

PDAF Tutorial

Implementation of the analysis step in offline mode



<http://pdaf.awi.de>

PDAF Parallel
Data Assimilation
Framework

Implementation Tutorial for PDAF offline

We demonstrate the implementation
of an offline analysis step with PDAF
using the template routines provided by PDAF

The example code is part of the PDAF source code package
downloadable at <http://pdaf.awi.de>

(This tutorial is compatible with PDAF V1.16 and later)

Implementation Tutorial for PDAF offline

This is just an example!

For the complete documentation of PDAF's interface
see the documentation
at <http://pdaf.awi.de>

Overview

Focus on Error Subspace Transform Kalman Filter
(ESTKF, Nerger et al., Mon. Wea. Rev. 2012)

4 Parts

1. Without parallelization
 - a) Global filter
 - b) Localized filter
(and OpenMP-parallelization)
2. With MPI-parallelization
 - a) Global filter
 - b) Localized filter

We recommend to first implement the global filter. The localized filter re-uses routines of the global filter.

We assume that 1a is implemented before 1b and 1a is implemented before 2a (1b before 2b).

Contents

0a) Files for the tutorial	6 - 9
0b) The model	10 - 15
0c) State vector and observation vector	16 - 18
0d) PDAF offline mode	19 - 22
1a) Global filter without parallelization	23 - 45
1b) Local filter without parallelization	46 - 71
1b.1) Use local filter with OpenMP-parallelization	72 - 79
2a) Parallelized global filter	80 - 96
2b) Parallelized local filter	97 - 111
3) Hints for adaptations for real models	112 - 116

0a) Files for the Tutorial

Tutorial implementation

Files are in the PDAF package

Directories:

`/tutorial/classical/offline_2D_serial` (OpenMP-parallelization)

`/tutorial/classical/offline_2D_parallel` (MPI parallelization)

- Fully working implementations of user codes
- PDAF core files are in `/src`
Makefile refers to it and compiles the PDAF library
- Only need to specify the compile settings (compiler, etc.) by environment variable `PDAF_ARCH`. Then compile with 'make'.

Templates for offline mode

Directory: `/templates/offline`

- Contains all required files
- Contains also
command line parser, memory counting, timers
(convenient but not required)

To generate your own implementation:

1. Copy directory to a new name
2. Complete routines for your model
3. Set base directory (`BASEDIR`) in Makefile
4. Set `$PDAF_ARCH`
5. Compile

PDAF library

Directory: `/src`

- The PDAF library is not part of the template
- PDAF is compiled separately as a library and linked when the assimilation program is compiled
- Makefile includes a compile step for the PDAF library
- One can also `cd` to `/src` and run 'make' there (requires setting of `PDAF_ARCH`)

`$PDAF_ARCH`

- Environment variable to specify the compile specifications
- Definition files in `/make.arch`
- Define by, e.g.

```
setenv PDAF_ARCH linux_gfortran (tcsh/csh)  
export PDAF_ARCH=linux_gfortran (bash)
```

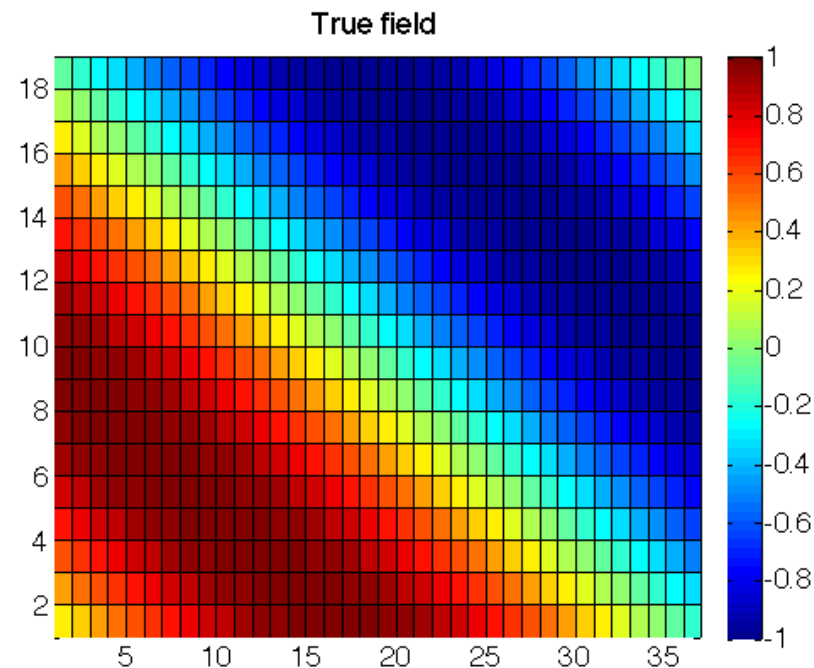
0b) The Model

Simple assimilation problem

- 2-dimensional model domain
- One single field (like temperature)
- Direct measurements of the field
- Data gaps (i.e. data at selected grid points)
- Same error estimate for all observations
- Observation errors are not correlated (diagonal observation error covariance matrix)
- Perform a single analysis step using input files for the ensemble and observations (offline mode: No time stepping in the assimilation program)

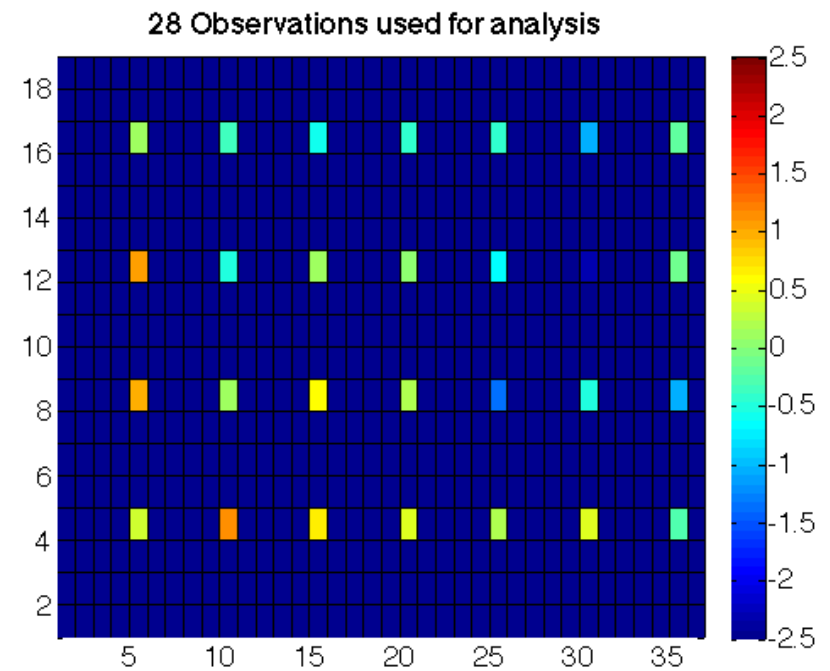
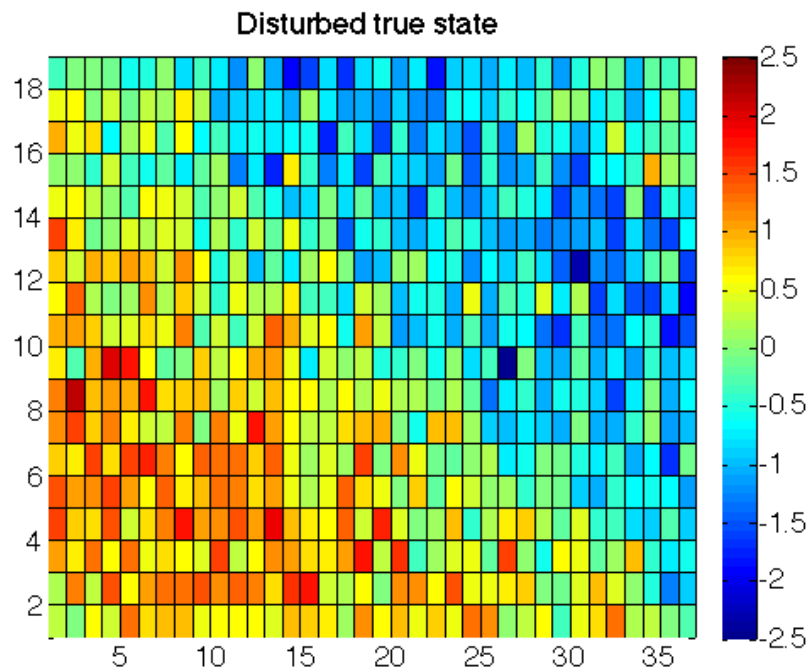
2D „Model“

- Simple 2-dimensional grid domain
- 36 x 18 grid points (longitude x latitude)
- True state: sine wave in diagonal direction
- No dynamics for offline mode
- Stored in text file (18 rows) – `true.txt`



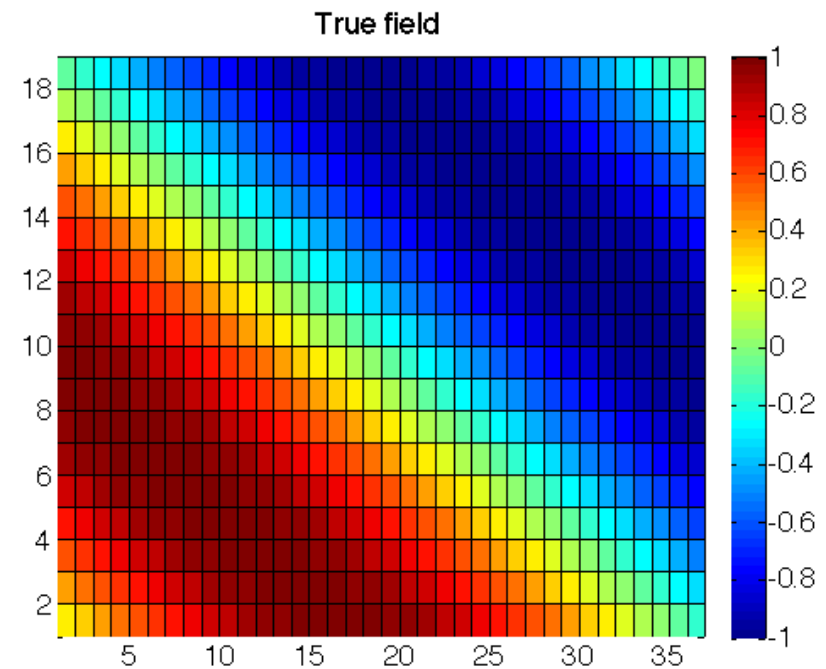
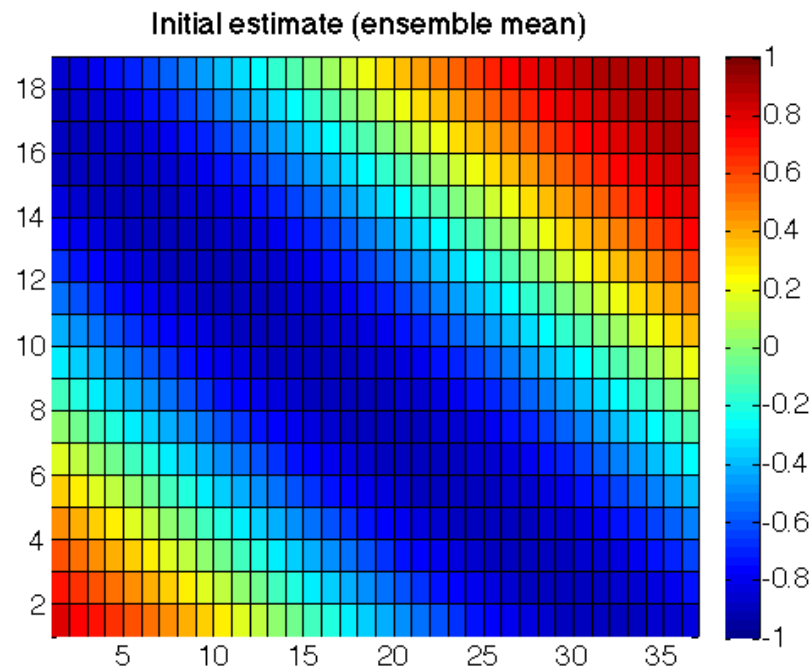
Observations

- Add random error to true state (standard deviation 0.5)
- Select a set of observations at 28 grid points
- File storage:
text file, full 2D field, -999 marks 'no data' – obs.txt

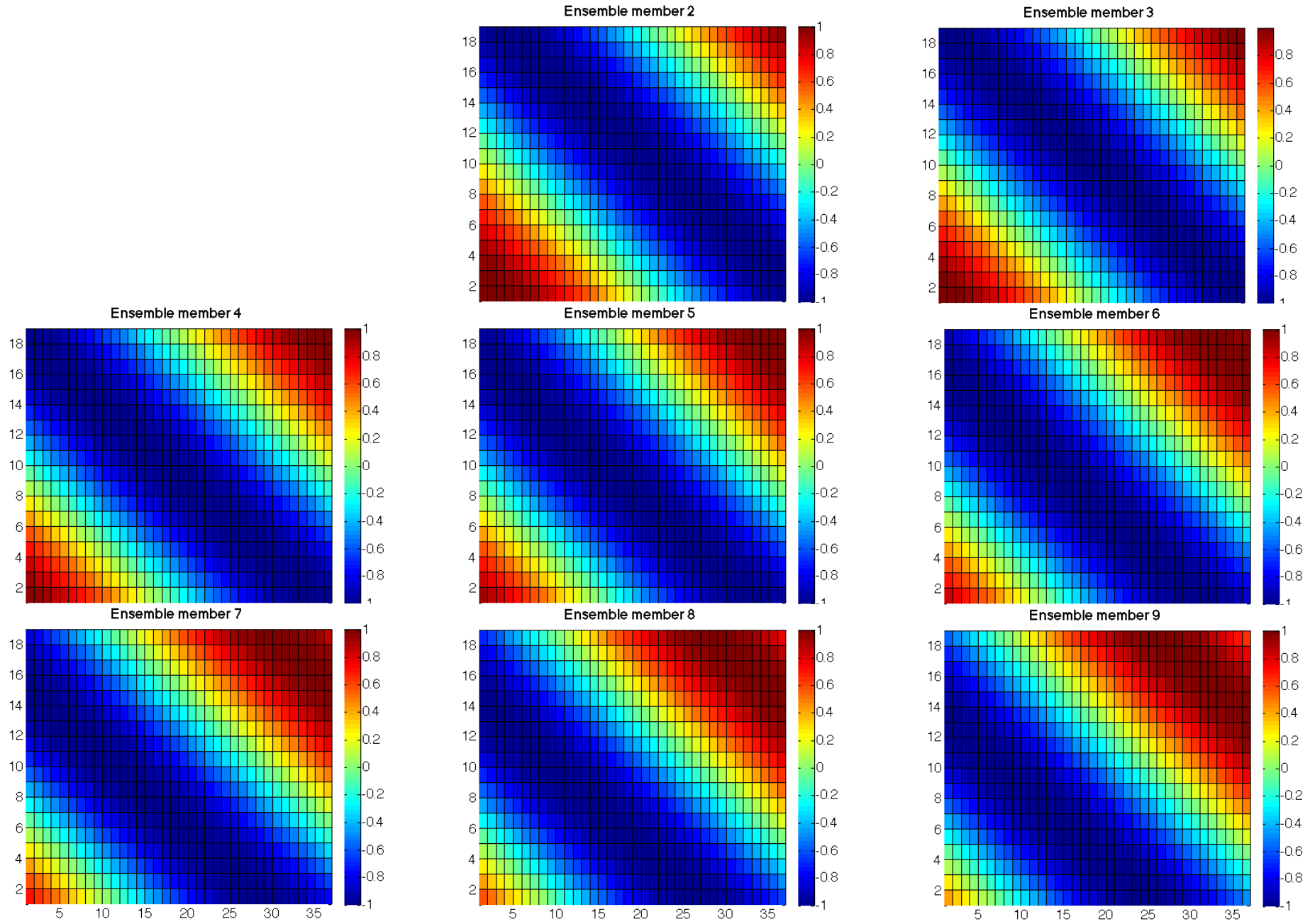


Ensemble

- Ensemble size 9
- Sine waves shifted along diagonal (truth not included)
- One text file per ensemble member – ens* .txt



Ensemble states



0c) state vector and observation vector

State vector – some terminology used later

- PDAF performs computations on state vectors
- **State vector**
 - Stores model fields in a single vector
 - Tutorial shows this for one 2-dimensional field
 - Multiple fields are just concatenated into the vector
 - All fields that should be modified by the assimilation have to be in the state vector
- **State dimension**
 - Is the length of the state vector
(the sum of the sizes of the model fields in the vector)
- **Ensemble array**
 - Rank-2 array which stores state vectors in its columns

Observation vector

- **Observation vector**
 - Stores all observations in a single vector
 - Tutorial shows this for one 2-dimensional field
 - Multiple observed fields are just concatenated into the vector
- **Observation dimension**
 - Is the length of the observation vector
(sum of the observations over all observed fields in the vector)
- **Observation operator**
 - Operation that computes the observed part of a state vector
 - Tutorial only selects observed grid points
 - The operation can involve interpolation or integration depending on type of observation

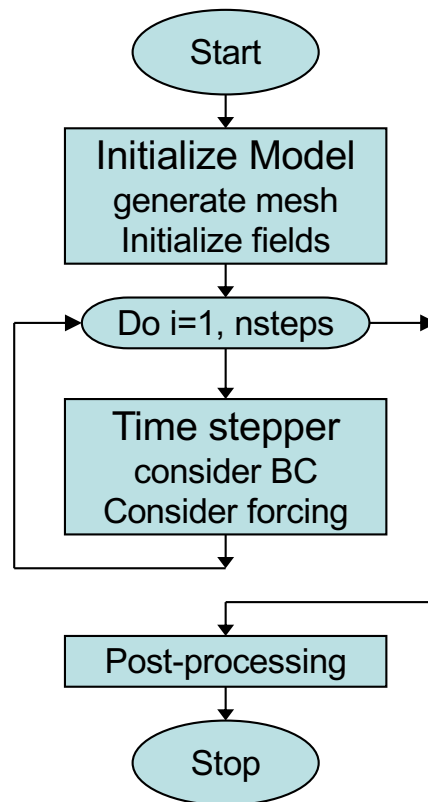
0d) PDAF offline mode

Offline mode

- Two separate programs
 - “Model” – performs ensemble integrations
 - “PDAF_offline” – perform analysis step
- Couple both programs through files
 1. “PDAF_offline” reads ensemble forecast files
 2. Performs analysis step
 3. Writes analysis ensemble files (restart files for “Model”)
 4. “Model” reads restart files and performs ensemble integration

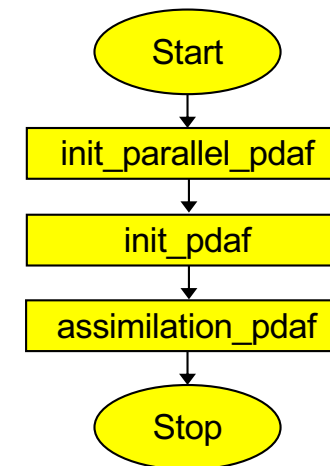
Programs in offline mode

Model



- Run for each ensemble member
- Write restart files

Assimilation program



- Read restart files (ensemble)
- Compute analysis step
- Write new restart files

PDAF_offline: General program structure

```
program main_offline
  init_parallel_pdaf
      initialize communicators
      (not relevant without parallelization)

  initialize
      initialize model information

  init_pdaf
      initialize parameters for PDAF
      and read ensemble

  assimilation_pdaf
      perform analysis
      (by call to PDAF_put_state_X)

end program
```

1a) Global filter without parallelization

Running the tutorial program

- Do `cd /tutorial/classical/offline_2D_serial`
- Set environment variable `PDAF_ARCH` or specify it when running `make` (e.g. `linux_gfortran`)
- Compile by running `'make'` (or `'make PDAF_ARCH=...'`) (next slide will discuss possible compile issues)
- Run the program with `./PDAF_offline`

- Inputs are read in from `/tutorial/inputs_offline`
- Outputs are written in `/tutorial/classical/offline_2D_serial`
- Plot result, e.g. with Python:

```
python ../plotting/plot_file.py state_ana.txt
```


Requirements for compiling PDAF

PDAF requires libraries for BLAS and LAPACK

- Libraries to be linked are specified in the include file for make in `/make.arch` (file according to `PDAF_ARCH`)
- For `$PDAF_ARCH=linux_gfortran` the specification is

```
LINK_LIBS =-L/usr/lib -llapack -lblas -lm
```
- If the libraries are at another non-default location, one has to change the directory name (`/usr/lib`)
- Some systems or compilers have special libraries (e.g. MKL for ifort compiler, or ESSL on IBM/AIX)

PDAF needs to be compiled for double precision

- Needs to be set at compiler time in the include file for make:
- For gfortran: `OPT = -O3 -fdefault-real-8`

Files in the tutorial implementation

/tutorial/inputs_offline

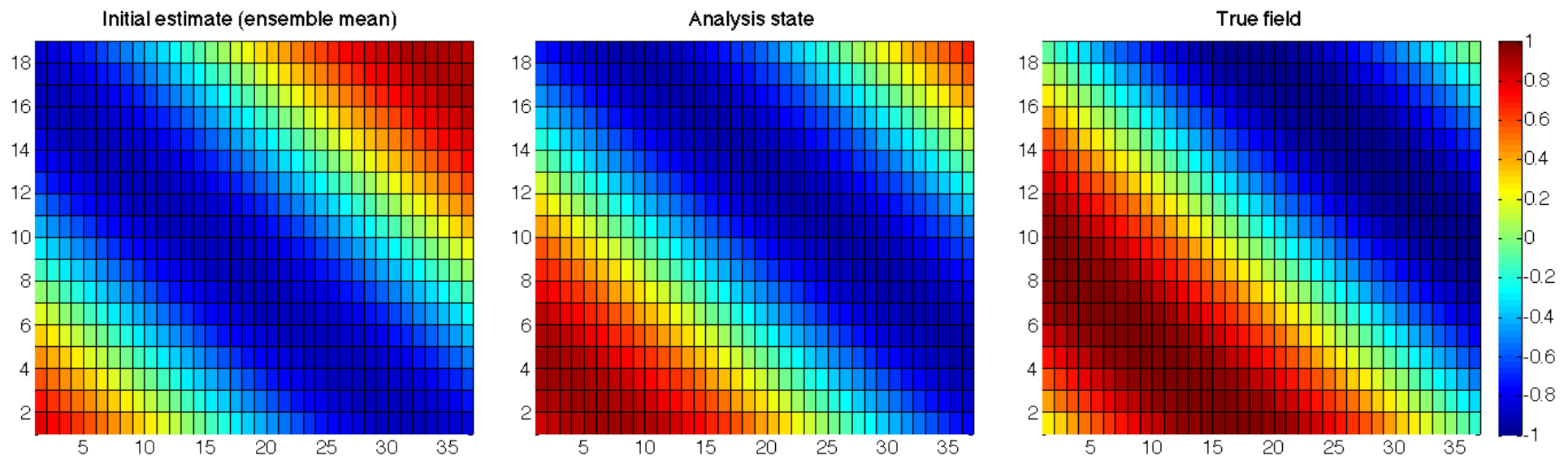
- `true.txt` true state
- `state_ini.txt` initial estimate (ensemble mean)
- `obs.txt` observations
- `ens_X.txt` ($X=1, \dots, 9$) ensemble members

/tutorial/classical/offline_2D_serial
(after running PDAF_offline)

- `state_ana.txt` analysis state estimate
- `ens_X_ana.txt` ($X=1, \dots, 9$) analysis ensemble members

Result of the global assimilation

- The analysis state is closer to the true field than the initial estimate
- Truth and analysis are not identical (the ensemble does not allow it)



Files to be changed

Template contains all required files

- just need to be filled with functionality

`mod_assimilation.F90`

} Fortran module

`initialize.F90`

`init_pdaf_offline.F90`

`init_ens_offline.F90`

} initialization

`init_dim_obs_pdaf.F90`

`obs_op_pdaf.F90`

`init_obs_pdaf.F90`

`prodrinva_pdaf.F90`

} analysis step

`prepoststep_ens_offline.F90`

} post step

mod_assimilation.F90

Fortran module

- Declares the parameters used to configure PDAF
- Add model-specific variables here
(see next slides)
- Will be included (with 'use') in the user-written routines

initialize.F90

Routine initializes the model information

1. Define 2D mesh in mod_assimilation.F90

```
integer :: nx, ny
```

2. In initialize.F90 include nx, ny, and dim_state_p with use mod_assimilation

3. Define mesh size in initialize.F90

```
nx = 36
```

```
ny = 18
```

4. Define state dimension in initialize.F90

```
dim_state_p = nx * ny
```

Note: Some variables end with `_p`.

It means that the variable is specific for a process.

(Not relevant until we do parallelization)

init_pdaf_offline.F90

Routine sets parameters for PDAF and calls `PDAF_init` to initialize the data assimilation:

Template contains list of available parameters (declared in and used from `mod_assimilation`)

For the example set :

1. `dim_ens = 9`
2. `rms_obs = sqrt(0.5)`
3. `filtertype = 6` (for ESTKF)

In call to `PDAF_init`, the name of the ensemble initialization routine is specified:

```
init_ens_offline
```

init_ens_offline.F90

A *call-back* routine called by PDAF_init:

- Implemented by the user
- Its name is specified in the call to PDAF_init
- It is called by PDAF through a defined interface:

```
SUBROUTINE init_ens_offline(filtertype, dim_p,  
                           dim_ens, state_p, Uinv, ens_p, flag)
```

Declarations in header of the routine shows “intent” (input, output):

```
REAL, INTENT(out)      :: ens_p(dim_p, dim_ens)
```

Note:

All call-back routines have a defined interface and show the intent of the variables. Their header comment explains what is to be done in the routine.

init_ens_offline.F90 (2)

Initialize ensemble matrix `ens_p`

1. Include `nx, ny` with `use mod_assimilation`
2. Declare and allocate `real :: field(ny, nx)`
3. Loop over ensemble files (`i=1,dim_ens`)

for each file:

- read ensemble state into `field`
- store contents of `field` in column `i` of `ens_p`

Note:

Columns of `ens_p` are state vectors.

Store following storage of `field` in memory (column-wise in Fortran)

The analysis step

At this point the initialization of PDAF is complete:

- Forecast ensemble is initialized
- Filter algorithm and its parameters are chosen

Next:

- Implement user-routines for analysis step
- All are call-back routines:
 - User-written, but called by PDAF

Note:

Some variables end with `_p`.

It means that the variable is specific for a process.
(Not relevant until we do parallelization)

init_dim_obs_pdaf.F90

Routine to

- read observation file
- count number of available observations
(direct output to PDAF: **dim_obs_p**)

Optional, also

- initialize array holding available observations
- initialize index array telling index of observation point
in full state vector

The most complicated routine in the example!
(but less than 100 lines)

init_dim_obs_pdaf.F90 (2)

Preparations and reading of observation file:

1. Include `nx`, `ny` with use `mod_assimilation`
2. declare and allocate real array `obs_field(ny, nx)`
3. read observation file:

```
OPEN (12, file='inputs_offline/obs.txt', &  
      status='old')  
  
DO i = 1, ny  
    READ (12, *) obs_field(i, :)  
END DO  
  
CLOSE (12)
```

init_dim_obs_pdaf.F90 (3)

Count available observations (**dim_obs_p**):

1. Declare integer `:: cnt, cnt0`
2. Now count

```
cnt = 0
DO j = 1, nx
  DO i= 1, ny
    IF (obs_field(i,j) > -999.0) cnt = cnt + 1
  END DO
END DO
dim_obs_p = cnt
```

init_dim_obs_pdaf.F90 (4)

Initialize observation vector (obs)
and index array (obs_index):

1. In mod_assimilation it is declared
`real, allocatable :: obs_p(:), obs_index_p(:)`
Include these variable with `use mod_assimilation`

2. Allocate

`obs_p(dim_obs_p), obs_index_p(dim_obs_p)`
(If already allocated, deallocate first)

3. Now initialize ...

Note:

The arrays only contain information about valid observations;
one could store observations already in files in this way.

init_dim_obs_pdaf.F90 (5)

3. Now initialize

```
cnt0 = 0           ! Count grid points
cnt = 0           ! Count observations
DO j = 1, nx
  DO i= 1, ny
    cnt0 = cnt0 + 1
    IF (obs_field(i,j) > -999.0) THEN
      cnt = cnt + 1
      obs_index_p(cnt) = cnt0           ! Index
      obs_p(cnt) = obs_field(i, j) ! observations
    END IF
  END DO
END DO
```

obs_op_pdaf.F90

Implementation of observation operator
acting on some state vector

Input: state vector `state_p`

Output: observed state vector `m_state_p`

1. Include `obs_index_p` by use `mod_assimilation`
2. Select observed grid points from state vector:

```
DO i = 1, dim_obs_p
    m_state_p(i) = state_p(obs_index_p(i))
END DO
```

Note:

`dim_obs_p` is an input argument of the routine

init_obs_pdaf.F90

Fill PDAF's observation vector

Output: vector of observations `observation_p`

1. Include obs by use `mod_assimilation`
2. Initialize `observation_p`:

```
observation_p = obs_p
```

Note:

This is trivial, because of the preparations in `init_dim_obs_pdaf`!

(However, the operations needed to be separate, because PDAF allocates `observations_p` after the call to `init_dim_obs_pdaf`)

prodrinva_pdaf.F90

Compute the product of the inverse observation error covariance matrix with some other matrix

- Input: Matrix $A_p(\text{dim_obs_p}, \text{rank})$
- Output: Product matrix $C_p(\text{dim_obs_p}, \text{rank})$
(rank is typically $\text{dim_ens}-1$)

1. Declare and initialize inverse observation error variance
`ivariance_obs = 1.0 / rms_obs**2`

2. Compute product:

```
DO j = 1, rank
  DO i = 1, dim_obs_p
    C_p(i, j) = ivariance_obs * A_p(i, j)
  END DO
END DO
```

prepoststep_ens_offline.F90

Post-step routine for the offline mode:

Already there in the template:

1. Compute ensemble mean state `state_p`
2. Compute estimated variance vector `variance`
3. Compute estimated root mean square error `rmsest`

Required extension:

4. Write analysis ensemble into files used for model restart
(Analogous to reading in `init_ens_pdaf_offline`)

Possible (useful) extension:

5. Write analysis state (ensemble mean, `state_ana.txt`)

Done!

The analysis step in offline mode is fully implemented now

The implementation allows you now to use the global filters ESTKF, ETKF, and SEIK

Not usable are EnKF and SEEK (The EnKF needs some other user files und SEEK a different ensemble initialization)

A complete analysis step

We now have a fully functional analysis step
- if no localization is required!

Possible extensions for a real application:

Adapt routines for

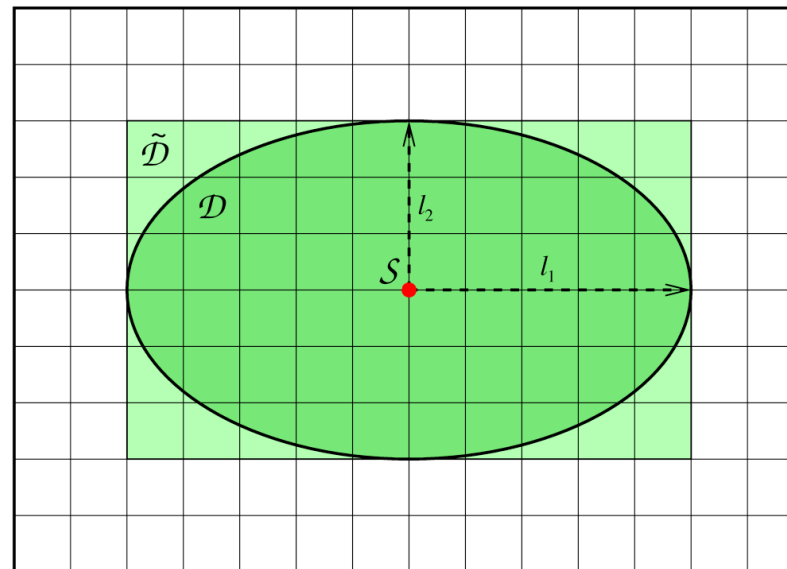
- Multiple model fields
 - Store full fields consecutively in state vector
- Third dimension
 - Extend state vector
- Different observation types
 - Store different types consecutively in observation vector
- Other file type (e.g. binary or NetCDF)
 - Adapt reading/writing routines

1b) Local filter without parallelization

Localization

Localization is usually required for high-dimensional systems

- Update small regions (S)
(e.g. single grid points, single vertical columns)
- Consider only observations within cut-off distance (D)
- Weight observations according to distance from S



The FULL observation vector

- A single local analysis at S (single grid point) need observations from domain D
- A loop of local analyses over all S needs all observations
 - This defines the *full* observation vector
- Why distinguish *full* and *all* observations?
 - They can be different in case of parallelization!
- Example:
 - Split domain in left and right halves
 - Some of the analyses in left half need observations from the right side.
 - Depending on localization radius not all observations from the right side might be needed for the left side analyses

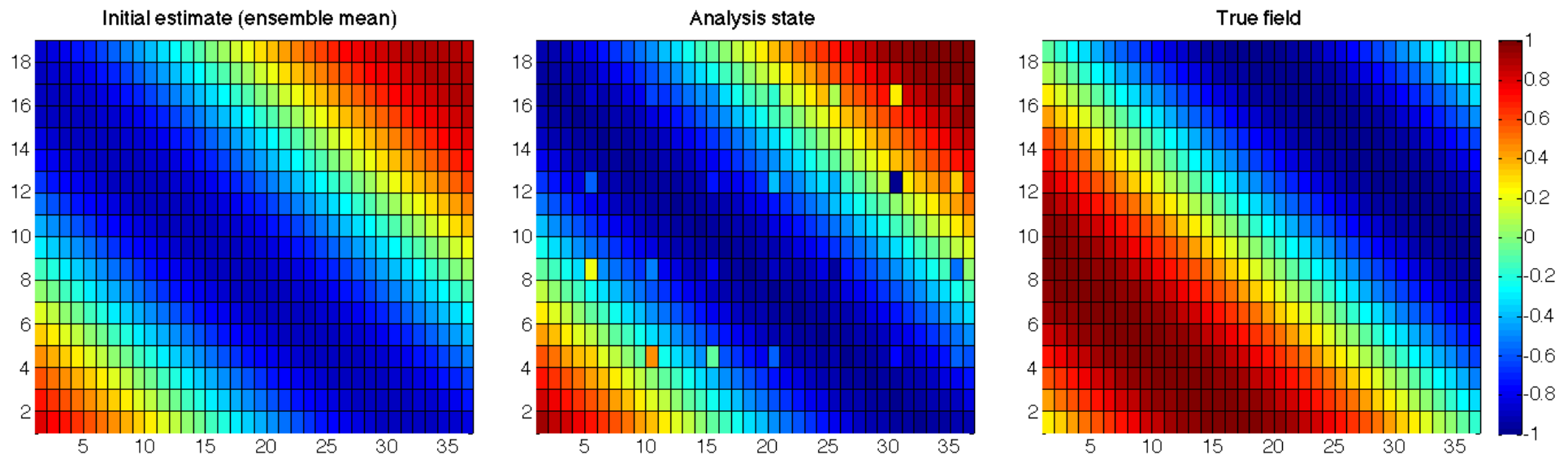
Running the tutorial program

- Compile as for the global filter
- Run the program with `./PDAF_offline OPTIONS`
- `OPTIONS` are always of type `–KEYWORD VALUE`
- Possible `OPTIONS` are
 - `-filtertype 7` (select LESTKF if not set in `init_pdaf_offline`)
 - `-local_range 5.0` (set localization radius, 0.0 by default, any positive value should work)
 - `-locweight 2` (set weight function for localization, default=0 for constant weight of 1; possible are integer values 0 to 4; see `init_pdaf_offline`)

Result of the local assimilation

```
./PDAF_offline -filtertype 7
```

- Default: zero localization radius (local_range=0.0)
- Change only at observation locations

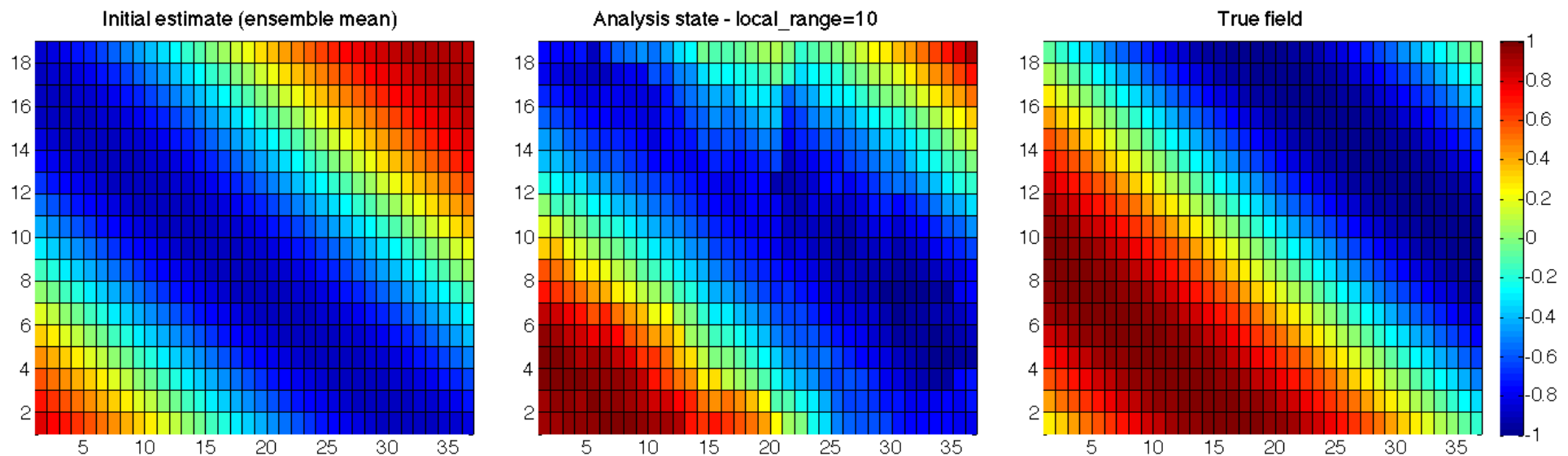


Result of the local assimilation (2)

```
./PDAF_offline -filtertype 7 -local_range 10.0
```

- All local analysis domains are influenced (all see observations)
- Up to 16 observations in a single local analysis (average 9.6)

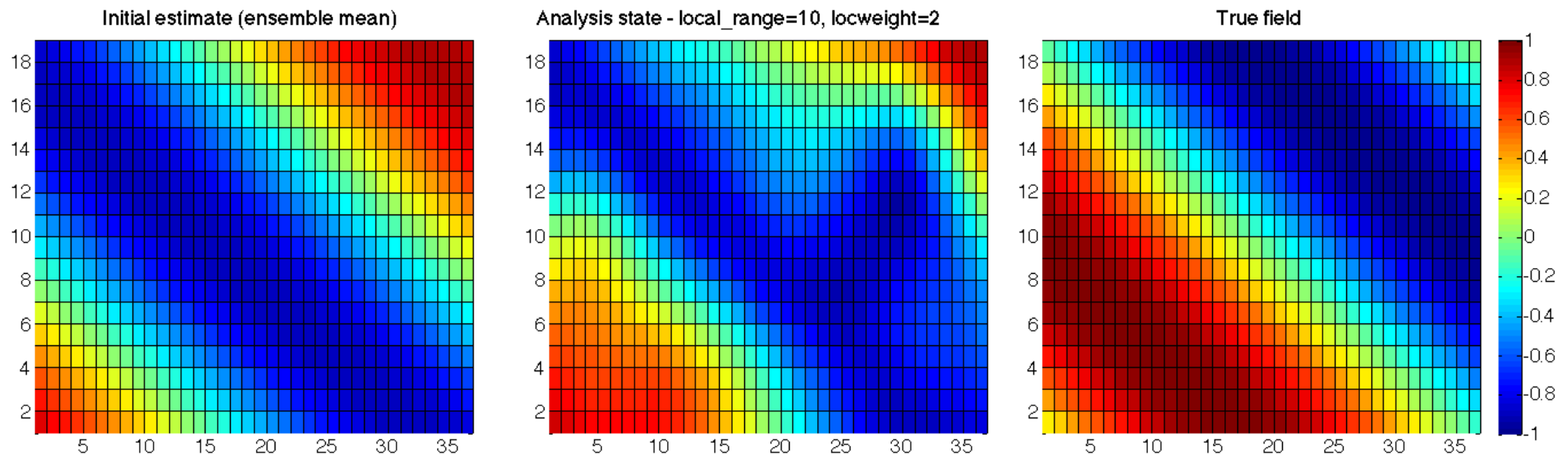
Note: The set up of the experiment favors the global filter because of the shape of the ensemble members



Result of the local assimilation (2)

```
./PDAF_offline -filtertype 7 -local_range 10.0 -locweight 2
```

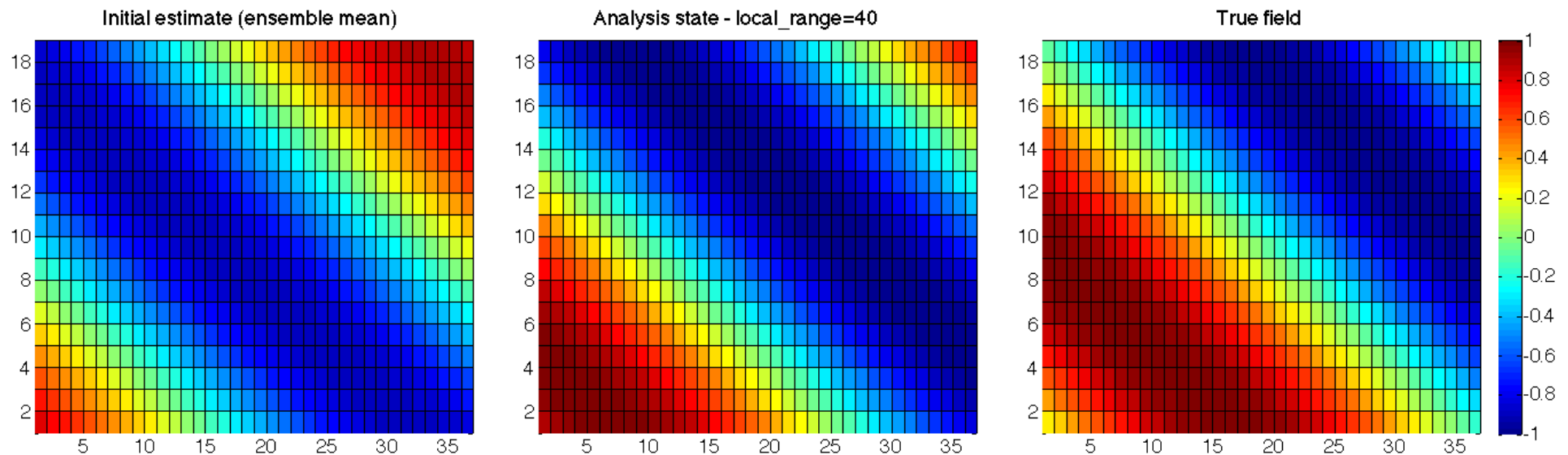
- Observation weighting by 5th-order polynomial
- Analysis field is smoother than before (because of weighting)



Result of the local assimilation (3)

```
./PDAF_offline -filtertype 7 -local_range 40.0
```

- Large radius: All local analysis domains see all observations
- Result identical to global filter



Local filter LESTKF

- Localized filters are a variant of the global filters
- User written files for global filter can be widely re-used
- Additional user-written files to handle local part

- No changes to:

```
initialize.F90
```

```
init_ens_offline.F90
```

```
prepoststep_ens_offline.F90
```

- Change in `init_pdaf_offline.F90`:

```
Set filtertype = 7
```

(You can also set it later on command line)

Local filter LESTKF (2)

Adapt files from global analysis

<code>init_dim_obs_pdaf.F90</code>	→	<code>init_dim_obs_f_pdaf.F90</code>
<code>obs_op_pdaf.F90</code>	→	<code>obs_op_f_pdaf.F90</code>
<code>init_obs_pdaf.F90</code>	→	<code>init_obs_f_pdaf.F90</code>
<code>prodrinva_pdaf.F90</code>	→	<code>prodrinva_l_pdaf</code>

Naming scheme:

`_f_` “full”: operate on all required observations
(without parallelization these are all observations)

`_l_` “local”: operation in local analysis domain or corresponding
local observation domain

Local filter LESTKF (3)

Additional files for local analysis step

`init_n_domains_pdaf.F90`

`init_dim_l_pdaf.F90`

`g2l_state_pdaf.F90`

`l2g_state_pdaf.F90`

`init_dim_obs_l_pdaf.F90`

`g2l_obs_pdaf.F90`

`init_obs_l_pdaf.F90`

localize
state vector

localize
observations

Discuss now the files in the order they are called

init_n_domains_pdaf.F90

Routine to set the number of local analysis domains

Output: `n_domains_p`

For the example: number of grid points (`nx * ny`)

To do:

1. Include `nx`, `ny` with `use mod_assimilation`
2. Set

```
n_domains_p = nx * ny
```

init_dim_obs_f_pdaf.F90

Initialize dimension of *full* observation vector

For the local filter:

1. Copy functionality from `init_dim_obs_pdaf.F90`
2. Rename `dim_obs_p` to **`dim_obs_f`** and `obs_p` to `obs_f`
3. Add storage of observation coordinates
 - a) Include `coords_obs_f` with `use mod_assimilation`
 - b) Where `obs_index_p` is allocated in the routine:
Allocate also `coords_obs_f(2,cnt)`
 - c) In the loop where `obs_index_p` is initialized add:

```
coords_obs_f(1,cnt)=REAL(j)
coords_obs_f(2,cnt)=REAL(i)
```

Note: We treat all coordinates as REAL variables even we use grid point indices here

obs_op_f_pdaf.F90

Implementation of observation operator
for full observation domain

1. Copy functionality from obs_op_pdaf.F90
2. Rename
 - `dim_obs_p` to `dim_obs_f`
 - `m_state_p` to `m_state_f`

Note:

The renaming is just for consistency. Quantities referring to the full observations should be recognizable by `_f`

init_obs_f_pdaf.F90

Fill PDAF's full observation vector

1. Copy functionality from `init_obs_pdaf.F90`
2. Rename
 - `dim_obs_p` to `dim_obs_f`
 - `observation_p` to `observation_f`

Note:

The renaming is just for consistency. Quantities referring to the full observations should be recognizable by `_f`

init_dim_1_pdaf.F90

Set the vector size `dim_1` of the local analysis domain

Further set the coordinates of the local analysis domain and the indices of the elements of the local state vector in the global state vector

Each single grid point is a local analysis domain in the example

1. Set `dim_1 = 1`

2. Compute the coordinates:

- Include `coords_1` with `use mod_assimilation`

```
coords_1(1) = REAL(CEILING(REAL(domain_p)/REAL(ny)))
```

```
coords_1(2) = REAL(domain_p) - (coords_1(1)-1)*REAL(ny)
```

Note: `coords_1` will be used later for computing the distance of observations from the local analysis domain in `init_dim_1_pdaf`

init_dim_1_pdaf.F90 (2)

3. Set indices of the elements of the local state vector in the global state vector

a) Include `id_lstate_in_pstate`
with `use mod_assimilation`

b) Allocate `id_lstate_in_pstate(dim_1)`
(Deallocate first if already allocated)

c) Specify the index: It's identical to `domain_p` here
(because we only have a single model variable)

```
id_lstate_in_pstate(1) = domain_p
```

init_dim_obs_l_pdaf.F90

Set the size of the observation vector for the local analysis domain

As for the global filter, this is the longest routine (~102 lines)

Only direct output: `dim_obs_l`

Operations:

1. Include coordinates `coords_l` with `use mod_assimilation`
2. Determine coordinate range for observations
3. Count observations within prescribed localization radius
4. Set index array for local observations (`id_lobs_in_fobs`) and array of distances of local observations (`distance_l`)

Note: The index array in step 4 is re-used for an efficient implementation of `g2l_obs_pdaf`. The local distance array initialized in step 4 is re-used in `prodrinva_l_pdaf` avoiding to recompute distances.

init_dim_obs_l_pdaf.F90 (2)

2. Determine coordinate range for local observations
 1. Declare `real :: limits_x(2), limits_y(2)`
 2. Include `local_range` with `use mod_assimilation`
 3. Set lower and upper limits. E.g. for x-direction

```
limits_x(1) = coords_l(1) - local_range
if (limits_x(1) < 1.0) limits_x(1) = 1.0
limits_x(2) = coords_l(1) + local_range
if (limits_x(2) > real(nx)) limits_x(2) = real(nx)
```

(analogous for y-direction)

Note: Using `limits_x`, `limits_y` is not strictly required, but it can make the search for local observations more efficient

If the localization is only based on grid point indices, the coordinates could be handled as integer values

init_dim_obs_l_pdaf.F90 (3)

3. Count local observations (within distance `local_range`)

```
dim_obs_l = 0
DO i = 1, dim_obs_f
  IF ("coords_obs(:,i) within coordinate limits") THEN
    Compute distance between coords_obs and coords_l
    IF (distance <= local_range) &
      dim_obs_l = dim_obs_l + 1
  END IF
END DO
```

Note:

For efficiency, we only compute `distance` for observations within coordinate limits `limits_x`, `limits_y`. Valid local observations reside within circle of radius `local_range` which is checked with `distance`.

init_dim_obs_l_pdaf.F90 (4)

4. Set index array for local observations

➤ Index of a local observation in the full observation vector

1. Include `id_obs_in_fobs` and `distance_l`
with `use mod_assimilation`

2. Allocate `id_obs_in_fobs(dim_obs_l)`

3. Fill index array:

```
cnt = 0
```

```
DO i = 1, dim_obs_f
```

```
  IF ("coords_obs(:,i) within coordinate limits") THEN
```

```
    Compute distance between coords_obs and coords_l
```

```
    IF (distance <= local_range) THEN
```

```
      cnt = cnt + 1
```

```
      id_obs_in_fobs(cnt) = i
```

```
      distance_l(cnt) = distance
```

```
  END ...
```

g2l_state_pdaf.F90 & l2g_state_pdaf.F90

g2l_state_pdaf: Initialize state vector for local analysis domain
from global state vector

l2g_state_pdaf: Initialize global state vector
from state vector for local analysis domain

- The templates provide a generic implementation
using the array `id_lstate_in_fstate`

→ We use the templates without any changes!

g2l_obs_pdaf.F90 & init_obs_l_pdaf.F90

g2l_obs_pdaf: Initialize local observed state vector from full observed vector

init_obs_l_pdaf: Initialize local vector of observations

- The templates provide a generic implementation using the array `id_lobs_in_fobs`

→ We use the templates with out any changes!

Note:

`init_obs_l_pdaf` requires that the full observation vector is stored in the array `obs_f`

prodrinva_l_pdaf.F90

Compute the product of the inverse observation error covariance matrix with some other matrix

+ apply observation localization (weighting)

➤ The weighting and the product are fully implemented for a diagonal observation error covariance matrix with constant variance

→ When we re-use the array `distance_l` initialized in `init_dim_obs_l_pdaf`, the template can be used without changes.

Done!

Now, the analysis step for local ESKTF in offline mode is fully implemented.

The implementation allows you now to use the local filters LESTKF, LETKF, and LSEIK

Not usable are EnKF and SEEK (PDAF does not have localization for these filters)

For testing one can vary localization parameters:

`local_range` – the localization radius

`locweight` – the weighting method

Default are `local_range=0.0` (observation at single grid point) and `locweight=1` (uniform weight)

A complete local analysis step

We now have a fully functional analysis step including localization

- It can be adapted to multiple model fields, 3 dimensions, different observations, etc.
- It can be used even with big models
 - if computing time is no concern
 - and if the computer has sufficient memory (e.g. ensemble array with dimension 10^7 and 20 members requires about 1.6 GB)
- Parallelization is required
 - if the problem is too big for a single process

1b.1) Use local filter OpenMP-parallelization

OpenMP

- OpenMP is so-called *shared-memory parallelization*
- Support for OpenMP is built into current compilers (needs to be activated by compiler-flag)
- Define OpenMP in the code by compiler directives: `!$OMP ...`
- Shared-memory parallelization:
 - Run several OpenMP “threads” (like processes in MPI)
 - All threads can access the same array in memory, but perform different operations
 - Typical is loop-parallelization: Each thread executes some part of a loop. For example, a fraction of a vector:

```
!$OMP parallel do
DO i = 1, 1000
    a(i) = b(i) + c(i)
ENDDO
```

With 2 threads, typically:

- thread 1 runs $i=1$ to 500
- thread 2 runs $i=501$ to 1000

OpenMP – what's relevant for PDAF

The local filters (LESTKF, LETKF, LSEIK, LNETF) are parallelized with OpenMP

- The loop over local analysis domains is distributed over threads

To make this work:

- Take into account, whether a variable is
 - *shared* (all threads see the same) or
 - *private* (each thread has its own copy)
- Variables referring to a local analysis domain (e.g. `coords_l`) have to be private
- This is ensured using the declaration 'THREADPRIVATE'

Running the tutorial program

Run analogously to case without parallelization

- `cd` to `/tutorial/classical/offline_2D_serial`
- Set environment variable `PDAF_ARCH` or set it in Makefile (e.g. `linux_gfortran`)
- Check and edit the make include file to activate OpenMP
 - for gfortran: `OPT = ... -fopenmp`
 - for Intel compiler: `OPT = ... -openmp`
- Compile by running `'make'`
- Set the number of OpenMP threads as environment variable, e.g.
 - for bash: `export OMP_NUM_THREADS=2`
 - for tcsh: `setenv OMP_NUM_THREADS 2`
- Run the program as [without OpenMP-parallelization](#)

Results from running with OpenMP parallelization

The results should be *identical* to those without parallelization

If the program is compiled with activated OpenMP-parallelization, you will see in the output of the analysis step the line

```
--- Use OpenMP parallelization with      2 threads
```

OpenMP in the local filter LESTKF

PDAF supports the use of OpenMP in the localized filters (LESTKF, LETKF, LSEIK, LNETF)

Settings to make OpenMP work are in two files:

```
prodrinva_l_pdaf.F90  
mod_assimilation.F90
```

The template files include the settings for OpenMP

prodrinva_l_pdaf.F90

Two variables have attribute 'save':

```
domain_save      mythread
```

Both variables are set private to the thread by

```
!$OMP THREADPRIVATE(mythread, domain_save)
```

(thus each OpenMP thread has a different value of the variables)

Both variables are used to ensure 'nice' screen output.

mod_assimilation.F90

Last line of `mod_assimilation.F90` is

```
!$OMP THREADPRIVATE(coords_l, id_lstate_in_pstate, id_lobs_in_fobs, ...  
                                                                distance_l)
```

- These variables are specific for each local analysis domain
- The variables are declared in `mod_assimilation.F90`
- The declaration 'THREADPRIVATE' ensures that each variable can have a different value in the different threads

2a) Parallelized global filter

Parallelize the analysis step

Implementation Strategy:

Take files from global analysis without parallelization and add the parallelization

Parallelization:

- Perform analysis step using multiple processors
- Split the state vector into equal parts to distribute the work

Notation for parallelization:

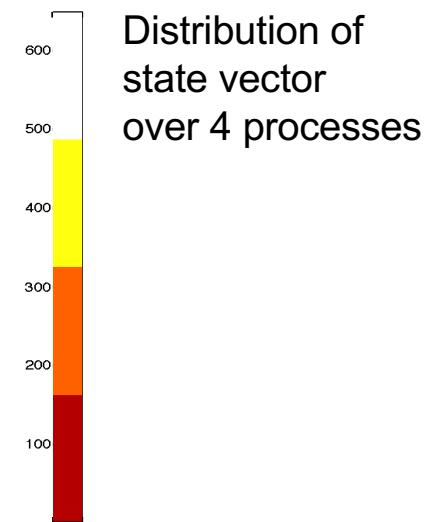
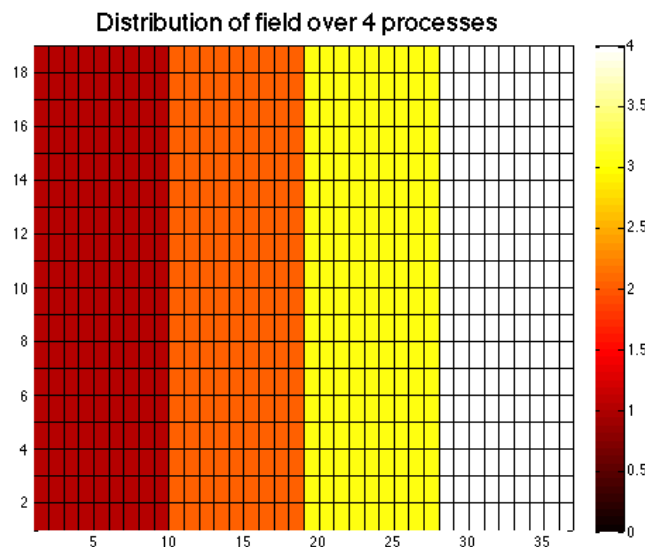
- Suffix `_p` marks variables with process-specific values
- Parallelization variables are declared in the module `mod_parallel`

Decomposition of model field

Want to distribute the state vector over the processes

- Split state vector into approximately equal continuous parts
- Corresponds to distribution along second index of model field (the first one in continuous in memory)

For 36 grid points we have uniform distributions for 2,3,4,6,or 9 processes (other numbers are possible)



Running the parallel tutorial program

- cd to /tutorial/classical/offline_2D_parallel
- Set environment variable PDAF_ARCH or set it in Makefile (e.g. linux_gfortran_openmpi)
- Clean existing files with 'make cleanall'
(This also removes the compiled PDAF library from previous tests)
- Compile by running 'make'
(this also builds the PDAF library again; now with parallelization)
- Run the program with

```
mpirun -np X ./PDAF_offline
```

($X > 0$; optimal are $x=1, 2, 3, 4, 6$ because then $ny=36$ is dividable by X)

Impact of the parallelization

- Ensemble array is distributed → less memory per process (visible in the memory display at the end of the screen output):

```
$ mpirun -np 1 ./PDAF_offline
```

PDAF Memory overview

```
-----  
Allocated memory (MB)  
state and A: 0.00543 MB (persistent)  
ensemble array: 0.04449 MB (persistent)  
analysis step: 0.02684 MB (temporary)
```

```
$ mpirun -np 4 ./PDAF_offline
```

```
Allocated memory (MB)  
state and A: 0.00172 MB (persistent)  
ensemble array: 0.01112 MB (persistent)  
analysis step: 0.01929 MB (temporary)
```

Impact of the parallelization (2)

Screen output shows some influence of the parallelization

```
Parallelization - Filter on model PEs:
```

```
    Total number of PEs:      4
```

```
    Number of parallel model tasks:  1
```

```
    PEs for Filter:          4
```

```
# PEs per ensemble task and local ensemble sizes:
```

```
    Task      1
```

```
    #PEs     4
```

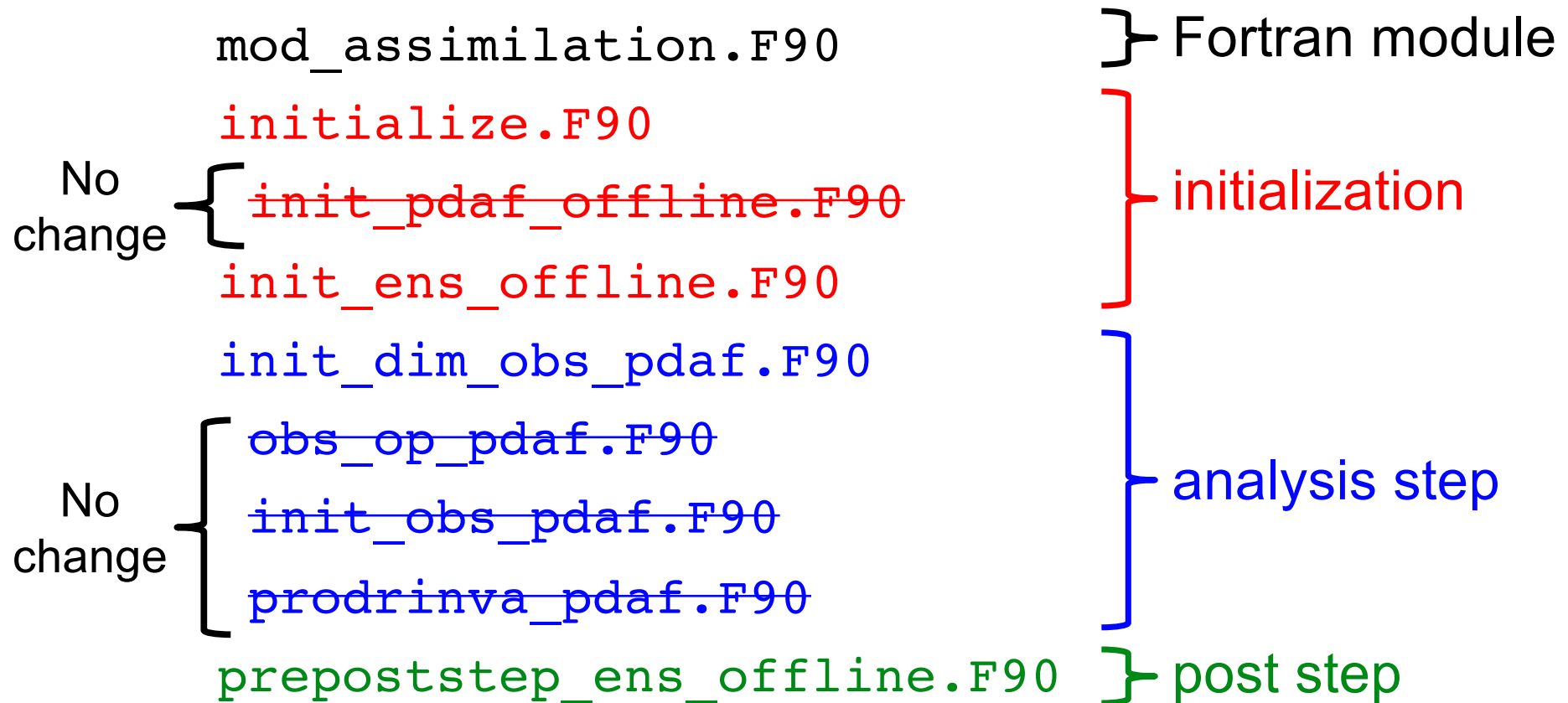
```
    N        9
```

At analysis step:

```
--- PE-domain 1 dimension of observation vector 8
--- PE-domain 2 dimension of observation vector 8
--- PE-domain 3 dimension of observation vector 8
--- PE-domain 4 dimension of observation vector 4
```

Note: The output lines might be unordered

Global ESTKF: Files to be changed for parallelization



initialize.F90 – parallelization

Initialize the model information – we have: `nx`, `ny`, `dim_state_p`

1. Use additional dimensions from `mod_assimilation`:

```
integer :: dim_state
integer, allocatable :: local_dims(:)
```

2. Rename `dim_state_p` to `dim_state` (global dimension)

3. Allocate `local_dims(npes_model)`

4. Set `dim_state_p` and `local_dims(:)`
– distribute `dim_state` over number of processes

```
local_dims = FLOOR(REAL(dim_state) / REAL(npes_model))
DO i = 1, (dim_state - npes_model * local_dims(1))
  local_dims(i) = local_dims(i) + 1
END DO
```

```
dim_state_p = local_dims(mype_model+1)
```

init_ens_offline.F90 – parallelization

Initialize ensemble matrix `ens_p`

Simple parallel variant:

1. Initialize global ensemble array (only one process)
 2. Distribute sub-states of ensemble array
(from the process doing step 1 to all others)
-
1. Required steps – only for `mype_filter==0`
 - Declare array `ens` and
allocate `ens(dim_state, dim_ens)`
 - Use serial implementation for initialize `ens`
(replace `ens_p` by `ens`)

init_ens_offline.F90 – parallelization (2)

2. Distribute sub-states of ensemble array

For `mype_filter=0`

a) Initialize local part of `ens_p` directly:

```
ens_p(1:dim_p,1:dim_ens) = ens(1:dim_p,1:dim_ens)
```

b) Distribute other sub ensembles

```
DO domain=2, npes_filter
```

```
allocate ens_p_tmp(local_dims(domain), dim_ens)
```

```
fill ens_p_tmp with part of ens for domain
```

```
MPI_Send ens_p_tmp from process 0 to process 'domain-1'
```

```
deallocate ens_p_tmp
```

init_ens_offline.F90 – parallelization (3)

2. Distribute sub-states of ensemble array

For all processes with `mype_filter>0`:

```
MPI_Recv ens_p_tmp into ens_p
```

Notes:

- “Classical” MPI communication: `MPI_Send/MPI_Recv`
- See tutorial code for MPI function calls
- Offset in state vector for `mype_filter=k` is
sum of `local_dims(i)` from `i=1` to `k`
- Size of state vector part is `local_dims(k)`
- The example code is not the most efficient possibility:
Each process could read its own local part of `ens_p`

init_dim_obs_pdaf.F90 – parallelization

Operations in case of parallelization:

- Read observation file
- Count number of observations for **process-local** part of state vector (**dim_obs_p**)
- Initialize array **obs_p** holding **process-local** available observations
- Initialize index array telling index of observation point in **process-local** state vector

Adapt serial implementation for these operations

init_dim_obs_pdaf.F90 – parallelization (2)

Count available process-local observations (**dim_obs_p**):

1. Get offset of local part in global state vector

`off_p` = Sum over local_dims(i) up to i=mype_filter

2. Now count

```
cnt = 0
cnt0 = 0
DO j = 1, nx
  DO i= 1, ny
    cnt0 = cnt0 + 1
    IF (cnt0>off_p .AND.
        cnt0<=off_p+local_dims(mype_filter+1)) THEN
      IF (obs_field(i,j) > -999.0) cnt = cnt + 1
    END IF; END DO; END DO
dim_obs_p = cnt
```

init_dim_obs_pdaf.F90 – parallelization (3)

Initialize obs_p and obs_index_p (now process-local parts)

```
cnt0 = cnt_p = cnt0_p = 0      ! Count grid points
DO j = 1, nx
  DO i= 1, ny
    cnt0 = cnt0 + 1
    IF (cnt0>off_p .AND. &
        cnt0<=off_p+local_dims(mype_filter+1)) THEN
      cnt0_p = cnt0_p + 1
      IF (obs_field(i,j) > -999.0) THEN
        cnt_p = cnt_p + 1
        obs_index_p(cnt_p) = cnt0_p      ! Index
        obs_p(cnt_p) = obs_field(i, j) ! observations
      END IF; END IF
    END DO
  END DO
END DO
```

prepoststep_ens_offline.F90 – parallelization

Post-step routine for the offline mode

Adapt writing of output files for parallelism

ensemble array `ens_p` is distributed

To do – inverse operations to `init_ens_offline`

- Use temporary array `ens_p_tmp`
- For `mype_filter>0`:
 - `MPI_Send ens_p` to `mype_filter=0`
- For `mype_filter=0`:
 - Do `domain=2, npes_filter`
 - `MPI_Recv` into `ens_p_tmp`
 - Initialize part of global array `ens` with `ens_p_tmp`
 - Write `ens` into files

prepoststep_ens_offline.F90 – parallelization (2)

Also in the tutorial implementation

- Collect local mean states (`state_p`) into a global analysis state and write to file.
- Collect vector of estimated variance (`variance_p`) into a global variance vector. Compute estimated RMS error from it.

Done!

The analysis step in offline mode with parallelization is fully implemented now

The implementation allows you now to use the global filters ESTKF, ETKF, and SEIK

- The parallel implementation can be compiled as before (without an MPI library) and run using 1 process
- To use the parallelization we have to compile with MPI library (see next slide)

2b) Parallelized local filter

Impact of the parallelization

- Ensemble array is distributed → less memory per process (visible in the memory display at the end of the screen output):

```
$ mpirun -np 1 ./PDAF_offline -filtertype 7
```

```
          Allocated memory  (MB)
state and A:  0.01038 MB (persistent)
ensemble array: 0.04449 MB (persistent)
analysis step: 0.01922 MB (temporary)
```

```
$ mpirun -np 4 ./PDAF_offline -filtertype 7
```

```
          Allocated memory  (MB)
state and A:  0.00296 MB (persistent)
ensemble array: 0.01112 MB (persistent)
analysis step: 0.01922 MB (temporary)
```

Note: Memory for analysis step is not changed!

Impact of the parallelization (2)

Screen output shows some influence of the parallelization

```
Parallelization - Filter on model PEs:  
Total number of PEs:      4
```

...

At analysis step:

```
--- PE-domain: 0 number of analysis domains: 162  
--- PE-domain: 1 number of analysis domains: 162  
--- PE-domain: 2 number of analysis domains: 162  
--- PE-domain: 3 number of analysis domains: 162  
  
--- PE-Domain: 2 dimension of PE-local full obs. vector 28  
--- PE-Domain: 3 dimension of PE-local full obs. vector 28  
--- PE-Domain: 0 dimension of PE-local full obs. vector 28  
--- PE-Domain: 1 dimension of PE-local full obs. vector 28
```

Note: The output lines might be unordered

Parallelize the local analysis step

Take files from global analysis without parallelization
and add the parallelization

Parallelization:

- Perform analysis step using multiple processors
- Split the state vector into equal parts to distribute the work

- Particular for localization:

Take care for local observation regions
(they can reach into state vector parts of other processes)

Notation for parallelization:

- Suffix `_p` marks variables with process-specific values

Local filter LESTKF – parallelization

Files to be parallelized

`init_n_domains_pdaf.F90`

`init_dim_obs_f_pdaf.F90`

`obs_op_f_pdaf.F90`

~~`init_obs_f_pdaf.F90`~~

~~`init_dim_l_pdaf.F90`~~

~~`init_dim_obs_l_pdaf.F90`~~

~~`g2l_state_pdaf.F90`~~

~~`g2l_obs_pdaf.F90`~~

~~`init_obs_l_pdaf.F90`~~

~~`prodrinva_l_pdaf`~~

~~`l2g_state_pdaf.F90`~~

No
Changes

Discuss now the files in the order they are called

Routines for initialization!

`initialize.F90`

`init_pdaf_offline.F90`

`init_ens_offline.F90`

These routines are identical for the global and local filters!
(Required changes are explained in the part about the parallelization of the global filter)

Only possible difference

- `filtertype = 7`

in `init_pdaf_offline.F90`

init_n_domains_pdaf.F90

Routine to set the number of local analysis domains

`n_domains_p`: now the number of local analysis domains for the particular process (according to part of state vector)

To do:

1. Include `local_dims` with use `mod_assimilation`

2. Set

```
n_domains_p = local_dims(mype_filter+1)
```

init_dim_obs_f_pdaf.F90 – parallelization

Operations in case of parallelization:

- Read observation file
- Count number of observations for **process-local** part of state vector (`dim_obs_p`)
- Initialize arrays holding **process-local** available observations (`obs_p`) and their coordinates (`obs_coords_p`)
- Initialize index array (`obs_index_p`) telling index of a **process-local** observation in **process-local** state vector
- Initialize **full** number of observations (`dim_obs_f`), vector of observations (`obs_f`), and coordinates (`coords_obs_f`)

“FULL” observation vector:

All observations required for all local analyses in process-local part of state vector (Here: *Full=All* observations for simplicity)

Adapt serial implementation ...

init_dim_obs_f_pdaf.F90 – parallelization (2)

Count process-local observations (dim_obs_p):

1. Include dim_obs_p with use mod_assimilation
2. Get offset off of local part in global state vector
(see global filter)
3. Now count

```
cnt0 = 0; cnt_p = 0
DO j = 1, nx; DO i= 1, ny
cnt0 = cnt0 + 1
  IF (cnt0>=off_p+1 .AND.
      cnt0<=off_p+local_dims(mytype_filter+1)) THEN
    IF (obs_field(i,j) > -999.0) cnt_p = cnt_p + 1
  END IF; END DO; END DO

dim_obs_p = cnt_p
```

init_dim_obs_f_pdaf.F90 – parallelization (3)

Initialize obs_p, obs_index_p, and coords_obs_p

1. Include obs_index_p, coords_obs_f and obs_f with use mod_assimilation
2. Add local arrays for obs_p(:) and coords_obs_p(:, :)
3. Adapt allocates to changed names and size dim_obs_p
4. In the loops rename the variables from _f to _p
5. Adapt the loop initializing the array by adding the check for the index range as for the counting loop

```
cnt0 = 0; cnt_p = 0; cnt0_p = 0
DO j = 1, nx; DO i= 1, ny
cnt0 = cnt0 + 1
  IF (cnt0>=off_p+1 .AND.
      cnt0<=off_p+local_dims(mype_filter+1)) THEN
    cnt0_p = cnt0_p + 1
    IF (obs_field(i,j) > -999.0) THEN ... END IF
  END IF; END DO; END DO
```

init_dim_obs_f_pdaf.F90 – parallelization (4)

Initialize full quantities (`dim_obs_f`, `obs_f`, `coords_obs_f`)

1. Obtain `dim_obs_f` by calling `PDAF_gather_dim_obs_f`
2. Allocate `obs_f` and `coords_obs_f`
(deallocate first if already allocated)
3. Obtain `obs_f` by calling `PDAF_gather_obs_f`
4. Obtain `coords_obs_f` by calling `PDAF_gather_obs_f`
5. Add `DEALLOCATE` for `obs_p` and `coords_obs_p`

Note: It is mandatory to call `PDAF_gather_dim_obs_f` once before using the two other functions because it stores dimension information.

Note: The three PDAF functions used here have been added with PDAF Version 1.13 to avoid that the user implementation needs calls to MPI functions.

Note: `coords_obs_f` has to be a REAL array

obs_op_f_pdaf.F90 – parallelization

Implementation of observation operator
for full observation domain

Difficulty:

- The state vector `state_p` is local to each process
- Full observed vector goes beyond process boundary

Adapt serial version:

1. Initialize process-local observed state
2. Get full observed state vector using `PDAF_gather_obs_f`

obs_op_f_pdaf.F90 – parallelization (2)

1. Initialize process-local observed state `m_state_p`

a) Include `dim_obs_p` and `obs_index_p`
with `use mod_assimilation`

b) Declare real allocatable array `m_state_p(:)`

c) Allocate
`m_state_p(dim_obs_p)`

d) Fill the array

```
DO i = 1, dim_obs_p
    m_state_p(i) = state_p(obs_index_p(i))
END DO
```

Note:

In the serial version the upper bound of the loop was `dim_obs_f`
and we filled `m_state_f` directly

obs_op_f_pdaf.F90 – parallelization (3)

2. Get full observed state vector

a) Add variable `INTEGER :: status`

b) Add call to `PDAF_gather_obs_f`:

```
CALL PDAF_gather_obs_f(m_state_p, m_state_f, status)
```

c) Deallocate `m_state_p`

Note: It is mandatory to call `PDAF_gather_dim_obs_f` once before using the two other functions because it stores dimension information. Usually this was already done in `init_dim_obs_f_pdaf`

Done!

Now, the analysis step for local ESKTF with parallelization in offline mode is fully implemented.

The implementation allows you now to use the local filters LESTKF, LETKF, and LSEIK

Not usable are EnKF and SEEK (PDAF doesn't have localization for SEEK and a different localization scheme for EnKF)

3) Hints for adaptations for real models

Implementations for real models

- Tutorial demonstrates implementation for simple model
- You can base your own implementation on the tutorial implementation or the templates provided with PDAF
- Need to adapt most routines, e.g.
 - Specify model-specific state vector and its dimension
 - Adapt routines handling observations
- Adapt file output:
 - need to read and write restart files from specific model
 - adapt writing of ensemble mean state in `prepoststep_pdaf`

Multiple fields in state vector

- Tutorial uses a single 2-dimensional field
- All fields that should be updated by the assimilation have to be part of the state vector
- For more fields:
 - concatenate them in the state vector
 - adapt state dimension in `init_pdaf`
 - adapt `init_ens_pdaf`, `collect_state_pdaf`, `distribute_state_pdaf`, `prepoststep_pdaf`
 - For local filters: Adapt full (`_f_`) and local (`_l_`) routines and `g2l_state_pdaf`, `l2g_state_pdaf`, `g2l_obs_pdaf`
- **Note**
 - It can be useful to define a vector storing the offset (position) of each field in the state vector

Multiple observed fields

- In tutorial: observed one field at some grid points
- For several observed fields adapt observation routines:
 - concatenate observed fields in observation vector
 - adapt all observation-handling routines
- **Note**
 - The observation errors can be set differently for each observed field (e.g. using an array `rms_obs`)
 - The localization radius can be set specific for each observed field (observation search in `init_dim_obs_1_pdaf` would use different `local_range` for different fields)
 - One can use spatially varying observation errors using an array `rms_obs` in `prodrinva(_1)_pdaf`

The End!

Tutorial described example implementations

- Offline mode of PDAF
- Simple 2D example
- Square root filter ESTKF
 - global and with localization
 - without and with parallelization
- Extension to more realistic cases possible with limited coding
- Applicable also for large-scale problems

For full documentation of PDAF
and the user-implemented routines
see <http://pdaf.awi.de>