

# WRF Software: Code and Parallel Computing

John Michalakes, WRF Software Architect

Dave Gill

## Outline

- WRF architecture – driver, mediation, model
- Need and design for parallelism
- Communication patterns to support parallelism
- Directory structure and file location overview
- Model layer interface
  - The “grid” struct
  - Indices
  - Dereferencing
- I/O

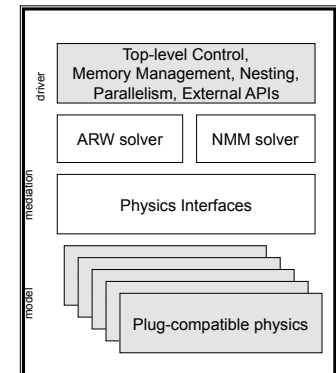
## Introduction – WRF Software Characteristics

- Developed from scratch beginning around 1998, primarily Fortran and C
- Requirements emphasize flexibility over a range of platforms, applications, users, performance
- WRF develops rapidly. First released Dec 2000
- Supported by flexible efficient architecture and implementation called the WRF Software Framework

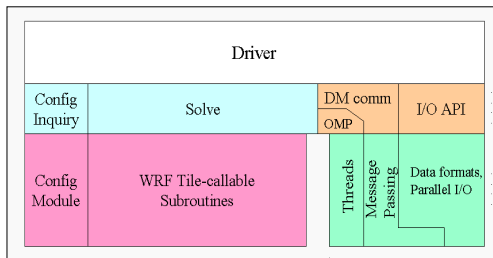
## Introduction - WRF Software Framework Overview

- Implementation of WRF Architecture
  - Hierarchical organization
  - Multiple dynamical cores
  - Plug compatible physics
  - Abstract interfaces (APIs) to external packages
  - Performance-portable
- Designed from beginning to be adaptable to today's computing environment for NWP

<http://mmm.ucar.edu/wrf/WG2/bench/>

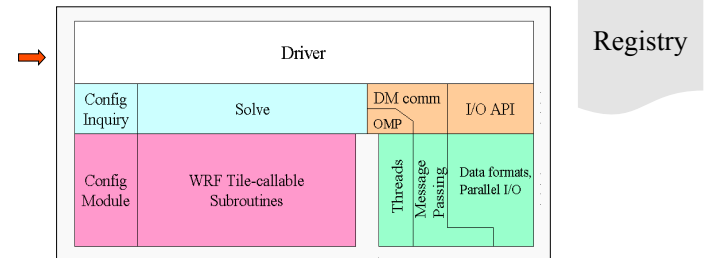


## WRF Software Architecture



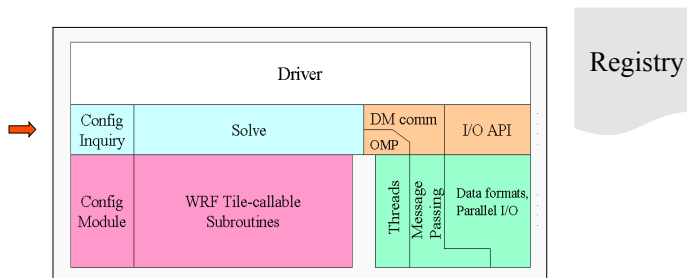
- **Hierarchical** software architecture
  - **Insulate** scientists' code from parallelism and other architecture/implementation-specific details
  - Well-defined **interfaces** between layers, and **external packages** for communications, I/O, and model coupling facilitates code reuse and exploiting of community infrastructure, e.g. ESMF.

## WRF Software Architecture



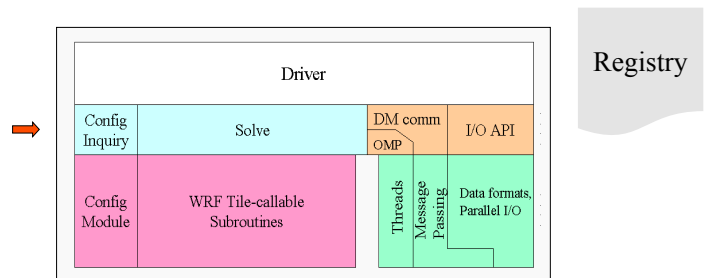
- **Driver** Layer
  - **Domains**: Allocates, stores, decomposes, represents abstractly as **single data objects**
  - **Time loop**: top level, algorithms for **integration over nest hierarchy**

## WRF Software Architecture



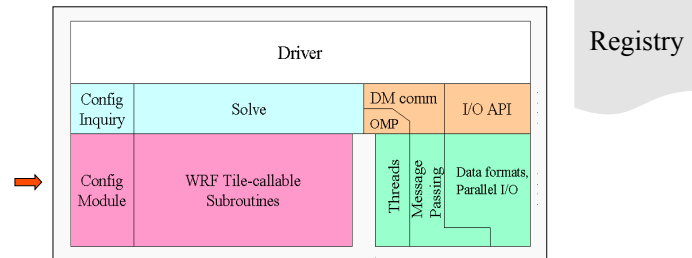
- **Mediation** Layer
  - **Solve** routine, takes a **domain object** and advances it **one time step**
  - **Nest** forcing, interpolation, and feedback routines

## WRF Software Architecture



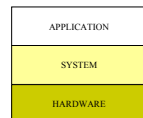
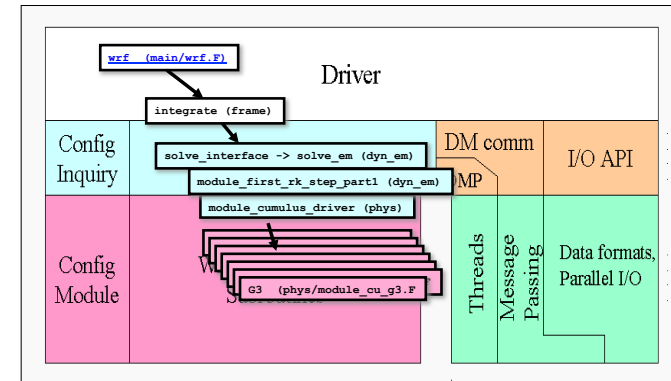
- **Mediation** Layer
  - The **sequence of calls** for doing a time-step for one domain is known in Solve routine
  - **Dereferences fields** in calls to physics drivers and dynamics code
  - Calls to **message-passing** are contained here as part of Solve routine

## WRF Software Architecture



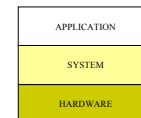
- Model Layer
  - **Physics and Dynamics:** contains the actual WRF model routines are written to **perform some computation** over an arbitrarily sized/ shaped, 3d, rectangular subdomain

## Call Structure Superimposed on Architecture



## Hardware: The Computer

- The 'N' in NWP
- Components
  - Processor
    - A program counter
    - Arithmetic unit(s)
    - Some scratch space (registers)
    - Circuitry to store/retrieve from memory device
    - Cache
  - Memory
  - Secondary storage
  - Peripherals
- The implementation has been continually refined, but the basic idea hasn't changed much



## Hardware has not changed much...

**IBM 7090**

A computer in 1960

6-way superscalar  
36-bit floating point precision  
~144 Kbytes  
~50,000 flop/s  
48hr 12km WRF CONUS in 600 years

A computer in 2013

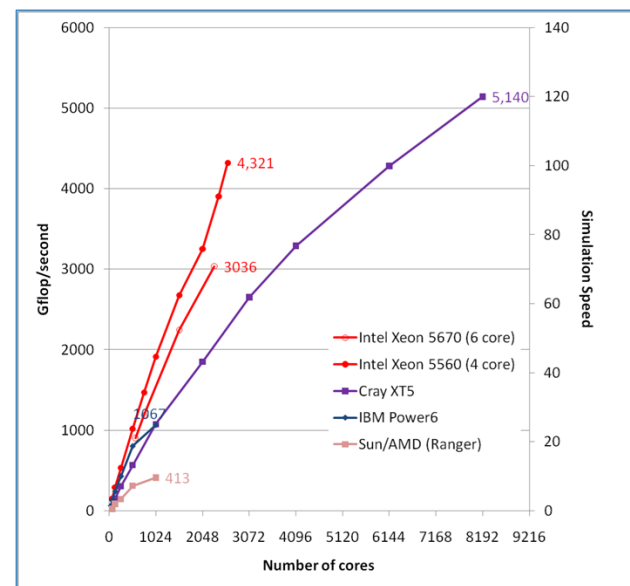
Dual core, 2.6 GHz chip  
64-bit floating point precision  
20 MB L3  
~5,000,000,000 flop/s  
48 12km WRF CONUS in 26 Hours

APPLICATION
SYSTEM
HARDWARE

...how we use it has

- Fundamentally, processors haven't changed much since 1960
- Quantitatively, they haven't improved nearly enough
  - 100,000x increase in peak speed
  - 100,000x increase in memory size
- We make up the difference with parallelism
  - Ganging multiple processors together to achieve  $10^{11-12}$  flop/second
  - Aggregate available memories of  $10^{11-12}$  bytes

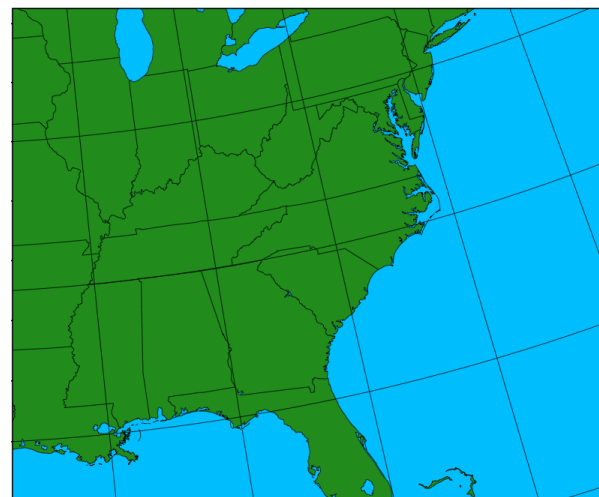
*~1,000,000,000,000 flop/s ~2500 procs  
48-h, 12-km WRF CONUS in under 15 minutes*



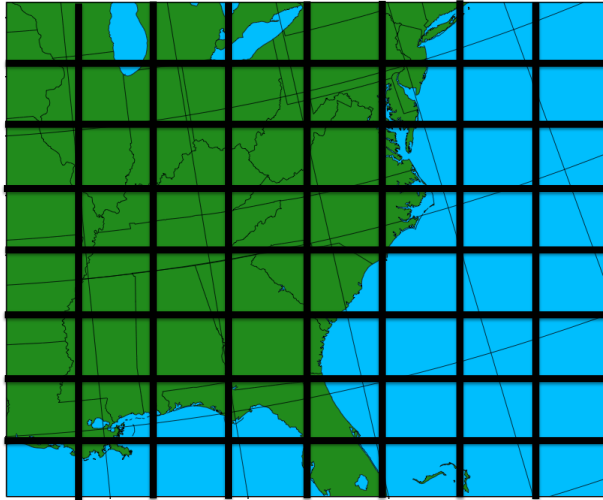
## WRF Domain Decomposition

- The WRF model decomposes domains horizontally
- For  $n$  MPI tasks, the two nearest factors ( $n = k * m$ ) are selected; the larger is used to decompose the y-direction, the smaller is used to decompose the x-direction
- Users may choose a preferred decomposition (nproc\_x, nproc\_y)
- Prime numbers and composites with large prime factors are usually to be avoided
- The behavior of 132 vs 131, and 200 vs 202 are quite different

## January 2000 Benchmark – 1 task: 74x61



### January 2000 Benchmark – 64 tasks: 10x8



### WRF Domain Decomposition

- As you increase the number of total MPI tasks, you reduce the amount of work inside of each MPI task
- The amount of time to process communication between MPI tasks tends to be *at best* constant
- As more MPI tasks are involved, more contention for hardware resources due to communication is likely
- As the computation time gets smaller compared to the communications time, parallel efficiency suffers

### January 2000 Benchmark

- 74x61 grid cells
- 1 hour forecast, 3 minute time step, 20 time step average
- IO excluded

Decomposed domain sizes    proc count: I-dim x J-dim

1: 74x61      2: 74x31      4: 37x31      8: 37x16

16: 19x16    32: 19x8      64: 10x8

### January 2000 Benchmark

Processor Count	SM – OpenMP % Efficiency	DM – MPI % Efficiency
1 74x61	100	100
2 74x31	72	98
4 37x31	65	91
8 37x16	31	83
16 19x16	16	70
32 19x8	8	56
64 10x8	3	40

## January 2000 Benchmark

- WRF timing estimates may be obtained from the model print-out

### Serial

Timing for main on domain 1: 32.16074 elapsed seconds

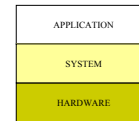
### OpenMP

Timing for main on domain 1: 8.56216 elapsed seconds

### MPI

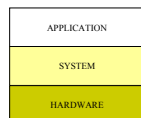
Timing for main on domain 1: 7.36243 elapsed seconds

- Get enough time steps to include “day-time” radiation, and to have the microphysics “active” for better estimates



## Application: WRF

- WRF can be run **serially** or as a **parallel** job
- WRF uses **domain decomposition** to divide total amount of work over parallel processes



## Parallelism in WRF: Multi-level Decomposition

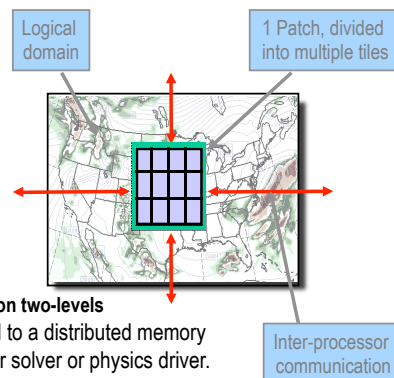
- Single version of code for efficient execution on:
  - Distributed-memory
  - Shared-memory (SMP)
  - Clusters of SMPs
  - Vector and microprocessors

### Model domains are decomposed for parallelism on two-levels

**Patch:** section of model domain allocated to a distributed memory node, this is the scope of a mediation layer solver or physics driver.

**Tile:** section of a patch allocated to a shared-memory processor within a node; this is also the scope of a model layer subroutine.

Distributed memory parallelism is over patches; shared memory parallelism is over tiles within patches



## Distributed Memory Communications

### When Needed?

Communication is required between patches when a horizontal index is incremented or decremented on the right-hand-side of an assignment.

### Why?

On a patch boundary, the index may refer to a value that is on a different patch.

Following is an example code fragment that requires communication between patches

### Signs in code

Note the tell-tale **+1** and **-1** expressions in indices for **rr**, **H1**, and **H2** arrays on right-hand side of assignment.

These are **horizontal data dependencies** because the indexed operands may lie in the patch of a neighboring processor. That neighbor's updates to that element of the array won't be seen on this processor.

## Distributed Memory Communications

```
(module_diffusion.F)

SUBROUTINE horizontal_diffusion_s (tendency, rr, var, . . .
. . .
DO j = jts,jte
DO k = kts,ktf
DO i = its,ite
  mrdx=msft(i,j)*rdx
  mrdy=msft(i,j)*rdy
  tendency(i,k,j)=tendency(i,k,j)-
    (mrdx*0.5*((rr(i+1,k,j)+rr(i,k,j))*H1(i+1,k,j)-
    (rr(i-1,k,j)+rr(i,k,j))*H1(i,k,j))+
    mrdy*0.5*((rr(i,k,j+1)+rr(i,k,j))*H2(i,k,j+1)-
    (rr(i,k,j-1)+rr(i,k,j))*H2(i,k,j))-
    msft(i,j)*(H1avg(i,k+1,j)-H1avg(i,k,j)+
    H2avg(i,k+1,j)-H2avg(i,k,j)
    )/dzetaw(k)
    )
ENDDO
ENDDO
ENDDO
. . .
```

## Distributed Memory Communications

```
(module_diffusion.F)

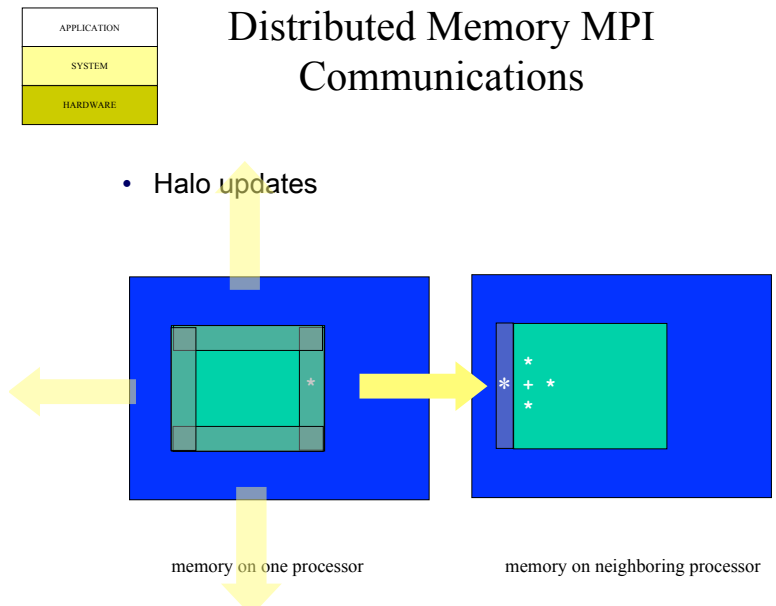
SUBROUTINE horizontal_diffusion_s (tendency, rr, var, . . .
. . .
DO j = jts,jte
DO k = kts,ktf
DO i = its,ite
  mrdx=msft(i,j)*rdx
  mrdy=msft(i,j)*rdy
  tendency(i,k,j)=tendency(i,k,j)-
    (mrdx*0.5*((rr(i+1,k,j)+rr(i,k,j))*H1(i+1,k,j)-
    (rr(i-1,k,j)+rr(i,k,j))*H1(i,k,j))+
    mrdy*0.5*((rr(i,k,j+1)+rr(i,k,j))*H2(i,k,j+1)-
    (rr(i,k,j-1)+rr(i,k,j))*H2(i,k,j))-
    msft(i,j)*(H1avg(i,k+1,j)-H1avg(i,k,j)+
    H2avg(i,k+1,j)-H2avg(i,k,j)
    )/dzetaw(k)
    )
ENDDO
ENDDO
ENDDO
. . .
```

## Distributed Memory Communications

```
(module_diffusion.F)

SUBROUTINE horizontal_diffusion_s (tendency, rr, var, . . .
. . .
DO j = jts,jte
DO k = kts,ktf
DO i = its,ite
  mrdx=msft(i,j)*rdx
  mrdy=msft(i,j)*rdy
  tendency(i,k,j)=tendency(i,k,j)-
    (mrdx*0.5*((rr(i+1,k,j)+rr(i,k,j))*H1(i+1,k,j)-
    (rr(i-1,k,j)+rr(i,k,j))*H1(i,k,j))+
    mrdy*0.5*((rr(i,k,j+1)+rr(i,k,j))*H2(i,k,j+1)-
    (rr(i,k,j-1)+rr(i,k,j))*H2(i,k,j))-
    msft(i,j)*(H1avg(i,k+1,j)-H1avg(i,k,j)+
    H2avg(i,k+1,j)-H2avg(i,k,j)
    )/dzetaw(k)
    )
ENDDO
ENDDO
ENDDO
. . .
```

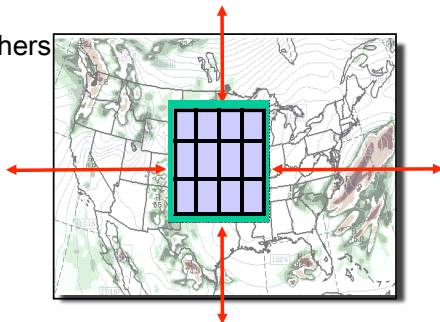
## Distributed Memory MPI Communications



APPLICATION
SYSTEM
HARDWARE

## Distributed Memory (MPI) Communications

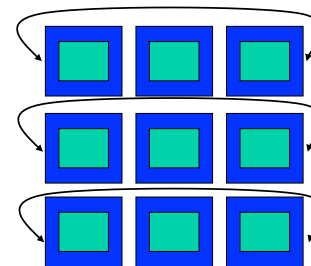
- Halo updates
- Periodic boundary updates
- Parallel transposes
- Nesting scatters/gathers



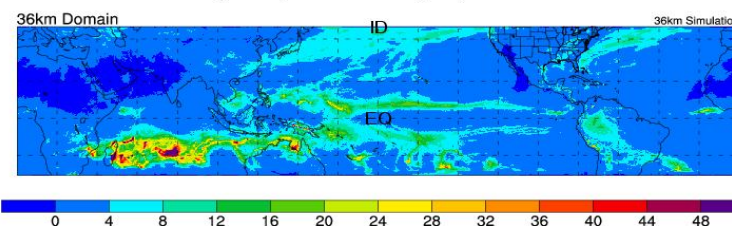
APPLICATION
SYSTEM
HARDWARE

## Distributed Memory (MPI) Communications

- Halo updates
- Periodic boundary updates
- Parallel transposes
- Nesting scatters/gathers



Average Daily Total rainfall (mm) - March 1997



APPLICATION
SYSTEM
HARDWARE

## Distributed Memory (MPI) Communications

- Halo updates
- Periodic boundary updates
- Parallel transposes
- Nesting scatters/gathers



all y on  
patch



all z on  
patch

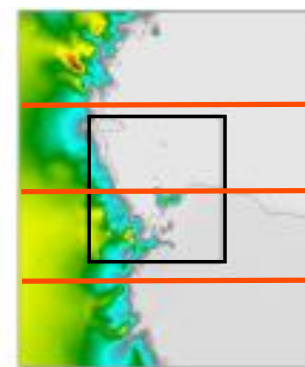


all x on  
patch

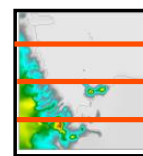
APPLICATION
SYSTEM
HARDWARE

## Distributed Memory (MPI) Communications

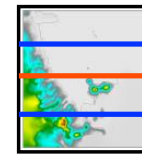
- Halo updates
- Periodic boundary updates
- Parallel transposes
- Nesting scatters/gathers



COARSE  
Ross Island  
6.66 km



NEST: 2.22 km



INTERMEDIATE: 6.66 km



## WRF Model Top-Level Directory Structure

[WRF Design  
and  
Implementation](#)  
Doc, p 5

DRIVER ●  
MEDIATION ●  
MODEL ●

Makefile		
README		
README_test_cases		
clean	}	build scripts
compile		
configure		
Registry/		CASE input files machine build rules
arch/		
● dyn_em/	}	source code directories
● dyn_nnm/		
external/		
● frame/		
inc/		
● main/		
● phys/		
● share/	}	execution directories
tools/		
run/		
test/		

Where are WRF source code files located?

```
$(RM) $@
```

```
$(CPP) -I$(WRF_SRC_ROOT_DIR)/inc \  
$(CPPFLAGS) $(OMPCPP) $*.F > $*.f90
```

```
$(FC) -o $@ -c $(FCFLAGS) $(MODULE_DIRS) \  
$(PROMOTION) $(FCSUFFIX) $*.f90
```

Where are WRF source code files located?

```
cpp -C -P file.F > file.f90
```

```
gfortran -c file.f90
```

Where are WRF source code files located?

- The most important command is the “find” command. If there is an error in the model output, you can find that location in the source code with the **find** command.

```
cd WRFV3
```

```
find . -name \*.F -exec grep -i “Flerchinger” {} \; -print
```

### Where are WRF source code files located?

- All of the differences between the .F and .f90 files are due to the included pieces that are manufactured by the Registry.
- These additional pieces are all located in the WRFV3/inc directory.
- For a serial build, almost 450 files are manufactured.
- Usually, most developers spend their time working with physics schemes.

### Where are WRF source code files located?

- The “main” routine that handles the calls to all of the physics and dynamics:
  - WRFV3/dyn\_em/solve\_em.F
- This “solver” is where the tendencies are initialized to zero, some pre-physics terms are computed, and the time stepping occurs
- The calls to most of the physics schemes are made from a further call down the call tree
  - dyn\_em/module\_first\_rk\_step\_part1.F

### Where are WRF source code files located?

- Inside of solve\_em and first\_rk\_step\_part1, all of the data is located in the “grid” structure: grid%ht.
- The dimensions in solve\_em and first\_rk\_step\_part1 are “d” (domain), and “m” (memory):
  - ids, ide, jds, jde, kds, kde
  - ims, ime, jms, jme, kms, kme
- The “t” (tile) dimensions are computed in first\_rk\_step\_part1 and passed to all drivers.
- WRF uses global indexing

### Where are WRF source code files located?

- If you are interested in looking at physics, the WRF system has organized the files in the WRFV3/phys directory.
- In WRFV3/phys, each type of physics has a driver:

module_cumulus_driver.F	cu
module_microphysics_driver.F	mp
module_pbl_driver.F	bl
module_radiation_driver.F	ra
module_surface_driver.F	sf

## Where are WRF source code files located?

- The subgrid-scale precipitation (\*\_cu\_\*.F)
 

module_cu_bmj.F	module_cu_camzm.F
module_cu_g3.F	module_cu_gd.F
module_cu_kf.F	module_cu_kfeta.F
module_cu_nsas.F	module_cu_osas.F
module_cu_sas.F	module_cu_tiedtke.F

## Where are WRF source code files located?

- Advection
 

WRFV3/dyn\_em/module\_advect\_em.F
- Lateral boundary conditions
 

WRFV3/dyn\_em/module\_bc\_em.F

## Where are WRF source code files located?

- Compute various RHS terms, pressure gradient, buoyancy, w damping, horizontal and vertical diffusion, Coriolis, curvature, Rayleigh damping
 

WRFV3/dyn\_em/module\_big\_step\_utilities\_em.F
- All of the sound step utilities to advance u, v, mu, t, w within the small time-step loop
 

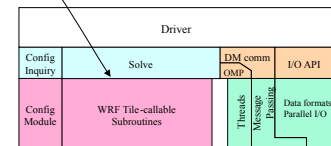
WRFV3/dyn\_em/module\_small\_step\_em.F

## WRF Model Layer Interface – The Contract with Users

All state **arrays** passed through argument list as simple (not derived) data types

Domain, memory, and run dimensions passed unambiguously in **three dimensions**

Model layer routines are called from mediation layer (physics drivers) in **loops over tiles**, which are multi-threaded



## WRF Model Layer Interface – The Contract with Users

### Restrictions on Model Layer subroutines:

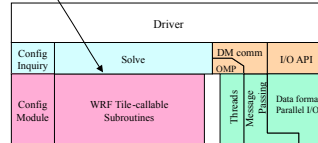
No I/O, communication

No stops or aborts

Use wrf\_error\_fatal

No common/module storage of decomposed data

Spatial scope of a Model Layer call is one “tile”



## WRF Model Layer Interface

```
SUBROUTINE driver_for_some_physics_suite (
    . . .
!$OMP DO PARALLEL
    DO ij = 1, numtiles
        its = i_start(ij) ; ite = i_end(ij)
        jts = j_start(ij) ; jte = j_end(ij)
        CALL model_subroutine( arg1, arg2, . . .
            ids , ide , jds , jde , kds , kde ,
            ims , ime , jms , jme , kms , kme ,
            its , ite , jts , jte , kts , kte )
    END DO
    . . .
END SUBROUTINE
```

## WRF Model Layer Interface

### template for model layer subroutine

```
SUBROUTINE model_subroutine ( &
    arg1, arg2, arg3, ... , argn, &
    ids, ide, jds, jde, kds, kde, & ! Domain dims
    ims, ime, jms, jme, kms, kme, & ! Memory dims
    its, ite, jts, jte, kts, kte ) ! Tile dims

IMPLICIT NONE

! Define Arguments (State and I1) data
REAL, DIMENSION (ims:ime,kms:kme,jms:jme) :: arg1, . . .
REAL, DIMENSION (ims:ime,jms:jme) :: arg7, . . .
. . .
! Define Local Data (I2)
REAL, DIMENSION (its:ite,kts:kte,jts:jte) :: loc1, . . .
. . .
```

## WRF Model Layer Interface

### template for model layer subroutine

```
. . .
! Executable code; loops run over tile
! dimensions
DO j = jts, MIN(jte,jde-1)
    DO k = kts, kte
        DO i = its, MIN(ite,ide-1)
            loc1(i,k,j) = arg1(i,k,j) + ...
        END DO
    END DO
END DO
```

```

template for model layer subroutine
SUBROUTINE model ( &
  arg1, arg2, arg3, ..., argn, &
  ids, ide, jds, jde, kds, kde, & ! Domain dims
  ims, ime, jms, jme, kms, kme, & ! Memory dims
  its, ite, jts, jte, kts, kte ) ! Tile dims

IMPLICIT NONE

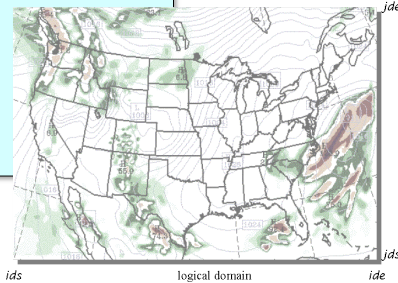
! Define Arguments (S and I1) data
REAL, DIMENSION (ims:ime,kms:kme,jms:jme) :: arg1, ...
REAL, DIMENSION (ims:ime,jms:jme) :: arg7, ...

! Define Local Data (I2)
REAL, DIMENSION (its:ite,kts:kte,jts:jte) :: loc1, ...

! Executable code; loops run over tile
! dimensions
DO j = MAX(jts,jds), MIN(jte,jde-1)
  DO k = kts, kte
    DO i = MAX(its,ids), MIN(ite,ide-1)
      loc1(i,k,j) = arg1(i,k,j) + ...
    END DO
  END DO
END DO

```

- Domain dimensions
  - Size of logical domain
  - Used for bdy tests, etc.



```

template for model layer subroutine
SUBROUTINE model ( &
  arg1, arg2, arg3, ..., argn, &
  ids, ide, jds, jde, kds, kde, & ! Domain dims
  ims, ime, jms, jme, kms, kme, & ! Memory dims
  its, ite, jts, jte, kts, kte ) ! Tile dims

IMPLICIT NONE

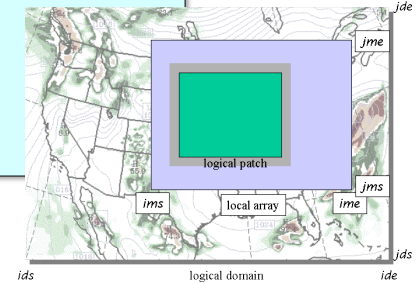
! Define Arguments (S and I1) data
REAL, DIMENSION (ims:ime,kms:kme,jms:jme) :: arg1, ...
REAL, DIMENSION (ims:ime,jms:jme) :: arg7, ...

! Define Local Data (I2)
REAL, DIMENSION (its:ite,kts:kte,jts:jte) :: loc1, ...

! Executable code; loops run over tile
! dimensions
DO j = MAX(jts,jds), MIN(jte,jde-1)
  DO k = kts, kte
    DO i = MAX(its,ids), MIN(ite,ide-1)
      loc1(i,k,j) = arg1(i,k,j) + ...
    END DO
  END DO
END DO

```

- Domain dimensions
  - Size of logical domain
  - Used for bdy tests, etc.
- Memory dimensions
  - Used to dimension dummy arguments
  - Do not use for local arrays



```

template for model layer subroutine
SUBROUTINE model ( &
  arg1, arg2, arg3, ..., argn, &
  ids, ide, jds, jde, kds, kde, & ! Domain dims
  ims, ime, jms, jme, kms, kme, & ! Memory dims
  its, ite, jts, jte, kts, kte ) ! Tile dims

IMPLICIT NONE

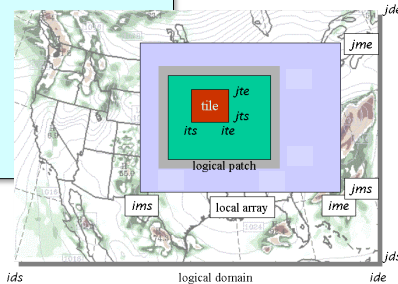
! Define Arguments (S and I1) data
REAL, DIMENSION (ims:ime,kms:kme,jms:jme) :: arg1, ...
REAL, DIMENSION (ims:ime,jms:jme) :: arg7, ...

! Define Local Data (I2)
REAL, DIMENSION (its:ite,kts:kte,jts:jte) :: loc1, ...

! Executable code; loops run over tile
! dimensions
DO j = MAX(jts,jds), MIN(jte,jde-1)
  DO k = kts, kte
    DO i = MAX(its,ids), MIN(ite,ide-1)
      loc1(i,k,j) = arg1(i,k,j) + ...
    END DO
  END DO
END DO

```

- Domain dimensions
  - Size of logical domain
  - Used for bdy tests, etc.
- Memory dimensions
  - Used to dimension dummy arguments
  - Do not use for local arrays
- Tile dimensions
  - Local loop ranges
  - Local array dimensions



```

template for model layer subroutine
SUBROUTINE model ( &
  arg1, arg2, arg3, ..., argn, &
  ids, ide, jds, jde, kds, kde, & ! Domain dims
  ims, ime, jms, jme, kms, kme, & ! Memory dims
  its, ite, jts, jte, kts, kte ) ! Tile dims

IMPLICIT NONE

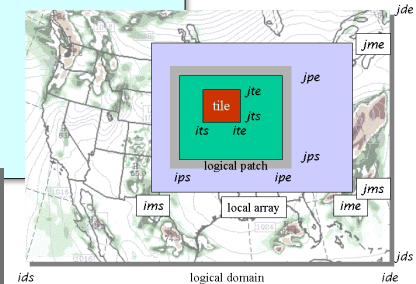
! Define Arguments (S and I1) data
REAL, DIMENSION (ims:ime,kms:kme,jms:jme) :: arg1, ...
REAL, DIMENSION (ims:ime,jms:jme) :: arg7, ...

! Define Local Data (I2)
REAL, DIMENSION (its:ite,kts:kte,jts:jte) :: loc1, ...

! Executable code; loops run over tile
! dimensions
DO j = MAX(jts,jds), MIN(jte,jde-1)
  DO k = kts, kte
    DO i = MAX(its,ids), MIN(ite,ide-1)
      loc1(i,k,j) = arg1(i,k,j) + ...
    END DO
  END DO
END DO

```

- Domain dimensions
  - Size of logical domain
  - Used for bdy tests, etc.
- Memory dimensions
  - Used to dimension dummy arguments
  - Do not use for local arrays
- Tile dimensions
  - Local loop ranges
  - Local array dimensions



- Patch dimensions
  - Start and end indices of local distributed memory subdomain
  - Available from mediation layer (solve) and driver layer; not usually needed or used at model layer

## WRF I/O

- Streams (similar to Fortran units): pathways into and out of model
- Can be thought of as files, though that is a restriction
  - History + auxiliary output streams (10 and 11 are reserved for nudging)
  - Input + auxiliary input streams (10 and 11 are reserved for nudging)
  - Restart, boundary, and a special DA in-out stream
  - Currently, 24 total streams
  - Use the large values and work down to stay away from “used”
  - Non-chemistry: use history streams 13-22, 24
  - Chemistry: use history streams 20, 21, 22, 24

## WRF I/O

- Attributes of streams
  - Variable set
    - The set of WRF state variables that comprise one read or write on a stream
    - Defined for a stream at compile time in Registry
  - Format
    - The format of the data outside the program (e.g. NetCDF), split
    - Specified for a stream at run time in the namelist

## WRF I/O

- Attributes of streams
  - Additional namelist-controlled attributes of streams
    - Dataset name
    - Time interval between I/O operations on stream
    - Starting, ending times for I/O (**specified as intervals from start of run**)

## WRF I/O

- Attributes of streams
  - **Mandatory for stream to be used:**
    - Time interval between I/O operations on stream
    - Format: `io_form`

### Example 1: Add output without recompiling

- Edit the namelist.input file, the time\_control namelist record

```
iofields_filename = "myoutfields.txt" (MAXDOM)
io_form_auxhist24 = 2 (choose an available stream)
auxhist24_interval = 10 (MAXDOM, every 10 minutes)
```
- Place the fields that you want in the named text file `myoutfields.txt`

```
+:h:24:RAIN,RAINNC
```
- Where “+” means ADD this variable to the output stream, “h” is the history stream, and “24” is the stream number

## Outline

- WRF architecture — driver, mediation, model
- Need and design for parallelism
- Communication patterns to support parallelism
- Directory structure and file location overview
- Model layer interface
  - The “grid” struct
  - Indices
  - Dereferencing
- I/O