

# Metos3D

## model

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## 1 Model interface

Metos3D can be coupled to every (biogeochemical) model that conforms to the following interface:

```
subroutine metos3dbgc(n, ny, m, nb, nd, dt, q, t, y, u, b, d)
  integer :: n          ! tracer count
  integer :: ny         ! layer count
  integer :: m          ! parameter count
  integer :: nb         ! boundary condition count
  integer :: nd         ! domain condition count
  real*8  :: dt         ! ocean time step
  real*8  :: q(nz, n)   ! bgc model output
  real*8  :: t          ! point in time
  real*8  :: y(nz, n)   ! bgc model input
  real*8  :: u(m)        ! parameters
  real*8  :: b(nb)      ! boundary conditions
  real*8  :: d(nz, nd)  ! domain conditions
end subroutine
```

The interface decouples biogeochemical models and driver routines (ocean circulation, forcing, geometry) programmatically. It gives you the possibility to provide a free number of tracers, parameters, boundary and domain conditions. It suits well an optimization as well as an Automatic Differentiation (AD) context.

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## 2 BGC Models

Generally, for every model implementation that is coupled to the transport driver via the interface a new executable must be compiled. Metos3D provides an automatic compile scheme along with a convention for the model directory structure. Within the `model` directory of the `model` repository a folder with the model name must be created. Regarding the two models presented here, the directory is named `I-Cs` respectively `MITgcm-P04-D0P`. Within such a model directory a source file named `model.F` is stored. Overall, whereas here the file suffix implies a pre-processed Fortran fixed format, every programming language that is supported by the PETSc library will be accepted.

Finally, to compile all sources invoke

```
$> metos3d simpack MITgcm-P04-D0P
```

to create an executable named

```
metos3d-simpack-MITgcm-P04-D0P.exe
```

respectively

```
$> metos3d simpack I-Cs
```

to create

```
metos3d-simpack-I-Cs.exe
```

Specific settings are provided via option files. Exemplary, each of the two presented model archives contain an `option` directory. You can find a test option file therein. Use it as a starting point for your own work. To start a test run just type:

```
$> ./metos3d-simpack-I-Cs.exe model/I-Cs/option/test.I-Cs.option.txt
```

respectively

```
$> ./metos3d-simpack-MITgcm-P04-D0P.exe model/MITgcm-P04-D0P/option/test.MITgcm-P04-D0P.option.txt
```

### 2.1 I-Cs

The Iodine (I) and Caesium (Cs) model was implemented to predict the Caesium distribution after the Fukushima accident.

#### 2.1.1 Equations

The model equations describe the radioactive decay of the  $I^{131}$  and  $Cs^{137}$  isotopes named  $y_1$  and  $y_2$ , respectively. The decay depends on the half-life. The tracers do not react with each other. The equations read:

$$\begin{aligned}q_1(y_1, y_2) &= \log(0.5) 360.0/8.02070 y_1 \\q_2(y_1, y_2) &= \log(0.5) 1.0/30.17 y_2\end{aligned}$$

## 2.2 MITgcm-PO4-DOP

The MITgcm-PO4-DOP model is an *original* implementation of a biogeochemical model that is used for the MIT General Circulation Model [cf. 2, MITgcm] biogeochemistry tutorial and described in detail in [1]. The model comprises five biogeochemical variables, namely dissolved inorganic carbon (DIC), alkalinity (ALK), phosphate (PO4), dissolved organic phosphorous (DOP) and oxygen (O2). In fact, here, for verification just PO4 and DOP are used.

## References

- [1] Stephanie Dutkiewicz, Andrei P. Sokolov, Jeffery Scott, and Peter H. Stone. A three-dimensional ocean-seaice-carbon cycle model and its coupling to a two-dimensional atmospheric model: Uses in climate change studies. Technical Report 122, MIT Joint Program on the Science and Policy of Global Change, 2005.
- [2] J. Marshall, A. Adcroft, C. Hill, L. Perelman, and C. Heisey. A finite-volume, incompressible navier stokes model for studies of the ocean on parallel computers. *Journal of Geophysical Research*, 102:5753–5766, 1997.