Data Assimilation

DART/CAM-Chem

Summary
The CAM-Chem model has been interfaced to the Data Assimilation Research Testbed (DART) in order to assimilate both chemical and meteorological observations.

In this case, the data assimilation scheme is an Ensemble Adjustment Kalman Filter (EAKF).

This requires an ensemble (or multiple instances) of CAM-Chem simulations, which is currently done within the multi-instance feature of the CESM coupler.

The DART documentation is available online and the code is on GitHub, below are some specifics needed for chemistry.

Adding a variable
If you want to add a variable to the state vector for evaluation or assimilation. You have to define a DART quantity (QTY Xxx).

The interface between the model and DART for chemistry variables is defined in chem_table_mod.f90.

For instance, if you want to add the ozone variable ('O3' in CAM-Chem), you will need to add this to the chem_table_mod.f90.

```fortran
call add_entry('O3', 47.9982_r8, 'QTY_O3')
```

The molecular weight (47.9982_r8) is used to convert the mass mixing ratio (in CAM initial files) into volume mixing ratio.

The created quantity needs to be defined in model_mod.f90 by adding:

```fortran
obs_kind_mod, only: use QTY_O3
```

The quantity (QTY_O3) also needs to be defined in ../assimilation_code/modules/observations/DEFAULT_obs_kind_mod.F90.

The procedure to add a quantity in obs_kind_mod is explained in the DART documentation of the preprocess program.

Other resources

Datasets available:

A reanalysis of MOPITT Carbon Monoxide (2002-2013) has been performed (Gaubert et al., 2016; Gaubert et al., 2017) and monthly means outputs are available here.

References:

DART-CAM-Chem (CAM6)
DART-CAM-Chem (CAM5)
DART-CAM (CAM4)
DART-CAM-Chem (CAM3)

Contact: Benjamin Gaubert