

Incremental caffeination of a terrestrial hydrological modeling framework using Fortran 2018 teams

Extended Abstract

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ABSTRACT

We present Fortran 2018 teams (grouped processes) running a parallel ensemble of simulations built from a pre-existing Message Passing Interface (MPI) application. A challenge arises around the Fortran standard's eschewing any direct reference to lower-level communication substrates, such as MPI, leaving any interoperability between Fortran's parallel programming model, Coarray Fortran (CAF), and the supporting substrate to the quality of the compiler implementation. Our approach introduces CAF incrementally, a process we term "caffeination." By letting CAF initiate execution and exposing the underlying MPI communicator to the original application code, we create a one-to-one correspondence between MPI group colors and Fortran teams. We apply our approach to the National Center for Atmospheric Research (NCAR)'s Weather Research and Forecasting Hydrological Model (WRF-Hydro). The newly caffeinated main program replaces batch job submission scripts and forms teams that each execute one ensemble member. To support this work, we developed the first compiler front-end and parallel runtime library support for teams. This paper describes the required modifications to a public GNU Compiler Collection (GCC) fork, an OpenCoarrays [1] application binary interface (ABI) branch, and a WRF-Hydro branch.

CCS CONCEPTS

• Software and its engineering → Parallel programming languages; • Applied computing → Environmental sciences;

KEYWORDS

coarray Fortran, computational hydrology, parallel programming

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1 INTRODUCTION

1.1 Motivation and Background

Since the publication of the Fortran 2008 standard in 2010 [4], Fortran supports a Single-Program Multiple-Data (SPMD) programming style that facilitates the creation of a fixed number of replicas of a compiled program, wherein each replica executes asynchronously after creation. Fortran refers to each replica as an image. The primary mechanism for distributing and communicating data between images involves defining coarrays, entities that may be referenced or defined on one image by statements executing on other images. As such, a coarray defines a partitioned global address space (PGAS) in which one image referencing or defining a coarray on another image causes inter-image communication.

The seminal role that coarrays played in the development of Fortran's intrinsic parallel programming model have made it common to refer to all of modern Fortran's parallel programming features under the rubric of CAF. To date, most published CAF applications involve scenarios wherein the parallelization itself poses one of the chief challenges and necessitates the custom development of parallel algorithms. These include ordinary and partial differential equation solvers in domains ranging from nuclear fusion [7] and weather [5] to multidimensional fast Fourier transforms and multigrid numerical methods [2]. Much of the effort involved in expressing parallel algorithms for these domains centers on designing and using various coarray data structures. In such settings, the moniker CAF seems appropriate.

Less widely appreciated are the ways Fortran's intrinsic parallel programming model supports embarrassingly parallel applications, wherein the division into independent sub-problems requires little coordination between the sub-problems. To support such applications, a parallel programming model might provide for explicit sub-problem disaggregation and independent sub-problem execution without any need for PGAS data structures such as coarrays. The draft Fortran 2018 standard (previously named "Fortran 2015"¹) offers several features that enable a considerable amount of parallel computation, coordination, and communication even without coarrays. A working definition of "embarrassingly parallel" Fortran might denote the class of use cases for which parallel algorithmic needs are met by the non-coarray parallel features, including

- Forming teams of images that communicate only with each other by default,
- Image synchronization: a mechanism for ordering the execution of program segments in differing images,

¹A Committee Draft is at <https://bit.ly/fortran-2015-draft>.

- Collective subroutines: highly optimized implementations of common parallel programming patterns,
- Image enumeration: functions for identifying and counting team members,
- Global error termination.

We anticipate that a common use case will encompass an ensemble of simulations, each member of which executes as a parallel computation in a separate team. This paper presents such a use case for WRF-Hydro, a terrestrial hydrological model developed at NCAR.

1.2 Objectives

The objectives of the current work are threefold:

- (1) To contribute the first-ever compiler front-end support for Fortran 2018 teams.
- (2) To contribute the parallel runtime library functionality required to support the new compiler capabilities.
- (3) To study and address issues arising from integrating teams into an existing MPI application, WRF-Hydro,

The compiler front-end described herein lives on the Sourcery Institute GCC fork's² teams branch. The parallel runtime library lives on the OpenCoarrays³ opencoarrays-teams ABI. We are unaware of any compiler support for Fortran 2018 teams other than that developed for the current project.

2 METHODOLOGY

Fortran 2018 will facilitate forming nonoverlapping image groups, allowing more efficient and independent subproblem execution. Multiphysics applications, e.g., climate and weather models, may benefit from the consequent reduction in off-node communication, particularly when an entire team fits in a single compute node.

2.1 Teams in Fortran 2018

A team is a set of images that can execute independently of other images outside the team. At program launch, all images comprise a team designated by a language-defined `initial_team` integer identifier. Except for the initial team, every team has a parent team in a one-to-many parent/child hierarchy. A program executes a `form team` statement to create a mapping for subsequent grouping of images into new child teams. Each new team has an integer identifier that Fortran produces as the result of invoking the `team_number()` intrinsic function. Information about the team to which the current image belongs can be determined by the compiler from the collective value of the team variables on the images of the team. All images execute the `form team` statement as a collective operation.

An image changes teams by executing `change team` or `end team`. The former moves the executing image from the current team to a team specified by a derived-type `team_type` variable. Subsequent execution of a corresponding `end team` statement restores the current team back to that team to which it belonged immediately prior to execution of the most recent `change team`.

The `form team` statement takes an `team_number` argument uniquely identifying the team and a `team_type` argument encapsulating other team information in private components. Successful

```

1 module assertions_module!Terminate globally if test fails
2 implicit none; contains
3 elemental subroutine assert(assertion, description)
4   logical, intent(in) :: assertion
5   character(len=*), intent(in), optional :: description
6   integer, parameter :: max_digits=12
7   character(len=max_digits) :: image_number
8   if (.not. assertion) then
9     write(image_number,*) this_image()
10    error stop description //" failed on " //image_number
11  end if
12 end subroutine
13 end module
14
15 program main !! Test team_number intrinsic function
16 use iso_fortran_env, only : team_type
17 use assertions_module, only : assertions
18 implicit none
19 integer, parameter :: standard_initial_value=-1
20 type(team_type), target :: home
21 call assert(team_number()==standard_initial_value)
22 associate(my_team=>mod(this_image(),2)+1)
23   form team(my_team,home)!Map even|odd images->teams 1|2
24   change team(home)
25   call assert(team_number()==my_team,"correct mapping")
26 end team
27 call assert(team_number()==standard_initial_value)
28 end associate
29 sync all; if (this_image()==1) print *, "Test passed."
30 end program

```

Figure 1: A unit test for the team-number function.

execution of a `form team` statement assigns the team-variable (of type `team_type`) on each participating image a value that specifies the new team to which the image will belong. The `change team` statement takes as argument a `team_type` variable that represents the new team to be used as current team. The execution of the `end team` statement restores the current team back to that immediately prior to execution of the `change team` statement. Figures 1-2 demonstrate the use of teams in two unit tests from the OpenCoarrays repository.

2.2 Teams in GCC and OpenCoarrays

A Fortran team is comparable to an MPI communicator. OpenCoarrays uses `MPI_Comm_split` to support `form team`, passing the team id as the color, and storing the resulting communicator in an available-teams list. Every list element tracks the team/communicator pairings. The function returns the available-teams list stored in the `team_type` variable.

We initialize the available- and used-teams lists to 1 (equivalent to `MPI_COMM_WORLD`) at the beginning of the execution. At a `change team`, the available-teams list element stored in the `team_type` variable gets passed to the corresponding OpenCoarrays function. The `current_team` variable used inside OpenCoarrays for representing the current communicator gets reassigned with the value contained into the element of the list passed as argument. Finally, a new element is added to the list of used teams. The list elements are pointers to the elements of the available-teams list. The insertion operation is always performed at the beginning of the list in order to keep track of the teams hierarchy. An execution of the `end team` statements is implemented by removing the first element of the list of used teams and reassigning the `current_team` to the new first element of the list of used teams.

²<https://github.com/sourceryinstitute/gcc>

³<https://github.com/sourceryinstitute/opencoarrays>

```

233 1 program main !! Test get_communicator language extension
234 2 use opencoarrays, only : get_communicator
235 3 use assertions_module, only : assert
236 4 use iso_fortran_env, only : team_type
237 5 type(team_type) :: league
238 6 integer, parameter :: num_teams=2 !! number of child teams to form
239 7 implicit none
240 8 call mpi_matches_caf(get_communicator()) !! verify rank & image numbering
241 9 associate(initial_image=>this_image(), initial_num_images=>num_images(), new_team=>mod(this_image()+1,num_teams)+1)
242 10 form team(new_team,league) !! create mapping
243 11 change team(league) !! join child team
244 12 call mpi_matches_caf(get_communicator()) !! verify new rank/image numbers
245 13 associate(my_team=>team_number())
246 14 call assert(my_team==new_team,"assigned team matches chosen team")
247 15 associate(new_num_images=>initial_num_images/num_teams+merge(1,0,my_team<=mod(initial_num_images,num_teams)))
248 16 call assert(num_images()==new_num_images,"block distribution of images")
249 17 end associate; end associate
250 18 end team
251 19 call assert([initial_image==this_image(),initial_num_images==num_images()],"correct rank/image remapping")
252 20 end associate
253 21 sync all; if (this_image()==1) print *,"Test passed."
254 22 contains
255 23 subroutine mpi_matches_caf(comm) !! verify num. ranks = num. images & image num. = rank num. + 1
256 24 use iso_c_binding, only : c_int
257 25 use mpi, only : MPI_COMM_SIZE, MPI_COMM_RANK
258 26 integer(c_int), intent(in) :: comm !! MPI communicator
259 27 integer(c_int) :: isize,ierror,irank
260 28 call MPI_COMM_SIZE(comm, isize, ierror)
261 29 call assert([ierror==0, isize==num_images()],"correct rank/image cardinality")
262 30 call MPI_COMM_RANK(comm, irank, ierror)
263 31 call assert([ierror==0, irank==this_image()-1,"correct rank/image numbering correspondence")
264 32 end subroutine
265 33 end program

```

Figure 2: A unit test for the get_communicator function.

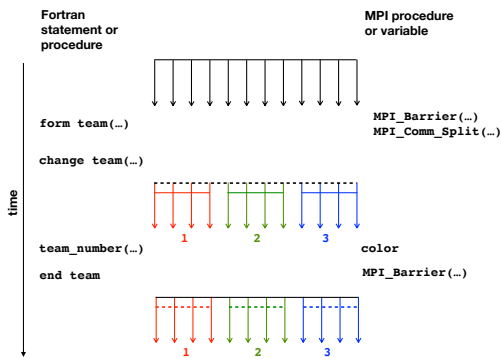


Figure 3: Schematic of program execution over time (left axis) in 12 images (top) communicating globally and within subgroups. Horizontal lines show communication mechanisms (default=solid, optional=dashed). Fortran concepts (left). Underlying MPI concepts (right).

Figure 3 depicts schematically an initial team of images (black arrows) executing over time (progressing downward) and able to coordinate and communicate through a global mechanism (black horizontal line). At the point of executing `form team` and `change team` statements, the compiler inserts references to the OpenCoarrays ABI into the executable program. Those references cause invocations of `MPI_Split`, which in turn creates the colored groupings that correspond to teams in Fortran 2018.

The teams unit tests in Figures 1–2 use a block distribution of images, dividing the initial team into three new teams, each with

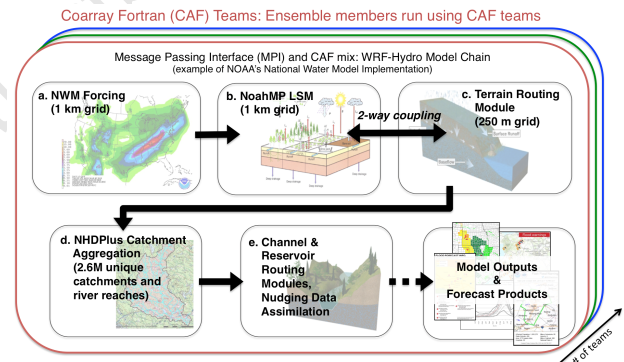


Figure 4: WRF-Hydro caffeination via Fortran 2018 teams: example components of the National Water Model. Different MPI colors represent independent teams, each of which is an ensemble member.

the same number of images except some teams with one extra. The number of teams with an extra image equals the remainder of integer division of the total number images by the number of teams. In Figure 2, an assertion procedure terminates across all images if assertion is false. The optional second argument in `assert` describes the checks performed.

2.3 A language extension

Line 2 in Figure 2 imports a `get_communicator()` function via Fortran's use-association mechanism for accessing entities in Fortran modules: an `opencoarrays` module that provides language

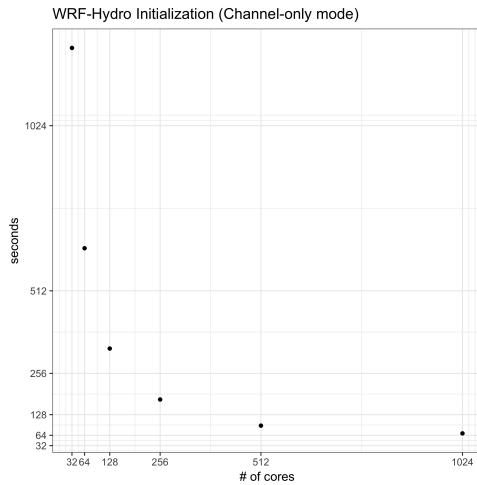


Figure 5: Scaling of the channel-only initialization time to support Amdahl's law calculation of potential speedup of ensemble computation with shared initialization.

extensions. Lines 8 and 12 invoke this function to provide the MPI communicator to the test subroutine `mpi_mpatches_caf`. Assertions in the latter subroutine verify the expected correspondences between MPI image and rank numbering. The MPI/CAF correspondence enables the newly caffeinated WRF-Hydro to interoperate safely with the existing WRF-Hydro MPI code.

3 DISCUSSION OF RESULTS

WRF-Hydro is a community hydrologic model providing a parallel-computing framework for coupling weather prediction (Figure 4a), land surface (Figure 4b), and hydrologic routing to handle spatial water redistribution via overland flow (Figure 4c), subsurface (soil column) flow (Figure 4c), baseflow (deep groundwater, 4d), and stream channel transport (Figure 4e) [3].

Developed to couple land hydrology to the atmosphere, WRF-Hydro usually runs “offline” with forcing from upper-boundary (weather) conditions (Fig. 4). The chief example is the National Weather Model (NWM) of the National Oceanic and Atmospheric Administration (NOAA) [6], a special configuration of WRF-Hydro providing operational, real-time analysis and forecasts over the U.S.

Ensemble forecasting and ensemble data assimilation are growing research areas for WRF-Hydro. Running ensembles under one executable program and job submission as shown in Fig. 4 can reduce the labor in workflow design and can open up possibilities for improving data flows and optimizing ensemble execution time.

An optimization that we expect teams to facilitate will involve sharing initialization work over all ensemble members using the full image set before changing teams. Amortizing common initialization work across all available computational resources rather than repeating common initialization work in each ensemble member using only that member's fraction of the resources might save significant computational costs in runs for which initialization occupies a large fraction of the run time such as in production runs of short-term NWM forecasts. We are exploring moving from deterministic (single ensemble member) runs to ensemble runs for

the stream channel submodel in isolation. Fig. 5 presents the scaling behavior of channel-only runs. Running 50 ensemble members concurrently in teams occupying 20 cores each, the initialization speedup will be approximately 20x. Currently, for the short-range forecasts, initialization requires approximately 50% of the run time on NCAR's yellowstone computer and greater than that on NOAA WCOSS system [8]. These numbers are similar for the channel-only model. In our proposed application for an ensemble size of 50, we estimate that approximately half the runtime ($f = .5$) can be sped up by a factor of about 20 ($S_f = 20$), based on Fig. 5. From these estimates, Amdahl's law yields a total speedup of 1.9:

$$S = 1/f/S_f + (1 - f) = 1/(.5/20 + (1 - .5)) = 1.9 \quad (1)$$

We converted the WRF-Hydro main program to a subroutine called in a loop over the ensemble members. We deleted `MPI_Init` and `MPI_Finalize` calls in the converted subroutine and instead pass a communicator to the subroutine. To loop over the ensemble also required deleting the stop statement in the subroutine.

4 CONCLUSIONS AND FUTURE WORK

We applied Fortran 2018 teams to WRF-Hydro ensemble simulation and forecasting. We developed the first-ever compiler front-end and parallel runtime library support for teams and a language extension that exposes CAF's underlying MPI communicator for use in WRF-Hydro. This approach facilitates incremental introduction of CAF, i.e., “caffeination,” of the pre-existing MPI program. We predict a speedup of 1.9 in ensemble execution when spreading common initializations across all images prior to changing teams. Such a speedup would greatly impact WRF-Hydro operational use cases.

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