation )
6
                      CALL HPF_SUBGRID_INFO( X, IERR,
7
                     & LB=LB, LB_EMBED=LB_EMBED,
8
                     & UB=UB, UB_EMBED=UB_EMBED)
9
                      IF (IERR.NE.O) STOP 'ERROR!'
10
                      CALL LOCAL1(
11
                     & LB(:,1), UB(:,1), LB_EMBED(:,1), UB_EMBED(:,1),
12
                     & LB(:,2), UB(:,2), LB_EMBED(:,2), UB_EMBED(:,2), X, X )
13
                      END
14
15
16
     G.6.2 FORTRAN 77 Callee
17
                      SUBROUTINE LOCAL1(
18
                     & LB1, UB1, LB_EMBED1, UB_EMBED1,
19
                     & LB2, UB2, LB_EMBED2, UB_EMBED2, X, X_DESC )
20
                      INTEGER LB1, UB1, LB_EMBED1, UB_EMBED1
21
                      INTEGER LB2, UB2, LB_EMBED2, UB_EMBED2
22
                ! The subgrid has been passed in its 'embedded' form
23
                      REAL X ( LB_EMBED1 : UB_EMBED1 , LB_EMBED2 : UB_EMBED2 )
^{24}
                ! Locally X_DESC is declared as an INTEGER
^{25}
                      INTEGER X_DESC
26
                ! Get the global extent of the first axis
27
                ! This is an HPF_LOCAL type of inquiry routine with an
28
                ! 'F77_' prefix
29
                      CALL F77_GLOBAL_SIZE(NX,X,1)
30
                ! Otherwise, initialize elements of the array
31
                ! Loop only over actual array elements
32
                      DO J = LB2, UB2
33
                        DO I = LB2, UB2
34
                          X(I,J) = I + (J-1) * NX
35
                        END DO
36
                      END DO
37
                      END
38
39
40
41
42
43
44
45
46
47
48
```