

How to Use the WRF Registry

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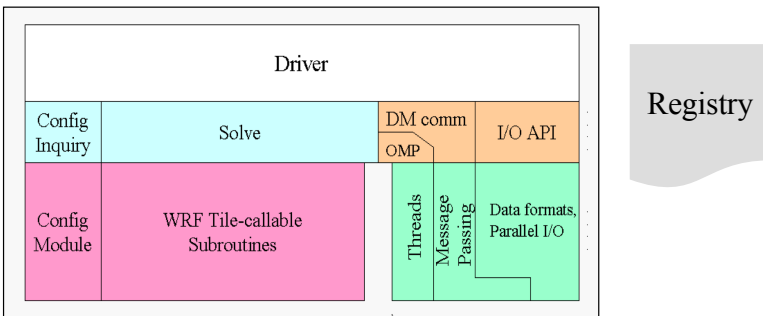
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Outline

- What is the WRF Registry
- Keyword syntax
- The BIG Three
- Examples
 - Runtime I/O mods
 - Adding a variable to the namelist
 - Adding an array to WRF
 - Compute a diagnostic
 - New physics scheme
 - Passive tracer

WRF Software Architecture



Text based file for real and WRF
Active data dictionary
Used with cpp to auto generate source
Controls/defines
Variables (I/O, comms, nesting)
Communications
namelist options

About 300k lines added to source
Easy – 3x the size since initial release
Compile-time option
./clean
./configure
./compile
Registry.EM_COMMON (else lost changes)

Registry Keywords

- Currently implemented as a text file: [Registry/Registry.EM_COMMON](#)
- Types of entry:
 - **Dimspec** – Describes dimensions that are used to define arrays in the model
 - **State** – Describes state variables and arrays in the domain structure
 - **I1** – Describes local variables and arrays in solve
 - **Typedef** – Describes derived types that are subtypes of the domain structure

Registry Keywords

- Types of entry:
 - Rconfig*—Describes a configuration (e.g. namelist) variable or array
 - Package*—Describes attributes of a package (e.g. physics)
 - Halo*—Describes halo update interprocessor communications
 - Period*—Describes communications for periodic boundary updates
 - Xpose*—Describes communications for parallel matrix transposes
 - include*—Similar to a CPP #include file

Registry State Entry

#	Type	Sym	Dims	Use	Tlev	Stag	IO	Dname	Descrip
state	real	tsk	ij	misc	1	-	i01rhud	"TSK"	"SKIN TEMP"

- Elements
 - Entry*: The keyword “state”
 - Type*: The type of the state variable or array (real, double, integer, logical, character, or derived)
 - Sym*: The symbolic name of the variable or array
 - Dims*: A string denoting the dimensionality of the array or a hyphen (-)
 - Use*: A string denoting association with a solver or 4D scalar array, or a hyphen
 - NumTlev*: An integer indicating the number of time levels (for arrays) or hyphen (for variables)

Registry State Entry

#	Type	Sym	Dims	Use	Tlev	Stag	IO	Dname	Descrip
state	real	tsk	ij	misc	1	-	i01rhud	"TSK"	"SKIN TEMP"

- Elements
 - Stagger*: String indicating staggered dimensions of variable (X, Y, Z, or hyphen)
 - IO*: String indicating whether and how the variable is subject to various I/O and Nesting
 - DName*: Metadata name for the variable
 - Units*: Metadata units of the variable
 - Descrip*: Metadata description of the variable

State Entry: Defining a variable-set for an I/O stream

- Fields are added to a variable-set on an I/O stream in the Registry

#	Type	Sym	Dims	Use	Tlev	Stag	IO	Dname	Descrip
state	real	tsk	ij	misc	1	-	i01rhud	"TSK"	"SKIN TEMP"

- IO* is a string that specifies if the variable is to be available to initial, restart, or history I/O. The string may consist of 'h' (subject to history I/O), 'i' (initial dataset), 'r' (restart dataset).
- The 'h', 'r', and 'i' specifiers may appear in any order or combination.

State Entry: Defining a variable-set for an I/O stream

- Fields are added to a variable-set on an I/O stream in the Registry

#	Type	Sym	Dims	Use	Tlev	Stag	IO	Dname	Descrip
state	real	tsk	ij	misc	1	-	i01rhud	"TSK"	"SKIN TEMP"

- The ‘h’ and ‘i’ specifiers may be followed by an optional integer string consisting of ‘0’, ‘1’, …, ‘9’
- Zero denotes that the variable is part of the principal input or history I/O stream.
- The characters ‘1’ through ‘9’ denote one of the auxiliary input or history I/O streams.
- Double digit streams require “{}” braces: **i01{19}{24}**

State Entry: Defining a variable-set for an I/O stream

- Fields are added to a variable-set on an I/O stream in the Registry

#	Type	Sym	Dims	Use	Tlev	Stag	IO	Dname	Descrip
state	real	tsk	ij	misc	1	-	i01rhud	"TSK"	"SKIN TEMP"

usdf refers to nesting options:

u = UP, d = DOWN, s = SMOOTH, f = FORCE

u – at end of each set of child time steps

d – instantiation of child domain

f – at beginning of each set of child time steps

s – after each feedback

State Entry: Defining a variable-set for an I/O stream

Only variables involved with I/O, communications, packages are required to be state

Local variables inside of physics packages are not controlled by the Registry

Rconfig Entry

#	Type	Sym	How set	Nentries	Default
rconfig	integer	spec_bdy_width	namelist,bdy_control	1	1

- This defines namelist entries
- Elements
 - Entry:** the keyword “rconfig”
 - Type:** the type of the namelist variable (integer, real, logical, string)
 - Sym:** the name of the namelist variable or array
 - How set:** indicates how the variable is set: e.g. namelist or derived, and if namelist, which block of the namelist it is set in

Rconfig Entry

```
#      Type      Sym      How set      Nentries   Default
rconfig integer spec_bdy_width  namelist,bdy_control  1          1
```

- This defines namelist entries
- Elements
 - *Nentries*: specifies the dimensionality of the namelist variable or array. If 1 (one) it is a variable and applies to all domains; otherwise specify max_domains (which is an integer parameter defined in module_driver_constants.F).
 - *Default*: the default value of the variable to be used if none is specified in the namelist; hyphen (-) for no default

Package Entry

- Elements

- *Entry*: the keyword “package”,
- *Package name*: the name of the package: e.g. “kesslerscheme”
- *Associated rconfig choice*: the name of a rconfig variable and the value of that variable that chooses this package

```
# specification of microphysics options
package  passiveqv    mp_physics==0    -    moist:qv
package  kesslerscheme mp_physics==1    -    moist:qv,qc,qr
package  linscheme     mp_physics==2    -    moist:qv,qc,qr,qi,qs,qq
package  ncepcloud3   mp_physics==3    -    moist:qv,qc,qr
package  ncepcloud5   mp_physics==4    -    moist:qv,qc,qr,qi,qs

# namelist entry that controls microphysics option
rconfig integer  mp_physics  namelist,physics  max_domains  0
```

Package Entry

- Elements
 - *Package state vars*: unused at present; specify hyphen (-)
 - *Associated variables*: the names of 4D scalar arrays (*moist*, *chem*, *scalar*) and the fields within those arrays this package uses, and the state variables (*state:u_gc*, ...)

```
# specification of microphysics options
package  passiveqv    mp_physics==0    -    moist:qv
package  kesslerscheme mp_physics==1    -    moist:qv,qc,qr
package  linscheme     mp_physics==2    -    moist:qv,qc,qr,qi,qs,qq
package  ncepcloud3   mp_physics==3    -    moist:qv,qc,qr
package  ncepcloud5   mp_physics==4    -    moist:qv,qc,qr,qi,qs

# namelist entry that controls microphysics option
rconfig integer  mp_physics  namelist,physics  max_domains  0
```

Outline

- Examples
 - 1) Add output without recompiling
 - 2) Add a variable to the namelist
 - 3) Add an array
 - 4) Compute a diagnostic
 - 5) Add a physics package
 - 6) Tracer

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:RAINC,RAINNC
```

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

Example 1: Zap output without recompiling

- Edit the namelist.input file, the time_control namelist record

```
iofields_filename = "myoutfields.txt"
```

- Place the fields that you want in the named text file myoutfields.txt

```
- :h:0:W,PB,P
```

- Where “-” means REMOVE this variable from the output stream, “**h**” is the history stream, and “**0**” is the stream number (standard WRF history file)

Example 1: What streams can I use?

- Generally history streams 10 – 24 are OK

- Avoid 21, 22, 23

- Need LOTS more streams?

- Edit WRFV3/arch/preamble

MAX_HISTORY = 25 *<-- right now*

- clean –a, configure, compile, re-run real and wrf

Outline

- Examples

- 1) Add output without recompiling
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Example 2: Add a variable to the namelist

- Use the examples for the **rconfig** section of the Registry
- Find a namelist variable similar to what you want
 - Integer *vs* real *vs* logical *vs* character
 - Single value *vs* value per domain
 - Select appropriate namelist record
- Insert your mods in all appropriate Registry files

Example 2: Add a variable to the namelist

- Remember that ALL Registry changes require that the WRF code be cleaned and rebuilt

```
./clean -a  
./configure  
./compile em_real
```

Example 2: Add a variable to the namelist

- Adding a variable to the namelist requires the inclusion of a new line in the Registry file:

```
rconfig integer my_option_1 namelist,time_control 1 0 - "my_option_1" "test namelist option"  
rconfig integer my_option_2 namelist,time_control max_domains 0
```

- Accessing the variable is through an automatically generated function:

```
USE module_configure  
INTEGER :: my_option_1 , my_option_2  
  
CALL nl_get_my_option_1( 1, my_option_1 )  
CALL nl_set_my_option_2( grid%id, my_option_2 )
```

Example 2: Add a variable to the namelist

- You also have access to the namelist variables from the grid structure ...

```
SUBROUTINE foo ( grid , ... )  
  
USE module_domain  
TYPE(domain) :: grid  
  
print *,grid%my_option_1
```

Example 2: Add a variable to the namelist

- ... and you also have access to the namelist variables from config_flags

```
SUBROUTINE foo2 ( config_flags , ... )  
  
  USE module_configure  
  TYPE(grid_config_rec_type) :: config_flags  
  
  print *,config_flags%my_option_2
```

Example 2: Add a variable to the namelist

- What your variable looks like in the namelist.input file

```
&time_control  
run_days = 0,  
run_hours = 0,  
run_minutes = 40,  
run_seconds = 0,  
start_year = 2006, 2006, 2006,  
my_option_1 = 17  
my_option_2 = 1, 2, 3
```

Outline

- Examples
 - 1) Add output without recompiling
 - 2) Add a variable to the namelist
 - **3) Add an array**
 - 4) Compute a diagnostic
 - 5) Add a physics package
 - 6) Tracer

Example 3: Add an Array

- Adding a state array to the solver, requires adding a single line in the Registry
- Use the previous Registry instructions for a **state** or **l1** variable

Example 3: Add an Array

- Select a variable **similar** to one that you would like to add
 - 1d, 2d, or 3d
 - Staggered (X, Y, Z, or not “-”, *do not leave blank*)
 - Associated with a package
 - Part of a 4d array
 - Input (012), output, restart
 - Nesting, lateral forcing, feedback

Example 3: Add an Array

- Copy the “**similar**” field’s line and make a few edits
- Remember, no Registry change takes effect until a “clean -a” and rebuild

```
state real h_diabatic ikj misc 1 - r \
      "h_diabatic" "PREVIOUS TIMESTEP CONDENSATIONAL HEATING" \
state real msft      ij  misc 1 - i012rhdus=(copy_fcnm) \
      "MAPFAC_M"   "Map scale factor on mass grid" \
state real ht       ij  misc 1 - i012rhdus \
      "HGT"        "Terrain Height" \
state real ht_input  ij  misc 1 - - \
      "HGT_INPUT"  "Terrain Height from FG Input File" \
state real TSK_SAVE  ij  misc 1 - - \
      "TSK_SAVE"   "SURFACE SKIN TEMPERATURE" "K"
```

Example 3: Add an Array

- Always modify Registry.**core_name**_COMMON or Registry.**core_name**, where **core_name** might be **EM**

```
state real h_diabatic ikj misc 1 - r \
      "h_diabatic" "PREVIOUS TIMESTEP CONDENSATIONAL HEATING" \
state real msft      ij  misc 1 - i012rhdus=(copy_fcnm) \
      "MAPFAC_M"   "Map scale factor on mass grid" \
state real ht       ij  misc 1 - i012rhdus \
      "HGT"        "Terrain Height" \
state real ht_input  ij  misc 1 - - \
      "HGT_INPUT"  "Terrain Height from FG Input File" \
state real TSK_SAVE  ij  misc 1 - - \
      "TSK_SAVE"   "SURFACE SKIN TEMPERATURE" "K"
```

Example 3: Add an Array

- Add a new 3D array that is sum of all moisture species, called **all_moist**, in the Registry.**EM**_COMMON
 - Type: real
 - Dimensions: 3D and ikj ordering, not staggered
 - Supposed to be output only: h
 - Name in netCDF file: ALL_MOIST

```
state real all_moist ikj \
      misc 1 - h \
      "ALL_MOIST" \
      "sum of all of moisture species" \
      "kg kg-1"
```

Example 3: Add an Array

- Registry **state** variables become part of the derived data structure usually called **grid** inside of the WRF model.
- WRF model top → integrate → solve_interface → solve
- Each step, the **grid** construct is carried along for the ride
- No source changes for new output variables required until below the solver routine when dereferenced by first_rk_step_part1 for the physics drivers

Example 3: Add an Array

- Top of **solve_em.F**
- **grid** is passed in
- No need to declare any new variables, such as **all_moist**

```
!WRF : MEDIATION_LAYER : SOLVER
```

```
SUBROUTINE solve_em ( grid , &  
config_flags , &
```

Example 3: Add an Array

- In **solve_em**, add the new array to the call for the microphysics driver
- Syntax for **variable=local_variable** is an association convenience
- All state arrays are contained within **grid**, and must be **de-referenced**

```
CALL microphysics_driver( &  
    QV_CURR=moist(ims,kms,jms,P_QV), &  
    QC_CURR=moist(ims,kms,jms,P_QC), &  
    QR_CURR=moist(ims,kms,jms,P_QR), &  
    QI_CURR=moist(ims,kms,jms,P_QI), &  
    QS_CURR=moist(ims,kms,jms,P_QS), &  
    QG_CURR=moist(ims,kms,jms,P_QG), &  
    QH_CURR=moist(ims,kms,jms,P_QH), &  
    all_moist=grid%all_moist , &
```

Example 3: Add an Array

- After the array is re-referenced from **grid** and we are **inside the microphysics_driver** routine, we need to
 - Pass the variable through the argument list
 - Declare our passed in 3D array

```
,all_moist &
```

```
REAL, DIMENSION(ims:ime ,kms:kme ,jms:jme ), &  
INTENT(OUT) :: all_moist
```

Example 3: Add an Array

- After the array is re-referenced from grid and we are **inside the microphysics_driver routine**, we need to
 - Zero out the array at each time step

```
! Zero out moisture sum.

DO j = jts,MIN(jde-1,jte)
DO k = kts,kte
DO i = its,MIN(ide-1,ite)
  all_moist(i,k,j) = 0.0
END DO
END DO
END DO
```

Example 3: Add an Array

- After the array is re-referenced from grid and we are **inside the microphysics_driver routine**, we need to
 - At the end of the routine, for each of the **moist species that exists**, add that component to **all_moist**

```
DO j = jts,MIN(jde-1,jte)
DO k = kts,kte
IF ( f_qv ) THEN
  DO i = its,MIN(ide-1,ite)
    all_moist(i,k,j) = all_moist(i,k,j) + &
      qv_curr(i,k,j)
  END DO
END IF
```

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 - 4) **Compute a diagnostic**
 - 5) Add a physics package
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Example 4: Compute a Diagnostic

- Problem: Output global average and global maximum and lat/lon location of maximum for 10 meter wind speed in WRF
- Steps:
 - Modify solve to compute wind-speed and then compute the local sum and maxima at the end of each time step
 - Use reduction operations built-in to WRF software to compute the global qualities
 - Output these on one process (process zero, the “monitor” process)

Example 4: Compute a Diagnostic

- Compute local sum and local max and the local indices of the local maximum

```
--- File: dyn_em/solve_em.F  (near the end) ---  
  
! Compute local maximum and sum of 10m wind-speed  
sum_ws = 0.  
max_ws = 0.  
DO j = jps, jpe  
  DO i = ips, ipe  
    wind_vel = sqrt( grid%u10(i,j)**2+ grid%v10(i,j)**2 )  
    IF ( wind_vel .GT. max_ws ) THEN  
      max_ws = wind_vel  
      idex = i  
      jdex = j  
    ENDIF  
    sum_ws = sum_ws + wind_vel  
  ENDDO  
ENDDO
```

Example 4: Compute a Diagnostic

- On the process that contains the maximum value, obtain the latitude and longitude of that point; on other processes set to an artificially low value.
- The use parallel reduction to store that result on every process

```
IF ( ips .LE. idex .AND. idex .LE. ipe .AND. &  
     jps .LE. jdex .AND. jdex .LE. jpe ) THEN  
  glat = grid%xlat(idex,jdex)  
  glon = grid%xlong(idex,jdex)  
ELSE  
  glat = -99999.  
  glon = -99999.  
ENDIF  
  
! Compute global maximum to find glat and glon  
glat = wrf_dm_max_real ( glat )  
glon = wrf_dm_max_real ( glon )
```

Example 4: Compute a Diagnostic

- Compute global sum, global max, and indices of the global max (WRF intrinsics)

```
! Compute global sum  
sum_ws = wrf_dm_sum_real ( sum_ws )  
  
! Compute global maximum and associated i,j point  
CALL wrf_dm_maxval_real ( max_ws, idex, jdex )
```

Example 4: Compute a Diagnostic

Example 4: Compute a Diagnostic

- Output the value on process zero, the “monitor”

```
! Print out the result on the monitor process  
IF ( wrf_dm_on_monitor() ) THEN  
  WRITE(outstring,*) 'Avg. ',sum_ws/((ide-ids+1)*(jde-jds+1))  
  CALL wrf_message ( TRIM(outstring) )  
  WRITE(outstring,*) 'Max. ',max_ws,' Lat. ',glat,&  
                   ' Lon. ',glon  
  CALL wrf_message ( TRIM(outstring) )  
ENDIF
```

Example 4: Compute a Diagnostic

- Output from process zero of a multi-process run

```
--- Output file: rsl.out.0000 ---
.
.
.
Avg. 5.159380
Max. 15.09370 Lat. 37.25022 Lon. -67.44571
Timing for main: time 2000-01-24_12:03:00 on domain 1: 8.96500 elapsed secs.
Avg. 5.166167
Max. 14.97418 Lat. 37.25022 Lon. -67.44571
Timing for main: time 2000-01-24_12:06:00 on domain 1: 4.89460 elapsed secs.
Avg. 5.205693
Max. 14.92687 Lat. 37.25022 Lon. -67.44571
Timing for main: time 2000-01-24_12:09:00 on domain 1: 4.83500 elapsed secs.
.
```

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Example 5: Input periodic SSTs

- Add a new physics package with time varying input source to the model
- This is how we could supply a time varying value to the model for a field that is traditionally fixed
- Example is sea surface temperature

Example 5: Input periodic SSTs

- Problem: adapt WRF to input a time-varying lower boundary condition, e.g. SSTs, from an input file for a new surface scheme
- Given: Input file in WRF I/O format containing 12-hourly SST's
- Modify WRF model to read these into a new state array and make available to WRF surface physics

Example 5: Input periodic SSTs

- Steps
 - Add a new state variable and definition of a new surface layer package (that will use the variable) to the Registry
 - Add to variable stream for an unused Auxiliary Input stream
 - Adapt physics interface to pass new state variable to physics
 - Setup namelist to input the file at desired interval

Example 5: Input periodic SSTs

- Add a new state variable to Registry/Registry.EM_COMMON and put it in the variable set for input on Auxiliary Input Stream #4

```
#      type   symbol dims use   tl stag   io      dname      description      units
state real    nsst    ij   misc   1   -   i4h   "NEW_SST"  "Time Varying SST"  "K"
```

- Also added to History and Restart
- Result:
 - 2-D variable named grid%nsst defined and available in solve_em
 - Dimensions: ims:ime, jms:jme
 - Input and output on the AuxInput #4 stream will include the variable under the name NEW_SST

Example 5: Input periodic SSTs

- Pass new state variable to surface physics

```
-- File: dyn_em/module_first_rk_step_part1.F ---

CALL surface_driver(
  . .
  ! Optional
  & ,QV_CURR=moist(ims,kms,jms,P_QV), F_QV=F_QV
  & ,QC_CURR=moist(ims,kms,jms,P_QC), F_QC=F_QC
  & ,QR_CURR=moist(ims,kms,jms,P_QR), F_QR=F_QR
  & ,QI_CURR=moist(ims,kms,jms,P_QI), F_QI=F_QI
  & ,QS_CURR=moist(ims,kms,jms,P_QS), F_QS=F_QS
  & ,QG_CURR=moist(ims,kms,jms,P_QG), F_QG=F_QG
  & ,NSST=grid%nsst & ! new
  & ,CAPG=grid%capg, EMISS=grid%emiss, HOL=hол, MOL=grid%mol
  & ,RAINBL=grid%rainbl, SR=grid%em_sr
  & ,RAINNCV=grid%rainncv, REGIME=regime, T2=grid%t2, THC=grid%thc
  . . .
```

Example 5: Input periodic SSTs

- Add new variable nsst to Physics Driver in Mediation Layer

```
--- File: phys/module_surface_driver.F ---

SUBROUTINE surface_driver(
  . .
  ! Other optionals (more or less em specific)
  & ,nsst
  & ,capg,emiss,hол,mол
  & ,rainncv,rainbl,regime,t2,thс
  & ,qsg,qvg,qcg,soiltl,tsnav
  & ,smfr3d,keepfr3dfлаг
  . .
  ))
```

REAL, DIMENSION(ims:ime, jms:jme), OPTIONAL, INTENT(INOUT) :: nsst

- By making this an “Optional” argument, we preserve the driver’s compatibility with other cores and with versions of WRF where this variable hasn’t been added.

Example 5: Input periodic SSTs

- Add call to Model-Layer subroutine for new physics package to Surface Driver

```
--- File: phys/module_surface_driver ---  
  
!$OMP PARALLEL DO  &  
!$OMP PRIVATE ( ij, i, j, k )  
DO ij = 1 , num_tiles  
    sfclay_select: SELECT CASE(sf_sfclay_physics)  
  
        CASE (SFCLAYScheme)  
  
            CASE (NEWSFCScheme) ! <- This is defined by the Registry "package" entry  
                IF (PRESENT(nsst)) THEN  
                    CALL NEWSFCSCHEME(  
                        nsst,  
                        ids,ide, jds,jde, kds,kde,  
                        ims,ime, jms,jme, kms,kme,  
                        i_start(ij),i_end(ij), j_start(ij),j_end(ij), kts,kte      )  
                ELSE  
                    CALL wrf_error_fatal('Missing argument for NEWScheme in surface driver')  
                ENDIF  
  
            END SELECT sfclay_select  
        ENDDO  
    !$OMP END PARALLEL DO
```

- Note the PRESENT test to make sure new optional variable nsst is available

Example 5: Input periodic SSTs

- Add definition for new physics package NEWScheme as setting 4 for namelist variable sf_sfclay_physics

rconfig	integer	sf_sfclay_physics	namelist,physics	max_domains	0
package	sfclayscheme	sf_sfclay_physics==1	-	-	
package	myjsfcscsche	sf_sfclay_physics==2	-	-	
package	gfssfcscsche	sf_sfclay_physics==3	-	-	
package	newsfcscsche	sf_sfclay_physics==4	-	-	

- This creates a defined constant NEWSFCSCHEME and represents selection of the new scheme when the namelist variable sf_sfclay_physics is set to '4' in the namelist.input file
- **clean -a** and recompile so code and Registry changes take effect

Example 5: Input periodic SSTs

- Setup namelist to input SSTs from the file at desired interval

```
--- File: namelist.input ---  
  
&time_control  
    . . .  
    auxinput4_inname      = "sst_input"  
    auxinput4_interval_h = 12  
    . . .  
/  
    . . .  
&physics  
    sf_sfclay_physics  = 4, 4, 4  
    . . .  
/
```

- Run code with sst_input file in run-directory

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Tracer Example

Modify Registry for new fields.

Use the “tracer” array with a new 3D component

Use existing NML option

Initialize data in real.

Identify (i,j) location

Spread in “PBL”

Set values in solver.

“Release” per time step



Tracer Example

Registry/Registry.EM add our new field “PLUME” as part of “TRACER” array.

```
# New tracer for example
state real plume ikjftb tracer \
1 - irhusdf=(bdy_interp:dt) \
"PLUME" "Fukushima Tracer" " "

# 4D arrays need an associated package
package tracer_test3 tracer_opt==3 - \
tracer:plume
```

Tracer Example

Modify the real and WRF programs to initialize and continuously re-supply the

“PLUME” array

dyn_em/module_initialize_real.F (initial value from real.exe)

dyn_em/solve_em.F (continuous plume in wrf.exe)

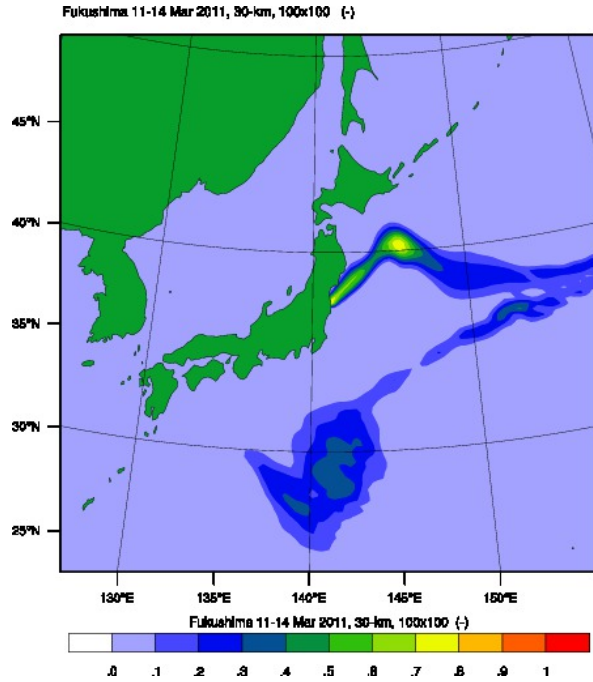
```
! Add in the Fukushima initial venting.
```

```
IF ( ( its .LE. 50 ) .AND. ( ite .GE. 50 ) .AND. &
( jts .LE. 50 ) .AND. ( jte .GE. 50 ) ) THEN
tracer(50,1:5,50,P_plume) = 1.
END IF
```

Tracer Example

- Modify the test/em_real/namelist.input file
- Include the new settings for the tracer option required from the Registry file

```
&dynamics
tracer_opt = 3, 3, 3,
```



Outline

- What is the WRF Registry
- Keyword syntax
- The BIG Three
- Examples
 - Runtime I/O mods
 - Adding a variable to the namelist
 - Adding an array to WRF