



OASIS3-MCT v6 User Guide
TR-CMGC-25-161

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Chapter 1

Introduction

In 1991, CERFACS started the development of a software interface to couple existing ocean and atmosphere numerical General Circulation Models. Today, different versions of the OASIS3-MCT coupler are used by at least 65 modelling groups all around the world to couple more than 80 applications on different computing platforms¹. OASIS3-MCT sustained development is ensured by a collaboration between CERFACS and the Centre National de la Recherche Scientifique (CNRS) and its maintenance and user support is regularly reinforced with additional resources coming from European and national projects.

OASIS3-MCT is a portable set of Fortran 77, Fortran 90 and C routines. Low-intrusiveness, portability and flexibility are OASIS3-MCT key design concepts. After compilation OASIS3-MCT is a coupling library to be linked to the component models, and which main function is to interpolate and exchange the coupling fields between the components to form a coupled system. OASIS3-MCT supports coupling of 2D logically-rectangular fields but 3D fields and 1D fields expressed on unstructured grids are also supported using a one dimension degeneration of the structures.

Internally, OASIS3-MCT uses MCT, the Model Coupling Toolkit² [Larson et al 2005] [Jacob et al 2005], developed by the Argonne National Laboratory in the USA. MCT implements fully parallel remapping, as a parallel matrix vector multiplication on the set of source or target component processes, and parallel distributed exchanges of the coupling fields via Message Passing Interface (MPI). Its design philosophy, based on flexibility and minimal invasiveness, is close to the OASIS3-MCT approach.

To communicate with another component, a component model needs to include few specific calls to OASIS3-MCT Fortran Application Programming Interface (API). OASIS3-MCT also has python and C language bindings. For remapping the coupling fields from the source to the target grid, OASIS3-MCT v6 includes a modified version of the SCRIP library [Jones 1999] and is interfaced with YAC [Hanke et al 2016] [Hanke and Redler 2019] [Jonville et al 2024]. OASIS3-MCT also supports basic non-parallel file I/O using NetCDF. The configuration of the coupling exchanges needs to be described externally by the user in a text file, the *namcouple*.

The developments realised between the different versions of OASIS3-MCT are described in Appendix B. We believe that OASIS3-MCT will still provide an efficient and easy-to-use coupling solution for many climate modelling groups in the few years to come.

1.1 Step-by-step use of OASIS3-MCT

To use OASIS3-MCT for coupling codes, one has to follow these steps:

1. Obtain OASIS3-MCT source code (see chapter 1.2).

¹A list of coupled models realised with OASIS3-MCT can be found at <https://oasis.cerfacs.fr/en/oasis-coupled-models/>

²<https://web.cels.anl.gov/projects/climate/mct/>

2. Get familiar with OASIS3-MCT, either by following the Short Private Online Course (SPOC, see <https://cerfacs.fr/en/online-training-sessions/>) or by going through the tutorial steps. A tutorial is also available in directory `examples/tutorial_communication` and all explanations are provided in the document `tutorial_communication.pdf` therein.
3. Identify the coupling or I/O fields and adapt the codes to implement the coupling exchanges with the OASIS3-MCT coupling library based on MPI message passing. The OASIS3-MCT coupling library uses NetCDF and therefore can also be used to perform I/O actions from/to disk files. For more detail on how to use the OASIS3-MCT API in the codes, see chapter 2.
4. Define all coupling and I/O parameters and transformations required to adapt each coupling field from the source model grid to the target model grid. On this basis, prepare OASIS3-MCT configuring file `namcouple`. OASIS3-MCT supports different interpolation algorithms as described in chapter 4. Remapping files can be computed online using the SCRIP or YAC library, or offline using either the SCRIP, YAC, ESMF or XIOS (see `examples/regrid_environment`, section 6.3.3) and read in during the run using the MAPPING transformation.

We strongly recommend to tests the quality of the chosen transformations and remappings offline using the environment available in `examples/regrid_environment` and explanations provided in the document `regrid_environment.pdf` therein.

5. Generate required auxiliary data files , if needed (see chapter 5).
6. Compile OASIS3-MCT, the component models and start the coupled experiment. For details on how to compile and run a coupled model with OASIS3-MCT, see section 6.

If you need extra help, do not hesitate to contact us at [oasishelp\(at\)cerfacs.fr](mailto:oasishelp@cerfacs.fr) .

1.2 OASIS3-MCT sources

OASIS3-MCT sources are available from CERFACS git server. To obtain more detail on downloading the sources, please fill in the registration form at <https://oasis.cerfacs.fr/en/downloads/> .

OASIS3-MCT directory structure is the following one:

- `oasis3-mct/lib/cbindings` C language bindings source code
 - `/mct` Model Coupling Toolkit Coupling Software
 - `/psmile` OASIS3-MCT coupling library
 - `/scrip` SCRIP remapping library
 - `/yac` YAC remapping library
 - `/yaxt` communication layer used by YAC
- `oasis3-mct/doc` OASIS3-MCT User Guide
- `oasis3-mct/util/make_dir` Utilities to compile OASIS3-MCT
 - `/load_balancing` Tool for load balancing analysis
 - `/env_tests` Script to compile and launch toy models
- `oasis3-mct/pyoasis` Python wrapper source code
- `oasis3-mct/examples` Environment to compile, run and use different toy coupled models.

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Chapter 2

Interfacing a component code with OASIS3-MCT

At run-time, OASIS3-MCT performs parallel exchange of coupling data between parallel components and sub-components and allows regridding (also called remapping or interpolation), time integration or accumulation and other transformations of these coupling fields.

This chapter describes how to adapt the component codes to couple them through OASIS3-MCT.

OASIS3-MCT supports coupling exchanges between parallel components and sub-components deployed in diverse configurations. The different possibilities and how to use the OASIS3-MCT library accordingly are described in section 2.1.

The OASIS3-MCT Application Programming Interface (API) includes different classes of modules or routines that are described in detail in sections 2.2, 2.3 and 2.4 for Fortran, C and python codes respectively. In section 2.5, the reader will also find an overview of the MCT library (see 2.5.4) and additional notes on how to exchange scalars (see 2.5.1), how to reproduce different coupling algorithms with OASIS3-MCT using the `LAG` index (see 2.5.2), and on the `SEQ` index (see 2.5.3).

2.1 Configurations of components supported

Since OASIS3-MCT_3.0 release, coupling exchanges between components and sub-components can be deployed in a much larger diversity of configurations than before. This is illustrated on figure 2.1 and how to use the OASIS3-MCT library accordingly is detailed on figure 2.2. All OASIS3-MCT API routines are also described in details in sections 2.2, 2.3 and 2.4.

We call here an “executable” a compiled code forming a part of or the whole coupled system. A “component” is the ensemble of processes, or tasks, within the coupled system calling `oasis_init_comp` with the same `comp_name` argument (see section 2.2.2). A “sub-component” is the subset of tasks within a component sending or receiving coupling fields on a specific grid. Of course, a component may have only one sub-component that gathers all its tasks.

In `examples/tutorial_communication`, `examples/regrid_environment` and `pyoasis/examples`, one finds practical examples of how to use the OASIS3-MCT library (see sections 6.3.1 and 6.3.3).

With OASIS3-MCT, it is currently possible to implement coupling exchanges (in the following paragraph, the text between [and] refers to figure 2.1) :

- between two components or sub-components running concurrently on separate sets of tasks within two different executables [A, D, E, J];
- between two components or sub-components running concurrently on separate sets of tasks within one same executable [B, F, I];

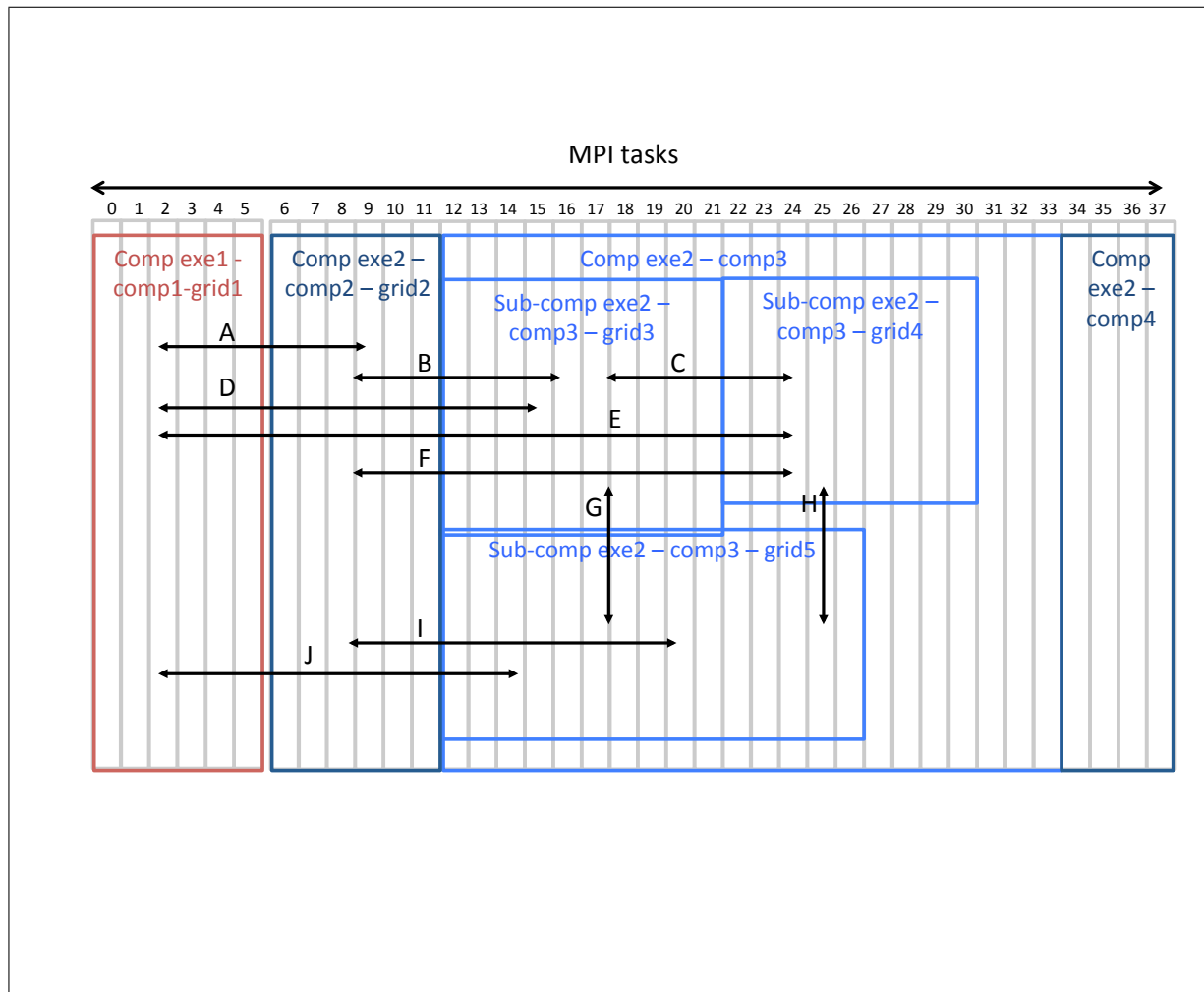


Figure 2.1: The different configuration of components supported by OASIS3-MCT. Two executables exe1 and exe2 are running concurrently on separate sets of MPI tasks (0-5 for exe1 and 6-37 for exe2). Executable exe1 includes only one component comp1 that has coupling fields defined on only one grid grid1 (decomposed on all of its 6 tasks). Executable exe2 includes 3 components, comp2, comp3, and comp4 running concurrently respectively on tasks 6-11, 12-33 and 34-37. Component comp2 participates in the coupling with fields defined on only one coupling grid grid2 (decomposed on all of its 5 tasks) while comp4 does not participate at all in the coupling. Component comp3 has 3 sub-components, respectively exchanging coupling fields defined on grid3 (tasks 12-21), grid4 (tasks 22-30) and grid5 (tasks 12-26, therefore overlapping with both grid3 and grid4); finally, comp3 has 3 tasks (31-33) not involved in the coupling. Sub-components exe2-comp3-grid3 and exe2-comp3-grid5, or sub-components exe2-comp3-grid4 and exe2-comp3-grid5 are examples of coupling between sub-components running sequentially on overlapping sets of tasks.

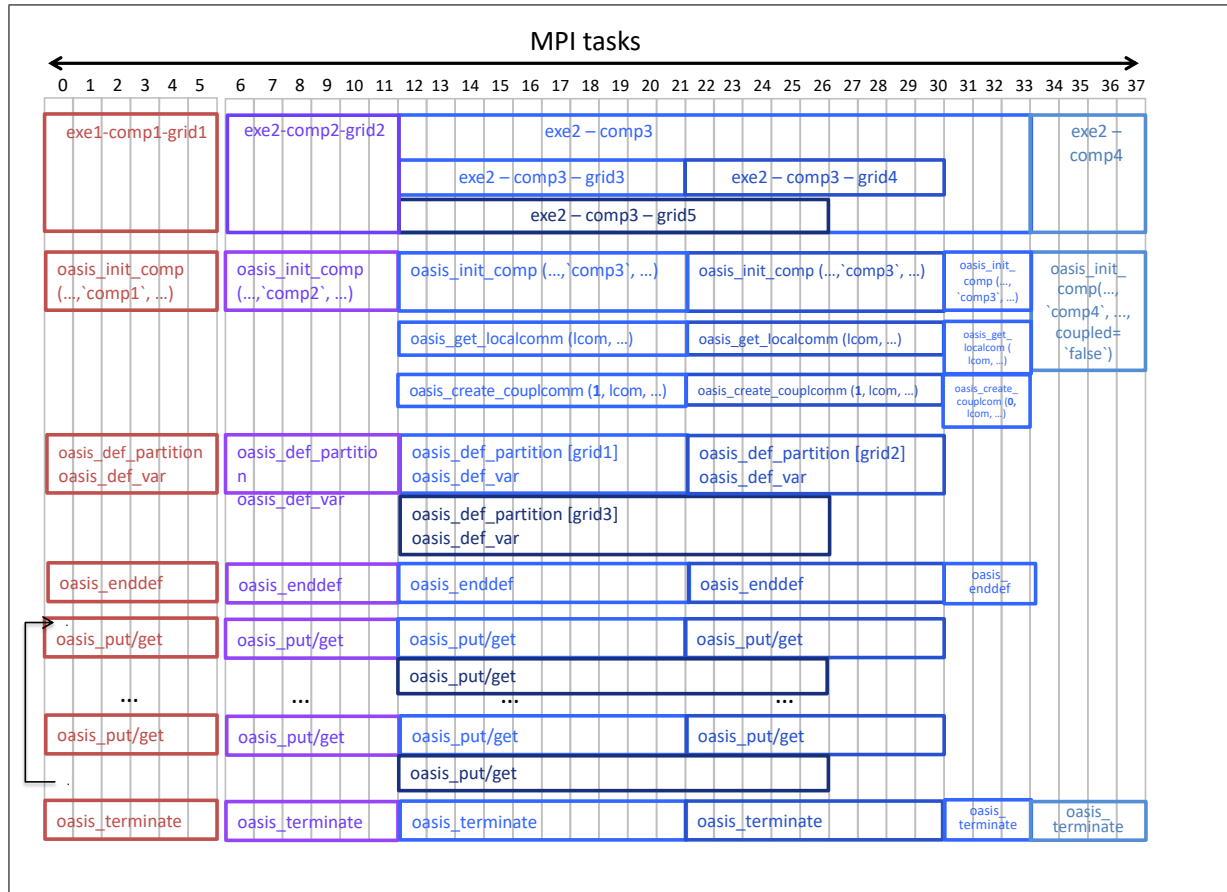


Figure 2.2: The sequence of OASIS3-MCT calls that have to be implemented in the codes so to allow the configuration of components described on figure 2.1. Each MPI tasks has to call `oasis_init_comp` once with the name of its component as 2^{nd} argument and `oasis_terminate` before exiting. As none of `comp4` tasks is participating to the coupling, `comp4` tasks calls `oasis_init_comp` with `coupled=.false.` as 4^{th} argument and does not call any other OASIS3-MCT routine except `oasis_terminate`. As some of `comp3` tasks are participating in the coupling, all `comp3` tasks have to call `oasis_init_comp`, `oasis_get_localcomm`, `oasis_create_couplcomm`, `oasis_undef` and `oasis_terminate` (these are the only routine to be called by `comp3` tasks 31-33 not participating to the coupling). To initialise the coupling exchanges, the tasks of a sub-component holding a field decomposed on a specific grid have to call the `oasis_def_partition` to express the decomposition of the grid, `oasis_def_var` to declare the coupling field and `oasis_undef`. Finally, the tasks of a sub-component exchanging coupling fields have to call `oasis_put` and `oasis_get` accordingly.

- within one executable between two concurrent sub-components [C]
- within one executable between two sub-components running sequentially on overlapping sets of tasks (i.e. a task can be coupled to itself calling both the “put” and the “get” of the exchange) [G, H]
- or to have some tasks of a component not participating to the coupling exchanges [tasks 31-33 of `exe2-comp3`]
- or to have all processes of a component not participating to the coupling exchanges [`exe2-comp4`, tasks 34-37]

The sequence of OASIS3-MCT API routines that have to be called in the different cases is detailed on figure 2.2. These routines are also described in detail in the next section.

2.2 OASIS3-MCT Fortran API

To interact with the rest of the coupled system, few calls of the OASIS3-MCT library routines, which sources can be found in `oasis3-mct/lib/psmile` directory, have to be implemented in the Fortran codes of the components. The interface for C or C++ codes and for python codes are described in section 2.3 and section 2.4 respectively. They belong to the following classes:

1. Module to use (section 2.2.1)
2. Initialisation (section 2.2.2)
3. Partition definition (section 2.2.3)
4. Grid data file definition (section 2.2.4)
5. Coupling-I/O field declaration (section 2.2.5)
6. End of definition phase (section 2.2.6)
7. Coupling-I/O field sending and receiving (section 2.2.7)
8. Termination (section 2.2.8)
9. Optional auxiliary routines (section 2.2.9)

2.2.1 Module to use in the code

To use OASIS3-MCT library, a user needs to add in his code:

- `USE mod_oasis`

Use of module `mod_oasis` provides access to all methods required that will continue to evolve in the future, always ensuring backward compatibility. In the following sections, the interface names are defined and the description of the interface arguments is provided.

2.2.2 Initialisation

Coupling initialisation

- `CALL oasis_init_comp (compid, comp_name, kinfo, coupled, commworld)`
 - `compid` [INTEGER; OUT]: returned component ID
 - `comp_name` [CHARACTER; IN]: component name; maximum length of 80 characters
 - `kinfo` [INTEGER; OUT]: returned error code
 - `coupled` [LOGICAL, OPTIONAL; IN]: flag to specify if the calling task is participating or not to the coupling (`.true.` by default).

- `commworld` [INTEGER, OPTIONAL; IN] : optional argument to specify the global communicator gathering the components of the coupled model. If not specified, `MPI_COMM_WORLD` will be used as the default communicator to startup. All components of the coupled model must specify the same `commworld` argument.

This routine must be called by all tasks of all components whether or not they are involved in the coupling. This routine calls `MPI_Init`. The component may also call `MPI_Init` explicitly, but if so, has to call it before calling `oasis_init_comp`; in this case, the component also has to call `MPI_Finalize` explicitly, but only after calling `oasis_terminate`.

A component is defined as the ensemble of tasks calling `oasis_init_comp` with the same `comp_name` argument. If, and only if, all tasks of a component are excluded from the coupling, the logical `coupled` can be set to `.false.` for the tasks of this component; in this case, `oasis_init_comp` and `oasis_terminate` are the only API routine that need to be called by the component tasks (see component `exe2-comp4` on figure 2.2). If at least one tasks of a component is participating to the coupling, all component tasks have to call `oasis_init_comp` with `coupled=.true.` (which is the default); in this case, the component tasks not participating to the coupling will also have to call `oasis_enddef` and `oasis_terminate`, and possibly `oasis_get_localcomm` or `oasis_create_couplcomm` if these routines are called by the other tasks of the component (see tasks 31-33 of `exe2-comp3` on figure 2.2).

Communicator for internal parallelisation

- CALL `oasis_get_localcomm (local_comm, kinfo)`
 - `local_comm` [INTEGER; OUT]: value of local communicator
 - `kinfo` [INTEGER; OUT]: returned error code.

This routine returns the value of a local communicator gathering only the tasks of the component (i.e. the tasks that called `oasis_init_comp` with the same `comp_name` argument).

This may be needed as all executables of the coupled system are usually started in a pseudo-MPMD mode with MPI1 and therefore share automatically the same `MPI_COMM_WORLD` communicator. Another communicator has therefore to be used for the internal parallelisation of each component. OASIS3-MCT creates this local communicator `local_comm` based on the value of the `comp_name` argument in the `oasis_init_comp` call.

Retrieving a local communicator `local_comm` is also needed if `oasis_create_couplcomm` is called, as `local_comm` is an argument of this routine (see below).

- CALL `oasis_create_couplcomm(icpl, local_comm, coupl_comm, kinfo)`
 - `icpl` [INTEGER; IN]: coupling process flag
 - `local_comm` [INTEGER; IN]: MPI communicator with all processes of the component
 - `coupl_comm` [INTEGER; OUT]: returned MPI communicator gathering only component processes participating in the coupling
 - `kinfo` [INTEGER; OUT; OPTIONAL]: returned error code

It is mandatory to call this routine that creates a coupling communicator for a subset of processes if only a subset of the component processes participate in the coupling (e.g. `exe2-comp3` in figure 2.2). In that case, the processes involved in the coupling have to call it with `icpl=1` while the other have to call it with `icpl = MPI_UNDEFINED`. Argument `local_comm` is the MPI communicator associated with all processes of the component returned by `oasis_get_localcomm`. The new coupling communicator is returned in `coupl_comm`.

If this communicator already exists in the code, the component should simply provide it to OASIS3-MCT with:

- CALL `oasis_set_couplcomm(coupl_comm, kinfo)`
 - `coupl_comm` [INTEGER; IN]: MPI communicator gathering only component processes participating in the coupling
 - `kinfo` [INTEGER; OUT; OPTIONAL]: returned error code

This routine allows users to provide a local coupling communicator to OASIS3-MCT, given that it already exists in the code. The value of `coupl_comm` must be the value of this local coupling communicator for the processes participating to the coupling and it must be `MPI_COMM_NULL` for processes not involved in the coupling.

2.2.3 Partition definition

The coupling fields sent or received by a component are usually scattered among the different component processes. With OASIS3-MCT, all processes exchanging coupling data have to describe, in a global index space spanning the full domain, the local partitioning of the different grids onto which the data is expressed (see 2.2.4 for the grid definition).

The choices detailed in the following provide a compact description of 2D partitions. In addition, a special case was introduced in OASIS3-MCT v6 to facilitate the description of 3D partitions. This does not mean that calculation of 3D interpolation weights are supported by OASIS3-MCT. In particular, the `cube` partition (see below) requires pre-computed 3D interpolation weights and forbids any other pre- or post-processing transformation that supposes a 2D structure of the exchanged data.

The processes not implied in the coupling do not have to call this routine (for backward compatibility with OASIS3-MCT_2.0, they may still call it describing a null partition, i.e. with `ig_parallel(:)=0`).

The partitions for the different processes cannot overlap, i.e. haloes are not supported and it is not possible to define a partition across multiple active OpenMP threads.¹

- CALL `oasis_def_partition(il_part_id, ig_parallel, kinfo, ig_size, name)`
 - `il_part_id` [INTEGER; OUT]: partition ID
 - `ig_parallel` [INTEGER, DIMENSION(:), IN]: vector of integers describing the local grid partition in the global index space spanning the 2D domain; has a different expression depending on the type of the partition (Serial -no partition, Apple, Box, Orange, Cube and Points, see below).
 - `kinfo` [INTEGER; OUT]: returned error code.
 - `ig_size` [INTEGER, OPTIONAL, IN]: Optional argument; mandatory if the coupling data is exchanged for only a subdomain of the global grid; in this case, `ig_size` must give the total number of grid points.
 - `name` [CHARACTER, OPTIONAL, IN]: Optional argument associating a name to the partition, mandatory if `oasis_def_partition` is called either for a grid decomposed not across all the processes of a component or if the related `oasis_def_partition` are not called in the same order on the different component processes; this argument is new since OASIS3-MCT_3.0 release and is linked to the greater flexibility in the configuration of components supported (see 2.1); it has a maximum length of 120 characters.

Serial (no partition)

This is the choice for a grid entirely supported by only one process. In this case, we have `ig_parallel(1:3):`

- `ig_parallel(1) = 0` (indicates a Serial partition, i.e. no partition)

¹Note also that with releases earlier than OASIS3-MCT_5.2 (2022/11/02), it was not allowed to associate more than one grid to one partition (this could cause problems with the global CONSERV operation.)

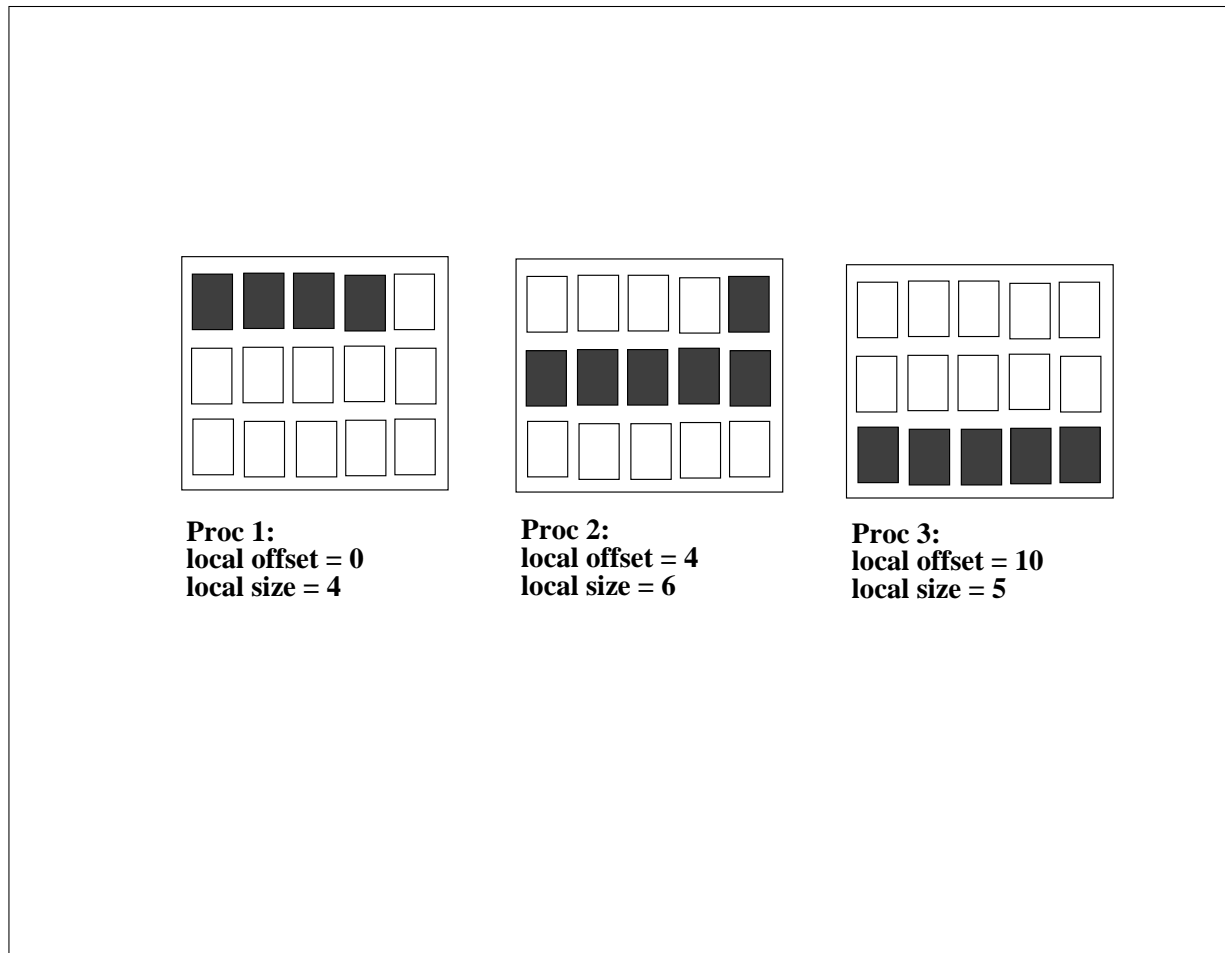


Figure 2.3: Apple partition. We assume here that the global index starts at 0 in the upper left corner.

- `ig_paral(2) = 0`
- `ig_paral(3) =` the total grid size.

Apple partition

Each partition is a segment of the global domain, described by its global offset and its local size. In this case, we have `ig_paral(1:3)`:

- `ig_paral(1) = 1` (indicates an Apple partition)
- `ig_paral(2) =` the segment global offset
- `ig_paral(3) =` the segment local size

Figure 2.3 illustrates an Apple partition over 3 processes.

Box partition

Each partition is a rectangular region of the global domain, described by the global offset of its upper left corner, and its local extents in the X and Y dimensions. The global extent in the X dimension must also be given. In this case, we have `ig_paral(1:5)`:

- `ig_paral(1) = 2` (indicates a Box partition)
- `ig_paral(2) =` the upper left corner global offset
- `ig_paral(3) =` the local extent in x
- `ig_paral(4) =` the local extent in y

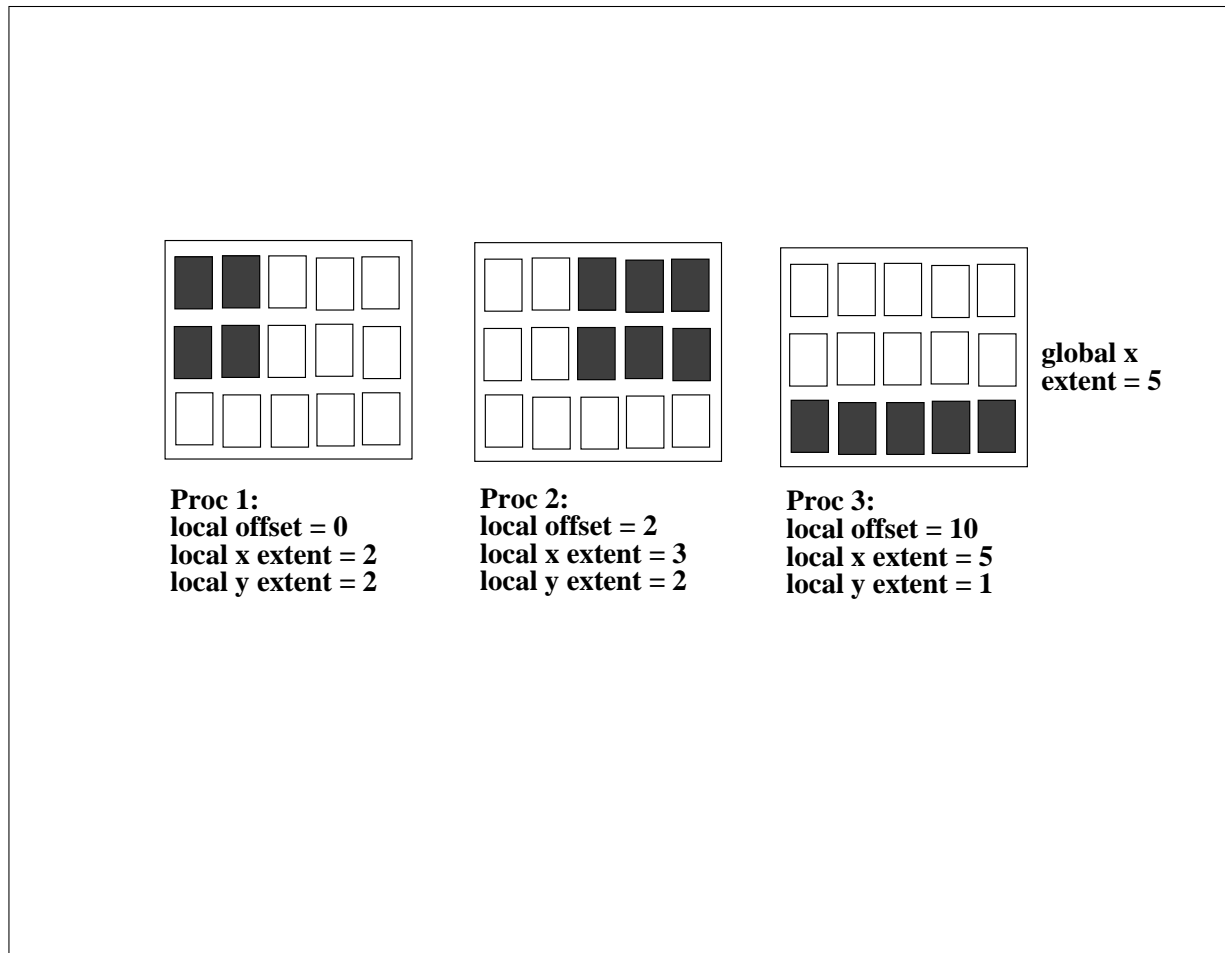


Figure 2.4: Box partition. We assume here that the global index starts at 0 in the upper left corner.

- `ig_parallel(5)` = the global extent in x.

Figure 2.4 illustrates a Box partition over 3 processes.

Orange partition

Each partition is an ensemble of segments in the global domain. Each segment is described by its global offset and its local extent. In this case, we have `ig_parallel(1:N)` where $N = 2 + 2 * \text{number of segments}$

- `ig_parallel(1)` = 3 (indicates a Orange partition)
- `ig_parallel(2)` = the total number of segments for the partition (limited to 200 presently, see note for `ig_parallel(4)` for Box partition above)
- `ig_parallel(3)` = the first segment global offset
- `ig_parallel(4)` = the first segment local extent
- `ig_parallel(5)` = the second segment global offset
- `ig_parallel(6)` = the second segment local extent
- ...
- `ig_parallel(N-1)` = the last segment global offset
- `ig_parallel(N)` = the last segment local extent

Figure 2.5 illustrates an Orange partition with 3 segments for one process. The other process partitions are not illustrated.

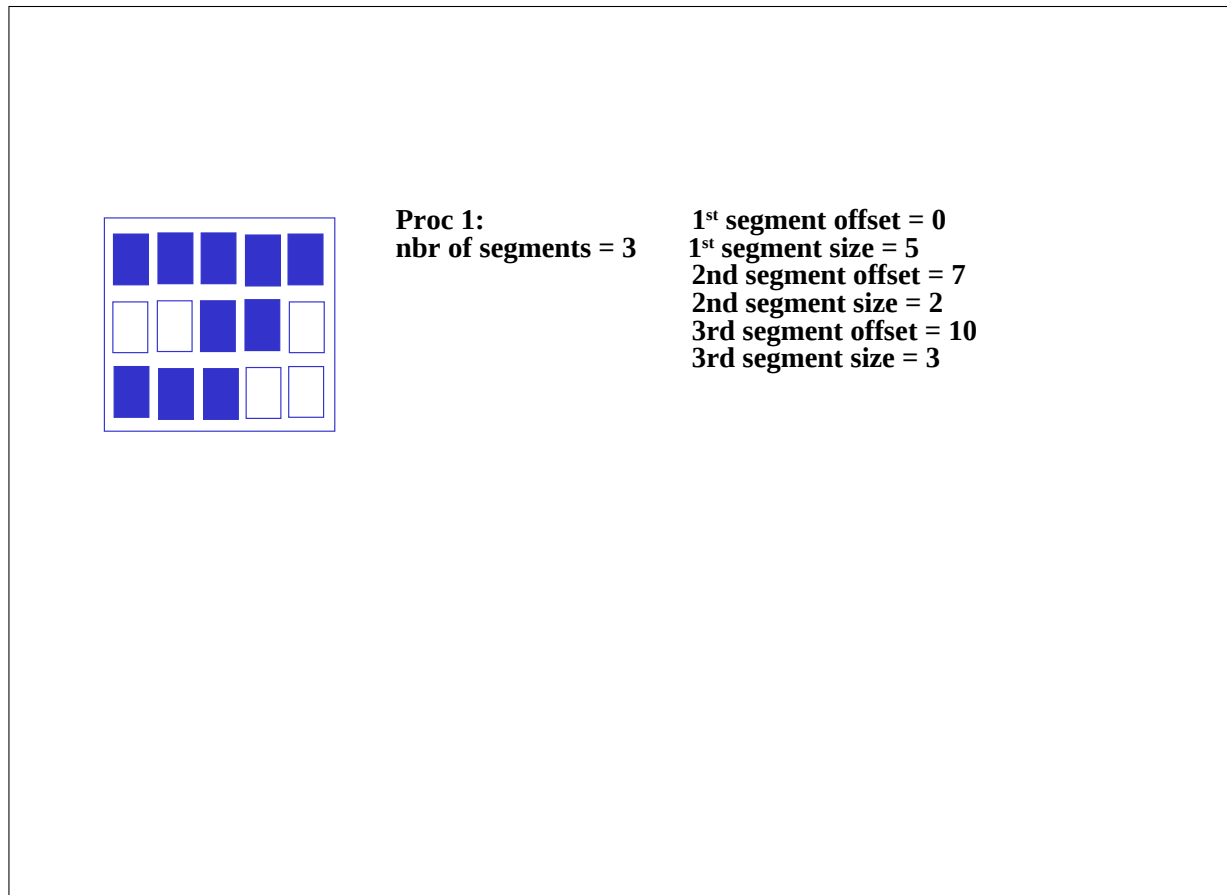


Figure 2.5: Orange partition for one process. We assume here that the global index starts at 0 in the upper left corner.

Note that each partition has to be defined with monotonically increasing global index values.

Points partition

This partition is a list of global indices associated with each process. The index naming is arbitrary but must be consistent between all processes involved in the partition description. In this case, we have `ig_parallel(1:N)` where $N = 2 + \text{number of points}$. As for Orange, each partition has to be defined with monotonically increasing global index values.

- `ig_parallel(1) = 4` (indicates a Points partition)
- `ig_parallel(2) = number of points in the partition`
- `ig_parallel(3) = the first global index`
- `ig_parallel(4) = the second global index`
- ...
- `ig_parallel(N) = the last global index`

Cube partition

For the special case of 3D fields interpolation with user-provided remapping weights, OASIS3-MCT supports a partition which acts as a Box partition at every vertical level without the need of repeating it. On every 2D level, each local partition is a rectangular region of the 2D global domain, described by the global offset of its upper left corner, and its local extents in the X and Y 2D dimensions. The global extents in

the X and Y 2D dimensions and the number of vertical levels must also be given. In this case, we have `ig_paral(1:7)`:

- `ig_paral(1) = 5` (indicates a Cube partition)
- `ig_paral(2) = the upper left corner global offset in the 2D domain`
- `ig_paral(3) = the local extent in X`
- `ig_paral(4) = the local extent in Y`
- `ig_paral(5) = the global extent in X`
- `ig_paral(6) = the global extent in Y`
- `ig_paral(7) = the number of vertical levels`

This type of partition is adapted to specific couplings, e.g. horizontal coarsening in the exchanges between a high-resolution ocean model and a lower-resolution bio-geo-chemistry model. An example of how to use this type of partition is given in `/pyoasis/examples/14-test-inter3d`.

2.2.4 Grid data file definition

Grid data files are required by OASIS3-MCT for specific operations, see sections 4 and 5.1. These grid data files can be created by the user before the run or can be written directly at run time by the components with the routines described in this section. In that last case, the corresponding routine will create a grid data files if it does not exist. If the grid data file does exist, the routine can still be used to **add** grid definition in that file, but it will not **overwrite** a grid definition if it already exist in the file with the same grid name.

These routines can be called only by one component process to write the whole grid or by each process holding a part of a grid. In the former case, optional argument `il_part_id` is not needed and the arrays handling the longitudes of the grid points or corners, the latitudes of the grid points or corners, the masks, fractions and areas of the grid cells need to cover the whole grid. In the later case, the `il_part_id` returned by `oasis_def_partition` needs to be provided as input argument and the arrays need to cover only the local partition of the grid.

The field names in the `grids.nc`, `masks.nc`, and `areas.nc` follow a well-defined convention (see also section 5.1). The fields are two-dimensional, and each field name consists of a grid acronym followed by a string that identifies the field. For instance, the center latitudes and longitudes for the grid `torc` will be called respectively `torc.lat` and `torc.lon`. The `grids.nc` file contains the center latitudes (`.lat`) and longitudes (`.lon`) as well as the corner latitudes (`.cla`) and corner longitudes (`.clo`). The corner fields have a third dimension associated with the number of corners per gridcell. The `area.nc` file contains the area field (`.srf`). The `masks.nc` file contains the mask (`.msk`) and static fraction (`.frc`) fields.

Note that if all remappings are based on user-defined weights, the files `grids.nc` and `areas.nc` are not required, while `masks.nc` is only necessary for the CHECKIN or CHECKOUT diagnostics (see sections 4).

- CALL `oasis_start_grids_writing (flag)`
 - `flag [INTEGER; OUT]: always 1`

Must be called to start the grid writing process.
- CALL `oasis_write_grid (cgrid, nx_global, ny_global, lon, lat, il_part_id)`
 - `cgrid [CHARACTER; IN]: grid name prefix (see 3.3 and 5.1); maximum length of 64 characters (4 are usually used for historical reasons)`
 - `nx_global [INTEGER; IN] : first dimension of the global grid`
 - `ny_global [INTEGER; IN] : second dimension of the global grid (=1 if the grid is expressed as a 1D vector)`
 - `lon [REAL, DIMENSION(nx,ny); IN] : single or double real array of longitudes`

covering the whole grid ($nx=nx_global, ny=ny_global$) or only the local partition (degrees East).

- `lat` [REAL, DIMENSION(nx, ny); IN] : single or double real array of latitudes covering the whole grid ($nx=nx_global, ny=ny_global$) or only the local partition (degrees North)
- `il_part_id` [INTEGER, OPTIONAL; IN] : partition ID returned by `oasis_def_partition`, see 2.2.3; needed if each component task writes its own part of the grid.

Writes the component grid longitudes and latitudes. Longitudes must be given in degrees East in the interval -360.0 to 720.0. Latitudes must be given in degrees North in the interval -90.0 to 90.0. Note that if some grid points overlap, it is recommended to define those points with the same number (e.g. 90.0 for both, not 450.0 for one and 90.0 for the other) to ensure automatic detection of overlap by OASIS3-MCT (which is essential to have a correct conservative remapping SCRIPR/CONSERV, see section 4.3).

- CALL `oasis_write_corner` (`cgrid, nx_global, ny_global, nc, clon, clat, il_part_id`)
 - `cgrid, nx_global, ny_global, il_part_id` : as for `oasis_write_grid`
 - `nc` [INTEGER; IN] : number of corners per grid cell (can be any number)
 - `clon` [REAL, DIMENSION (nx, ny, nc); IN] : single or double real array of corner longitudes covering the whole grid ($nx=nx_global, ny=ny_global$) or only the local partition (in degrees East)
 - `clat` [REAL, DIMENSION (nx, ny, nc); IN] : single or double real array of corner latitudes covering the whole grid ($nx=nx_global, ny=ny_global$) or only the local partition (in degrees North)

Writes the grid cell corner longitudes and latitudes (counterclockwise sense). Longitudes must be given in degrees East in the interval -360.0 to 720.0. Latitudes must be given in degrees North in the interval -90.0 to 90.0. Note also that cells larger than 180.0 degrees in longitude are not supported. Writing of corners is optional as corner information is needed only for SCRIPR/CONSERV (see section 4.3). If called, needs to be called after `oasis_write_grid`.

- CALL `oasis_write_mask` (`cgrid, nx_global, ny_global, mask, il_part_id, companion`)
 - `cgrid, nx_global, ny_global, il_part_id` : as for `oasis_write_grid`
 - `mask` [INTEGER, DIMENSION(nx, ny); IN] : mask array covering the whole grid ($nx=nx_global, ny=ny_global$) or only the local partition. Be careful about OASIS3-MCT historical convention (!): 0 = not masked (i.e. active), 1 = masked (i.e. not active).
 - `companion` [CHARACTER ; IN; OPTIONAL] : the character string value that will be associated with the mask field attribute `coherent_with_grid`. This will be written to the `masks.nc` netcdf file with the mask field. It is purely informational and used in cases where the mask field is derived from or consistent with another grid.

Writes the component grid mask. The mask field should be consistent with the static fraction field (see below) and will define the 0/1 mask of the grid cell. The mask field is used by both the SCRIPR remappings and in the global CONSERV operations if defined. The mask field is written to the `masks.nc` file.

- CALL `oasis_write_frac` (`cgrid, nx_global, ny_global, frac, il_part_id, companion`)
 - `cgrid, nx_global, ny_global, il_part_id` : as for `oasis_write_grid`

- `frac` [REAL, DIMENSION(nx,ny) ; IN] : single or double real fraction array covering the whole grid (nx=nx_global, ny=ny_global) or only the local partition.
- `companion` [CHARACTER ; IN; OPTIONAL] : the character string value that will be associated with the fraction field attribute “coherent_with_grid”. It should refer to the acronym of the grid which mask was used to define the fraction of the current grid (see section 4.4). This will be written to the *masks.nc* NetCDF file with the fraction field. It is purely informational and used in cases where the fraction field is derived from or consistent with another grid mask.

Writes the component grid cell static fractions. This should be consistent with the mask field and defines the fraction of the grid cell that is active (i.e. not masked), e.g. the water fraction of an atmospheric cell having water and land fractions. The fraction field is only used in the global CONSERV operations. Either the mask or fractions must be defined for that operation. If both are defined, they must be consistent; OASIS3-MCT will abort if they are not coherent or if both are missing. Note that by OASIS3-MCT conventions for the mask, a gridcell with mask=0 (active) should have a fractions greater than 0 and a gridcell with mask=1 (inactive) should have a fractions equal to 0. The fraction field is written to the *masks.nc* file.

- CALL `oasis.write_area` (cgrid, nx_global, ny_global, area, il_part_id)
 - `cgrid`, `nx_global`, `ny_global`, `il_part_id` : as for `oasis.write_grid`
 - `area` [REAL, DIMENSION(nx,ny) ; IN] : single or double real array of grid cell areas covering the whole grid (nx=nx_global, ny=ny_global) or only the local partition

Writes of the component grid cell areas. Needed for some SCRIPR options and for the CONSERV operation (see section 4.4). The area field is written to the *areas.nc* file. The surfaces may be given in any units but they must be coherent on the source and target sides. Furthermore they must be in square radians if the True Area (TR) correction is activated, see section 4.3.

- CALL `oasis.write_angle` (cgrid, nx_global, ny_global, angle, il_part_id)
 - `cgrid`, `nx_global`, `ny_global`, `il_part_id` : as for `oasis.write_grid`
 - `angle` [REAL, DIMENSION(nx,ny) ; IN] : single or double real array of grid cell angles covering the whole grid (nx=nx_global, ny=ny_global) or only the local partition

Writes of the component grid cell angles. The angle field is written to the *grids.nc* file. This field does not play a role in OASIS3-MCT implementation and is never needed.

- CALL `oasis.terminate_grids_writing()`

The creation of the different grid data files is completed in the routine `oasis_enddef`.

2.2.5 Coupling field declaration

All processes of a component that send or receive a coupling field, or a part of it, needs to declare the coupling field.

Processes not implied in the coupling do not have to call this routine at all but for backward compatibility with OASIS3-MCT.2.0, they may still call it with any name and `il_part_id`.

- CALL `oasis_def_var` (var_id, name, il_part_id, var_nodims, kinout, var_actual_shape, var_type, kinfo)
 - `var_id` [INTEGER; OUT]: coupling field ID. Note that all coupling fields appearing in the *namcouple* must be declared with a call to `oasis_def_var`; not doing so would lead to an abort. But all fields defined with a call to `oasis_def_var` must not necessarily be activated in the *namcouple*. If a field does not appear in the *namcouple*, the `var_id` returned by the `oasis_def_var` will be -1. The value of the `var_id` should be tested and if `var_id`

- equals -1, the corresponding `oasis_put` and `oasis_get` should not be called. These constraints are imposed to avoid that a typo in the *namcouple* leads to coupling exchanges not corresponding to what the user intends to activate.
- `name` [CHARACTER; IN]: field symbolic name (as in the *namcouple*); maximum length of 80 characters
 - `il_part_id` [INTEGER; IN]: partition ID returned from `oasis_def_partition` (see section 2.2.3)
 - `var_nodims` [INTEGER, DIMENSION(2); IN]: is an integer array of size two. The first element, `var_nodims(1)`, is not used anymore in OASIS3-MCT, so its value can be anything. The second element, `var_nodims(2)`, is the number of fields in a bundle (this will be 1 for unbundle fields or greater than 1 for fields that are bundled; if `var_nodims(2)=0`, it will be automatically reset to 1 in the routine, to ensure backward compatibility).
 - `kinout` [INTEGER; IN]: OASIS_In (i.e. = 21) for fields received by the component; OASIS_Out (i.e. = 20) for fields sent by the component ².
 - `var_actual_shape` [INTEGER, DIMENSION(:) or DIMENSION(:, :), IN]: optional, not used anymore but can still be passed for backwards compatibility; the interface is overloaded to support a 1d or 2d array of any length.
 - `var_type` [INTEGER; IN]: type of coupling field array; put OASIS_Real (i.e. = 4) for single or double precision real arrays. All coupling data is treated as double precision in the coupling layer, but conversion to or from single precision data is supported by the API.
 - `kinfo` [INTEGER; OUT]: returned error code.

2.2.6 End of definition phase

All processes of components at least partly involved in the coupling (e.g. `comp3` in figure 2.2) have to close the definition phase. Different configurations of components and corresponding use of `oasis_enddef` are described in section 2.1 and on figures 2.1 and 2.2.

- CALL `oasis_enddef` (`kinfo`)
 - `kinfo` [INTEGER; OUT]: returned error code.

2.2.7 Sending “put” and receiving “get” actions

This section describes how to send (put) and receive (get) fields through OASIS-MCT API. This coupling interface supports several ranks and types of coupling fields. The field decomposition must be consistent with the grid partition (see 2.2.3)³ and the bundle fields are supported (i.e. have an extra non-spatial dimension, for example for different ice categories).

The bundle field option can be used to send or receive together multi-level variables, multiple related fields, and other types of fields. The fields must share a common partition and common *namcouple* settings (e.g. interpolation) to be bundled. While this is a useful feature for multi-level fields, this does not mean that 3D interpolation is supported. Each field in the bundle is treated internally as a separate field in the coupling layer without any information about the relationship between the fields in the bundle. In fact, the bundle field variables are internally renamed and a field number is appended to the variable name to keep track of the distinct fields in the bundle. That updated variable name will appear in restart and output files.

It is possible to define a 1D bundle or 2D bundle field with a bundle dimension of 1. So in general, `fld`, the field passed into the get and put interface can have rank 1, 2, or 3 and includes the following possible options:

²Parameters OASIS_In, OASIS_Out are defined in `oasis3-mct/lib/psmile/src/mod_oasis_parameters.F90`

³But the field rank does not necessarily have to match the rank of the grid partition description (i.e. it can be expressed in either 1D or 2D) as long as it provides overall the same number of elements.

- 1D, fld(:) = a single, unbundle field of rank 1.
- 2D, fld(:, :) = a single, unbundle field of rank 2.
- 3D, fld(:, :, :) = a single, unbundle field of decomposition rank 3 ; special case linked with the Cube partition (see section 2.2.3) adapted to specific couplings (e.g. horizontal coarsening in the exchanges between a high-resolution ocean model and a lower-resolution bio-geo-chemistry model) that requires user-defined interpolation remapping files, i.e. MAPPING (*cf.* section 4.3) but no vertical interpolation.
- 1D bundle, fld(:, :) = a bundle set of fields of rank 1 (the size of the second dimension must equal the number of fields in the bundle, defined by var_nodims(2) in the oasis_def_var interface).
- 2D bundle, fld(:, :, :) = a bundle set of fields of rank 2. (the size of the third dimension must equal the number of fields in the bundle, defined by var_nodims(2) in the oasis_def_var interface).

For a given bundle, the number of fields must match in the source and target models. This is explicitly checked within the coupling layer and will lead to an abort if the match is not effective.

Sending a coupling field (or writing a coupling restart file)

In the component time step loop, each process sends its part of the coupling (or I/O) field.

- CALL oasis_put (var_id, date, fld1, info, fld2, fld3, fld4, fld5, write_restart, fracwgt)
 - var_id [INTEGER; IN]: field ID (returned from corresponding oasis_def_var, see section 2.2.5)
 - date [INTEGER; IN]: number of seconds (or any other time units as long as they are the same in all components and in the *namcouple*) at the time of the call
 - fld1 [REAL, IN]: coupling (or I/O) field array; can be 4-byte or 8-byte 1D, 2D, 3D, bundle 1D, or bundle 2D, see above.
 - info [INTEGER; OUT]: returned info code:
 - * OASIS_Sent (=4) : the field was sent to another component
 - * OASIS_LocTrans (=5) : the field was only used in a time transformation (not sent, not output)
 - * OASIS_ToRest (=6) : the field was written to a restart file only
 - * OASIS_Output (=7) : the field was written to an output file only
 - * OASIS_SentOut (=8) : the field was both written to an output file and sent to another component
 - * OASIS_ToRestOut (=9): the field was written both to a restart file and to an output file.
 - * OASIS_WaitGroup (=14) : the field was not sent because it is part of a group (see "Support to group multiple fields in a single communication" in section 3.3.1. It will be sent only when the oasis_put of the last field in the group will be called. However, the field is buffered and therefore the field array can be modified in the component code when returning from the oasis_put call.
 - * OASIS_Ok (=0) : otherwise and no error occurred.
 - fld2/fld3/fld4/fld5 [REAL, IN, OPTIONAL]: optional higher-order 2nd/3rd/4th/5th coupling field array; 4-byte or 8-byte 1D, 2D, 3D, bundle 1D, or bundle 2D; rank and size must match fld1; needed for higher-order remapping, see details below.
 - write_restart [LOGICAL, IN, OPTIONAL]: optional argument to write the field array associated with var_id in an intermediate restart file at the current timestep, taking into account LOCTRANS operations; false by default. If multiple fields are grouped in a single operation in the model (see "Support to group multiple fields in a single communication" in

section 3.3.1), `write_restart` should be true for the `oasis_put` calls of all corresponding fields. Intermediate restart files are created with 9 digits in their name, corresponding to the time of the `oasis_put` call, e.g. `TC000014400_rst4.nc`, where "rst4.nc" is the coupling restart filename defined in the *namcouple*.

- `fracwgt` [REAL, IN, OPTIONAL]: optional fraction weight field array; 4-byte or 8-byte 1D, 2D, 3D, bundle 1D, or bundle 2D; it may vary in time and represents the fraction of the cell for which `fld1` is valid, like an ice fraction. Calling `oasis_put` with this optional argument is referred to as the `fracwgt` option in the rest of the text and is described in details in Appendix A.
 - * `fracwgt` rank must match `fld1` rank; `fracwgt` grid size must match `fld1` grid size; if `fld1` has a bundle dimension, then `fracwgt` must have a bundle dimension that can be the same than `fld1` bundle dimension (and in that case, `fracwgt` can be different for the different bundle members) or that can be equal to 1 (and in that case, `fracwgt` is the same for the different bundle members).
 - * `fracwgt` cannot be combined with global CONSERV (see section 4.4) if LOCTRANS AVERAGE or ACCUMUL is also activated, or with LOCTRANS T_MIN or T_MAX (see section 4.1).
 - * If the `fracwgt` option is activated for a field with a LAG (see section 3.3), special care have to be taken on the content of the coupling restart files; see section 5.2 for details.
 - * `fracwgt` option can only be carried out with remapping on the source model processors, (`$MAPLOC=src` in section 4.3).
 - * If higher-order transforms (CONSERV/SECOND or BICUBIC) are specified for the coupling fields, they are applied on the target grid without regard to `fracwgt`.

To ensure a proper use of the `oasis_put`, one has to take care of the following aspects:

- This routine may be called by the component at each timestep. The convention for the `date` argument is to indicate the time at the beginning of the timestep. The sending is actually performed if the time obtained by adding the field lag (LAG in the *namcouple*, if any, with LAG=0 by default) to the `date` corresponds to a coupling period, as specified by the user in the *namcouple* (see section 3). By convention, the first coupling of a run occurs at `date=0`.
- Trying to send with `oasis_put` a field declared with a `oasis_def_var` but not defined in the configuration file *namcouple* will lead to an abort. When a field is not defined in the *namcouple*, the field ID returned by the `oasis_def_var` is equal to -1 and the corresponding `oasis_put` should not be called.
- For a coupling field with a positive lag, a coupling restart file containing the initial values of the field must be provided and will be used at the beginning of the run (see section 5.2) is automatically overwritten by the `oasis_put` when the `date+LAG=runtime`. **It is the user's responsibility to ensure that the overwritten restart file is copied in the working directory of the next run.**
- The total run time should match an integer number of coupling periods. No warranty is given when a run terminates in the middle of a coupling period, although compiling with the CPP key LOCTRANS_RESTARTS ensures that partial time transformations, if any, are saved in the coupling restart file at the end of the run.
- If a local time transformation is indicated for the field in the *namcouple* (INSTANT, AVERAGE, ACCUMUL, T_MIN or T_MAX, see section 4), it is automatically performed below the `oasis_put` and the resulting field is sent at the coupling period.
- A 2nd, 3rd, 4th and 5th source field can be passed as optional arguments for higher-order remapping. If `fld3` is passed, `fld2` must also be passed; if `fld4` is passed, `fld3` and `fld2` must also be passed; if `fld5` is passed `fld4`, `fld3` and `fld2` must also be passed. If passed, the 2nd, 3rd, 4th and 5th set of weights present in the remapping file will be applied to these fields respectively

(see section 5.4 for the remapping file format). From OASIS3-MCT_4.0 onwards, the number of weights in the remapping file and the number of fields in the coupling restart file (when such a file is needed) must strictly match the number of source fields passed to the `oasis_put`. The higher-order restart fields in the coupling restart file must have the same name than the coupling field itself prefixed with either `av2_`, `av3_`, `av4_`, or `av5_`, see also section 5.2.

This possibility of passing higher-order fields must be used for example for the higher-order SCRIPR BICUBIC remapping for which a 1st, 2nd, 3rd, 4th set of weights should be respectively applied to the field value, its gradient in the first dimension, its gradient in the second dimension, and its cross-gradient in that order.

This will be used also for the higher-order remapping SCRIPR CONSERV/SECOND for which a 1st, 2nd, 3rd set of weights must be respectively applied to the field value, its gradient with respect to the latitude (θ) $\frac{\delta f}{\delta \theta}$ and its gradient with respect to the longitude (ϕ) $\frac{1}{\cos \theta} \frac{\delta f}{\delta \phi}$ in that order.

- A coupling field sent by a source component can be associated with more than one target field and component, with different entries in the *namcouple* configuration file. In that case, the source component needs to send the field only once and the corresponding data will arrive at multiple targets as specified in the *namcouple*. Different coupling frequencies and transformations are allowed for different coupling exchanges of the same field. If coupling restart files are required (either if a LAG is specified for that field), it is mandatory to specify different files for the different fields.
- The `oasis_put` routine cannot be called from OpenMP-threaded regions.

Receiving a coupling (or I/O) field

In the component time stepping loop, each process receives its part of the coupling field.

- CALL `oasis_get (var_id, date, fld, info)`
 - `var_id` [INTEGER; IN]: field ID (returned by the corresponding `oasis_def_var`)
 - `date` [INTEGER; IN]: number of seconds (or any other time units as long as the same are used in all components and in the *namcouple*) at the time of the call (by convention at the beginning of the timestep)
 - `fld` [REAL, OUT]: I/O or coupling field array; can be 1D, 2D, 3D, bundle 1D, or bundle 2D.
 - `info` [INTEGER; OUT]: returned info code:
 - * OASIS_Recvd(=3) if the field was received from another component
 - * OASIS_FromRest (=10) if the field was read from a restart file only
 - * OASIS_Input (=11) if the field was read from an input file only
 - * OASIS_RecvOut (=12) if the field was both received from another component and written to an output file
 - * OASIS_FromRestOut (=13) if the field was both read from a restart file and written to an output file
 - * OASIS_Ok (=0) otherwise and no error occurred.

To ensure a proper use of the `oasis_get`, one has to take care of the following aspects:

- This routine may be called by the component at each timestep. The `date` argument is automatically analysed and the receiving action is actually performed only if `date` corresponds to a time for which it should be activated, given the period indicated by the user in the *namcouple*. An exchange at the beginning of the run at time=0 is expected.
- Trying to receive with `oasis_get` a field declared with a `oasis_def_var` but not defined in the configuration file *namcouple* will lead to an abort. In this case, the field ID returned by the `oasis_def_var` is equal to -1 and the corresponding `oasis_get` should not be called.

- If a coupling field has a positive lag, the coupling field that matches the `oasis_get` at `time=0` is automatically read in from a coupling restart file in the source model under its `oasis_undef` and sent to match that `oasis_get` (see section 2.5.2).
- The `oasis_get` routine cannot be called from an OpenMP threaded region.

2.2.8 Termination

- CALL `oasis_terminate (kinfo)`
 - `kinfo` [INTEGER; OUT]: returned error code.

All processes have to terminate the coupling by calling this routine. Note that if the process called `MPI_Init`, necessarily before calling `oasis_init_comp`, it must also call `MPI_Finalize` explicitly, but only after calling `oasis_terminate`. Different configurations of components and corresponding use of `oasis_terminate` are described in section 2.1 and on figures 2.1 and 2.2.

2.2.9 Auxiliary routines

The following auxiliary routines are currently available.

- CALL `oasis_abort (compid, routine_name, abort_message, file, line, rcode)`
 - `compid` [INTEGER; IN]: component ID (from `oasis_init_comp`)
 - `routine_name` [CHARACTER*; IN]: name of calling routine
 - `abort_message` [CHARACTER*; IN]: message to be written out
 - `file` [CHARACTER*; OPTIONAL; IN]: file from which `oasis_abort` is called from
 - `line` [INTEGER, OPTIONAL; IN]: line in file from which `oasis_abort` is called from
 - `rcode` [INTEGER, OPTIONAL; IN]: Optional argument. When OASIS3-MCT aborts, it returns `rcode` if it is present, else it returns 1

If a process needs to abort voluntarily, it should do so by calling `oasis_abort`. This will ensure a proper termination of all processes in the coupled model communicator. This routine writes the name of the calling component, the name of the calling routine, and the message to the process debug file (see `$NLOGPRT` in section 3.2). This routine cannot be called before `oasis_init_comp`.

- CALL `oasis_get_debug(debug, kinfo)`
 - `debug` [INTEGER; OUT]: output debug value
 - `kinfo` [INTEGER; OUT; OPTIONAL]: returned error code

This routine may be called at any time to retrieve the current OASIS3-MCT internal debug level (see `$NLOGPRT` in section 3.2).

- CALL `oasis_set_debug(debug, kinfo)`
 - `debug` [INTEGER; IN]: input debug value
 - `kinfo` [INTEGER; OUT; OPTIONAL]: returned error code

This routine may be called at any time to change the debug level in OASIS3-MCT.

- CALL `oasis_get_intercomm(new_comm, cdnam, kinfo)`
 - `new_comm` [INTEGER; OUT]: MPI inter-communicator
 - `cdnam` [CHARACTER*; IN]: other component name (i.e. 2nd argument of the call to `oasis_init_comp` in that component)
 - `kinfo` [INTEGER; OUT; OPTIONAL]: returned error code

This routine sets up an MPI inter-communicator between two components, the local component and the component associated with `cdnam`. This call is collective across the tasks of the two components

only. An MPI inter-communicator preserves the rank of the original communicators and does not allow collective communication within the communicator. It provides point to point communication between two non-overlapping MPI groups. This method must be called synchronously across all components involved to minimize the chance of a deadlock, and it should be called only after `oasis_enddef`.

- CALL `oasis_get_intracomm(new_comm, cdnam, kinfo)`
 - `new_comm` [INTEGER; OUT]: MPI intra-communicator
 - `cdnam` [CHARACTER*; IN]: single string argument containing the other component name (i.e. 2nd argument of the call to `oasis_init_comp` in that component).
 - `kinfo` [INTEGER; OUT; OPTIONAL]: returned error code

This routine sets up an MPI intra-communicator between two components, the local component and the component defined by the string `cdnam`. This call is collective across the tasks of the two components creating the intra-communicator only, and it must be called synchronously across all tasks of the two components to minimize the chance of a deadlock. It should be called only after `oasis_enddef`. This method creates a new communicator consisting of a new collective group of tasks with new ranks. This communicator supports collective communications and is more typically used in MPI applications than inter-communicators (see `oasis_get_intercomm` above).

- CALL `oasis_get_multi_intracomm(new_comm, cdnam, root_ranks, kinfo)`
 - `new_comm` [INTEGER; OUT]: MPI intra-communicator
 - `cdnam` [CHARACTER*; IN]: 1d array of character strings (i.e. `cdnam(:)`) containing the name of the component (i.e. 2nd argument of the call to `oasis_init_comp` in each component).
 - `root_ranks` [INTEGER; OUT]: 1d integer array (i.e. `root_ranks(:)`) of the same size than `cdnam` containing root ranks. This argument is a
 - `kinfo` [INTEGER; OUT]: returned error code

This routine sets up an MPI intra-communicator between two or more components defined by the component names passed in the `cdnam` array argument. The local model name **MUST BE** one of the models defined in the `cdnam` array. The component names must be valid names, but empty strings are allowed and ignored. This call is collective across all the tasks of the components defined in `cdnam`, and it must be called synchronously and consistently across all tasks of those components to minimize the chance of a deadlock. It should be called only after `oasis_enddef`. This method creates a new communicator consisting of a new collective group of tasks with new ranks. The root rank of each individual components relative to the new communicator is output in the `root_ranks` argument. The size of `cdnam` and `root_ranks` should be identical, and the values of `root_ranks` are consistent with the order of `cdnam`. This communicator supports collective communications and is more typically used in MPI applications than inter-communicators (see `oasis_get_intercomm` above).

- CALL `oasis_put_inquire(var_id, date, kinfo)`
 - `var_id` [INTEGER; IN]: field ID (from corresponding `oasis_def_var`)
 - `date` [INTEGER; IN]: as in `oasis_put`, number of seconds (or any other time units as long as the same are used in all components and in the *namcouple*) in the run at the time of the call
 - `kinfo` [INTEGER; OUT]: returned info code
 - * `OASIS_Sent`(=4) if the field would be sent to another component
 - * `OASIS_LocTrans` (=5) if the field would be only used in a time transformation (not sent, not output)
 - * `OASIS_ToRest` (=6) if the field would be written to a restart file only

- * OASIS_Output (=7) if the field would be written to an output file only
- * OASIS_SentOut (=8) if the field would be both written to an output file and sent to another component (directly or via OASIS3 main process)
- * OASIS_ToRestOut (=9) if the field would be written both to a restart file and to an output file.
- * OASIS_Ok (=0) otherwise and no error occurred.

This routine may be called at any time to inquire what would happen to the field corresponding to that `var_id` if it was sent with an `oasis_put` at that same date. This maybe useful if, for example, the calculation of a coupling field is costly and if one wants to compute it only when it is really sent out.

- CALL `oasis_get_ncpl(var_id, ncpl, kinfo)`
 - `var_id` [INTEGER; IN]: field ID (from corresponding `oasis_def_var`)
 - `ncpl` [INTEGER; OUT]: number of coupling exchanges in which the field is involved (i.e. when a field is sent to multiple targets)
 - `kinfo` [INTEGER; OUT]: returned info code

This routine returns the number of coupling exchanges in which the field with that `var_id` is involved. This number is needed to get the coupling frequencies with the routine `oasis_get_freqs`, see below.

- CALL `oasis_get_freqs(var_id, mop, ncpl, cpl_freqs, kinfo)`
 - `var_id` [INTEGER; IN]: field ID (from corresponding `oasis_def_var`)
 - `mop` [INTEGER; IN]: OASIS_Out or OASIS_In
 - `ncpl` [INTEGER; IN]: number of couplings in which the field is involved (i.e. when a field is sent to multiple targets)
 - `cpl_freqs` [INTEGER; DIMENSION(ncpl); OUT]: coupling period(s) of field `var_id`. There is one coupling period for each coupling exchange in which the field is involved
 - `kinfo` [INTEGER; OUT]: returned info code

This routine can be used to retrieve the coupling period(s) of field with corresponding `var_id`, as defined in the *namcouple*.

2.3 OASIS3-MCT C API

OASIS3-MCT is distributed with C bindings and can be called from models written in C and in C++. These bindings leverage the Fortran ISO_C_BINDING standard. The C bindings can be compiled into static or shared libraries by the OASIS3-MCT TopMakefileOasis3 as documented in 6.1.

The C interfaces largely match up with equivalent interfaces in Fortran. An interface named `oasis_interface` in Fortran can be expected to be named `oasis_c_interface` in C.

All of the C interfaces return an integer error code which can be tested against the `OASIS_Ok` (or the equivalent `OASIS_Success`) constant. The `OASIS_CHECK_ERR` macro aborts OASIS3-MCT with a meaningful message in the debug files in case of failure. For most of the functions, the return code is consistent with the `kinfo` argument in the Fortran interfaces (see section 2.2). However, for the put and get communication functions, the return error code only indicates success or failure, while the detailed status is returned by the `kinfo` argument (equivalent to the returned value of `info` argument for the Fortran `oasis_put` and `oasis_get` routines, see section 2.2.7).

For example, the following constructs are equivalent in the two languages:

```
call oasis_get_localcomm(localcomm, kinfo)
if (kinfo .ne. OASIS_Ok) &
    & call OASIS_Abort(comp_id, "oasis_get_localcomm", &
```

```
& "Runtime error", __FILE__, __LINE__)
```

and

```
OASIS_CHECK_ERR(oasis_get_localcomm(localcomm))
```

For convenience a similar macro `OASIS_CHECK_MPI_ERR` has been defined for testing the return code of any MPI function against `MPI_SUCCESS` and cleanly aborting OASIS3-MCT in case of failure.

Use of these C bindings are illustrated in practical examples in directories `/C` in the different subdirectories in `pyoasis/examples/`. Notes and deviations from the Fortran standard are noted below.

- To use the OASIS3-MCT C bindings, the statement


```
#include "oasis_c.h"
```

 needs to be added to the C model source code. All available parameters macros and interfaces are defined there.
- `int oasis_c_init_comp(int* compid, const char* comp_name, const bool coupled)`
 - This `oasis_c_init_comp` interface includes the coupled flag but no commworld argument; use `oasis_c_init_comp_with_comm` below to provide a communicator argument.
 - The coupled argument can receive the predefined mnemonic boolean constants `OASIS_Coupled` and `OASIS_Not_Coupled`
- `int oasis_c_init_comp_with_comm(int* compid, const char* comp_name, const bool coupled, const MPI_Comm commworld)`
 - This alternative `init_comp` interface has the coupled flag and a C `MPI_Comm` communicator
- `int oasis_c_get_localcomm(MPI_Comm* local_comm)`
 - Returns the local communicator by reference in a C `MPI_Comm*` type argument
- `int oasis_c_create_couplcomm(const int icpl, const MPI_Comm local_comm, MPI_Comm* coupl_comm)`
 - `icpl` is a C `int` type input argument (see `oasis_create_couplcomm` in section 2.2.2)
 - `local_comm` is a C `MPI_Comm` type input argument (by value)
 - `coupl_comm` is a C `MPI_Comm*` type output argument (by reference)
- `int oasis_c_set_couplcomm(const MPI_Comm coupl_comm)`
 - Takes a C `MPI_Comm` type as an input argument (by value)
- `int oasis_c_def_partition(int* il_part_id, const int ig_paral_size, const int* ig_paral, const int ig_size, const char* name)`
 - `ig_paral_size` is the size of the array `ig_paral`: the `oasis_c.h` header provides a set of constants and macros for the size of every partition strategy, namely `OASIS_Serial_Params`, `OASIS_Apple_Params`, `OASIS_Box_Params`, `OASIS_Orange_Params` (`n_segments`), `OASIS_Points_Params` (`n_points`)
 - the `oasis_c.h` header also provides a set of predefined constants for the storages positions in `ig_paral`, namely `OASIS_Strategy`, `OASIS_Segments`, `OASIS_Npoints`, `OASIS_Offset`, `OASIS_Length`, `OASIS_SizeX`, `OASIS_SizeY`, `OASIS_LdX`
 - the partition strategy, to be stored in the `OASIS_Strategy` position of `ig_paral`, can take one of the following predefined constants values: `OASIS_Serial`, `OASIS_Apple`, `OASIS_Box`, `OASIS_Orange`, `OASIS_Points`

- for the cases in which the `ig_size` and `name` arguments are not relevant for the partition definition, the placeholders `OASIS_No_Gsize` and `OASIS_No_Name` can be used instead

Here an example for the `OASIS_Part_Apple` strategy

```
int part_params[OASIS_Apple_Params];
part_params[OASIS_Strategy] = OASIS_Apple;
part_params[OASIS_Offset] = offset;
part_params[OASIS_Length] = local_size;
int part_id;
OASIS_CHECK_ERR(oasis_c_def_partition(&part_id, OASIS_Apple_Params,
                                     part_params, OASIS_No_Gsize,
                                     OASIS_No_Name));
```

- `int oasis_c_start_grids_writing()`
- `int oasis_c_write_grid(const char* cgrid, const int nx_global, const int ny_global, const int nx_loc, const int ny_loc, const double* lon, const double* lat, const int il_partid)`
 - `/local`
 - `nx_global` and `ny_global` are the first and second dimensions of the global grid
 - `nx_loc` and `ny_loc` are the two dimensions of the local arrays `lon` and `lat`
 - `lon` and `lat` are stored with a Fortran compatible (column major) ordering. For example, if they are declared as double pointers, they have to be `lon[ny_loc][nx_loc]`
 - `il_partid` is relevant only for parallel writing; in case of single proc invocations, the constant `OASIS_No_Part` can be passed as a placeholder
- `int oasis_c_write_corner(const char* cgrid, const int nx_global, const int ny_global, const int nc, const int nx_loc, const int ny_loc, const double* clon, const double* clat, const int il_partid)`
 - `nx_global` and `ny_global` are the first and second dimensions of the global grid
 - `nc` is the maximum number of corners per cell
 - `nx_loc` and `ny_loc` are the two dimensions of the local arrays `clon` and `clat`
 - `clon` and `clat` are stored with a Fortran compatible (column major) ordering. If they are declared as triple pointers, they have to be `clon[nc][ny_loc][nx_loc]`
 - `il_partid` is relevant only for parallel writing. In case of single proc invocations, the constant `OASIS_No_Part` can be passed as a placeholder
- `int oasis_c_write_mask(const char* cgrid, const int nx_global, const int ny_global, const int nx_loc, const int ny_loc, const int* mask, const int il_partid, const char* companion)`
 - `nx_global` and `ny_global` are the first and second dimensions of the global grid
 - `nx_loc` and `ny_loc` are the two dimensions of the local array `mask`
 - `mask` is stored with a Fortran compatible (column major) ordering. If it is declared as a double pointer, it has to be `mask[ny_loc][nx_loc]`
 - `il_partid` is relevant only for parallel writing. In case of single proc invocations, the constant `OASIS_No_Part` can be passed as a placeholder
 - if no companion grid attribute is needed, the constant `OASIS_No_Companion` can be passed as a placeholder
- `int oasis_c_write_frac(const char* cgrid, const int nx_global, const int ny_global, const int nx_loc, const int ny_loc, const double* frac,`

- ```
const int il_partid, const char* companion)
```
- `nx_global` and `ny_global` are the first and second dimensions of the global grid
  - `nx_loc` and `ny_loc` are the two dimensions of the local array `frac`
  - `frac` is stored with a Fortran compatible (column major) ordering. If it is declared as a double pointer, it has to be `frac[ny_loc][nx_loc]`
  - `il_partid` is relevant only for parallel writing. In case of single proc invocations, the constant `OASIS_No_Part` can be passed as a placeholder
  - if no companion grid attribute is needed, the constant `OASIS_No_Companion` can be passed as a placeholder
- `int oasis_c_write_area(const char* cgrid, const int nx_global, const int ny_global, const int nx_loc, const int ny_loc, const double* area, const int il_partid)`
    - `nx_global` and `ny_global` are the first and second dimensions of the global grid
    - `nx_loc` and `ny_loc` are the two dimensions of the local array `area`
    - `area` is stored with a Fortran compatible (column major) ordering. If it is declared as a double pointer, it has to be `area[ny_loc][nx_loc]`
    - `il_partid` is relevant only for parallel writing. In case of single proc invocations, the constant `OASIS_No_Part` can be passed as a placeholder
  - `int oasis_c_write_angle(const char* cgrid, const int nx_global, const int ny_global, const int nx_loc, const int ny_loc, const double* angle, const int il_partid)`
    - `nx_global` and `ny_global` are the first and second dimensions of the global grid
    - `nx_loc` and `ny_loc` are the two dimensions of the local array `angle`
    - `angle` is stored with a Fortran compatible (column major) ordering. If it is declared as a double pointer, it has to be `angle[ny_loc][nx_loc]`
    - `il_partid` is relevant only for parallel writing. In case of single proc invocations, the constant `OASIS_No_Part` can be passed as a placeholder
  - `int oasis_c_terminate_grids_writing()`
  - `int oasis_c_def_var(int* var_id, const char* name, const int il_part_id, const bundle_size, const int kinout, const int var_type)`
    - `bundle_size` is a scalar integer, corresponding to the second entry of the Fortran `var_nodims` array
    - `kinout` can receive one of the two predefined constants `OASIS_In` or `OASIS_Out` (also in the form `OASIS_IN` or `OASIS_OUT`)
    - `var_type` can receive one of the two predefined constants `OASIS_Real` or `OASIS_Double` (also in the form `OASIS_REAL` or `OASIS_DOUBLE`)
  - `int oasis_c_enddef()`
  - `int oasis_c_put(const int var_id, const int date, const int x_size, const int y_size, const int bundle_size, const int fkind, const int storage, const void* fld1, const bool write_restart, int* kinfo)`
    - This interface does not support higher order mapping through optional fields at this time.
    - `x_size` and `y_size` are the dimensions of the local portion of the domain (i.e. `fld1`). The order of these dimensions must be the same than the order of the dimensions of the arrays in the `grids.nc` file, which corresponds to the storage order in the corresponding internal

OASIS3-MCT Fortran work arrays.

Notice that for unstructured grids `y_size=1`

- `bundle_size` should be coherent with the argument of same name in the corresponding `oasis_c_def_var`
- `fkind` can take one of the two predefined constants `OASIS_Real` or `OASIS_Double` (also in the form `OASIS_REAL` or `OASIS_DOUBLE`) should be coherent with the argument `ktype` in `oasis_c_def_var`
- `storage` can take one of the two predefined constants `OASIS_COL_MAJOR` or `OASIS_ROW_MAJOR`.

In the first case, the array containing the field has to be declared as

`field[bundle_size][y_size][x_size]` (or any equivalent storage of total size `bundle_size*y_size*x_size` stored in column major order).

In the second case, the field has to be declared as

`field[x_size][y_size][bundle_size]` (or any equivalent row major storage).

Notice that this choice implies an extra memory copy performed internally by OASIS3-MCT before acting on the field and is therefore to be avoided whenever possible.

- `write_restart` can take one of the two predefined constants `OASIS_Write_Restart` or `OASIS_No_Restart`
  - `kinfo` contains a return status to be compared against the values of the `return_codes` enumeration in `oasis_c.h` (equivalent to the returned value of `info` argument for the Fortran `oasis_put` routine, see section 2.2.7)
- `int oasis_c_put_with_frac(const int var_id, const int date, const int x_size, const int y_size, const int bundle_size, const int fkind, const int storage, const void* fld1, const void* fracwgt, const bool write_restart, int* kinfo)`
    - Same as for `oasis_c_put` with the additional `fracwgt` argument, which gives a fraction weight field array (see `oasis_put` in section 2.2.7).
  - `int oasis_c_get(const int var_id, const int date, const int x_size, const int y_size, const int bundle_size, const int fkind, const int storage, void* fld1, int* kinfo)`
    - arguments are the same than for `oasis_c_put`, except that `kinfo` return status is equivalent to the returned value of `info` argument for the Fortran `oasis_get` routine, see section 2.2.7.
  - `int oasis_c_terminate()`
  - `int oasis_c_abort(const int compid, const char* routine_name, const char* abort_message, const char* file, const int line)`
  - `int oasis_c_get_debug(int* debug)`
  - `int oasis_c_set_debug(const int debug)`
  - `int oasis_c_get_intercomm(MPI_Comm* new_comm, char* cdnam)`
    - Returns a C `MPI_Comm` type by reference
  - `int oasis_c_get_intracomm(MPI_Comm* new_comm, char* cdnam)`
    - Returns a C `MPI_Comm` type as by reference
  - `int oasis_c_get_multi_intracomm(MPI_Comm* new_comm, const int cdnam_size, char** cdnam, int* root_ranks)`
    - Returns a C `MPI_Comm` type as by reference

- `cdnam.size` is the size of the array `cdnam`
- `int oasis_c_put_inquire(int var_id, int date, int* kinfo)`
  - `kinfo` contains a return status to be compared against the values of the `return_codes` enumeration in `oasis_c.h`
- `int oasis_c_get_ncpl(const int var_id, int* ncpl)`
- `int oasis_c_get_freqs(const int var_id, const int mop, const int ncpl, int* cpl_freqs)`

## 2.4 OASIS3-MCT python API

The source code of pyOASIS is in the directory `pyoasis/src`. Complete documentation is available in `pyoasis/pyoasis.pdf`. The `pyoasis` interface ultimately call the Fortran version. This is done by wrapping the Fortran in ISO-C bindings (see `lib/cbindings/fortran_isoc`), then wrapping the Fortran ISO-C bindings in C (see `lib/cbindings/c`), then wrapping the C bindings in python (see `pyoasis/src`). This method provides both the C and python bindings. Examples on how to use pyOASIS are provided in `pyoasis/examples`. To run all tests including python, C, and Fortran examples, use `make test`. The python wrapper functions are briefly described next.

- Creating a component using MPI

In pyOASIS, components are instances of the `Component` class. To initialise a component, its name has to be supplied. It is also possible to provide an optional `coupling_flag` argument which defaults to “True”, which means the component is coupled through OASIS3\_MCT.

OASIS3\_MCT couples models which communicate using MPI. If the global communicator at the start of the run is different from the default `MPI_COMM_WORLD` communicator, the global communicator has to be passed to the `Component` class through the third optional argument. By default, the `Component` class will set up MPI internally and provides methods to get access to information such as rank and number of processes in the local communicator gathering only the component processes.

---

```
import pyoasis
[...]
comm = my_global_comm
component_name = "component"
coupling_flag = True
comp = pyoasis.Component(component_name, coupling_flag, comm)
print("Hello world from process " + str(comp.localcomm.rank)
 + " of " + str(comp.localcomm.size))
```

---

To create a coupling communicator for a subset of processes, one can use the method `create_couplcomm`, with a flag being `True` for all these processes; see `pyoasis/examples/4-orange/python` for a practical example.

If such a communicator already exists in the code, it should simply be provided to OASIS3\_MCT with the method `set_couplcomm`; as in `pyoasis/examples/6-apple-and-orange/python`.

To set up an MPI intra communicator or inter communicator between the local component and another component, one can use the methods `get_intracomm` or `get_intercomm`; as in `pyoasis/examples/3-box/python`.

To set up an MPI intra-communicator among some of the coupled components, listed in the `comp_list` list, one can use the method `get_multi_intracomm`, as in `9-python_fortran_C-multi_intracomm`.

Also, the current OASIS3-MCT internal debug level (`$NLOGPRT` value in the *namcouple*), can be retrieved as a property of a component, namely `debug_level`, and can be changed by directly modifying this property, as in `pyoasis/examples/7-multiple-puts/python`.

- Creating a partition

The data can be partitioned in various ways. These correspond to the `SerialPartition`, `ApplePartition`, `BoxPartition`, `OrangePartition` and `PointsPartition` classes which are inherited from the `Partition` abstract class. For details on the different ways to describe the partitions, see section 2.2.3 and examples `1-serial`, `2-apple`, `3-box`, `4-orange`, `5-points` in `pyoasis/examples`.

The simplest situation is the serial partitioning where all the data is held by a single process and only the number of points has to be specified (see example `1-serial`)

In the case of the Apple partitioning, each process contains a segment of a linear domain. To initialise such a partitioning, an offset has to be supplied for each rank as well as the number of data points that will be stored locally (see example `2-apple`).

When we use the Box partitioning, a two-dimensional domain is split into several rectangles. The global offset, local extents in the x and y directions and the global extent in the x direction have to be supplied to the constructor. The global offset is the index of the corner of the local rectangle (see example in `3-box`).

The Orange partitioning consists of several segments of a linear domain (see an example with only one segment per process in `4-orange`.)

The last type of partitioning is Points, where we have to specify, in a list, the global indices of the points stored by the process (see example in `5-points`).

- Defining the coupling grids

The grid data files, containing the definition of the grids onto which the coupling data is defined, can be created by the user before the run or can be written directly at run time by the components, either by one component process to write the whole grid or by each process holding a part of a grid. Details about the grid definition can be found in section 2.2.4. A full example of writing a grid in sequential and parallel models can be found in `examples/10-grid`.

To initialise a grid and write the grid longitudes and latitudes, one has to create an instance of the `Grid` class. Then to write the grid cell corner longitudes and latitudes, areas, mask, cell valid fraction, angle, the `set_corners`, `set_area`, `set_mask`, `set_frac`, and `set_angle` methods can be used respectively.

- Declaring the coupling data

The coupling data is handled by the class `Var`. Its constructor requires its symbolic name, as it appears in the *namcouple* file, the partition and a flag indicating whether the data is incoming or outgoing. The latter is an enumerated type and can have the values `pyoasis.OasisParameters.OASIS_OUT` or `pyoasis.OasisParameters.OASIS_IN`.

The property `is_active` can be tested to check if the variable is activated in the *namcouple* configuring file (see example `3-box/python`).

The coupling period(s) of the data, as defined in the *namcouple*, can be accessed with the property `cpl_freqs` and the number of coupling exchanges in which the data is involved by `len(cpl_freqs)` (see example `7-multiple-puts/python`).

The property `put_inquire` of the variable tells what would happen to the corresponding data at that date below the corresponding send action. This may be useful if, for example, the calculation of a coupling field is costly and if one wants to compute it only when it is really sent out. The different possible return codes are listed in section 2.2.9.

- Ending the definition phase

Then the definition of the component must be ended by calling the `enddef()` method. This must be done only once the partitioning and the variable data have been initialised.

- Sending and receiving data

pyOASIS expects data to be provided as a `pyoasis.asarray` object:

---

```
field = pyoasis.asarray(range(n_points))
```

---

This is a numpy array but ordered in the Fortran way. In C, multidimensional arrays store data in row-major order where contiguous elements are accessed by incrementing the rightmost index while varying the other indices will correspond to increasing strides in memory as we use indices further towards the left. By default, numpy arrays use that ordering as well. Fortran, on the other hand, uses column-major order. In that case, contiguous elements are accessed by incrementing the leftmost index. `pyoasis.asarray` objects use the same ordering as Fortran. As a consequence, it is not necessary to transform data in order to use it in the OASIS3\_MCT Fortran library.

The sending and receiving actions may be called by the component at each timestep. The date argument is automatically analysed and actions are actually performed only if date corresponds to a time for which it should be activated, given the period indicated by the user in the namcouple. See section 2.2.7 for details.

The data is sent with the `put` function.

---

```
date = int(0)
variable.put(date, field)
```

---

Or, if the `write_restart` argument is passed,

---

```
variable.put(date, field, write_restart)
```

---

If the `fracwgt` option is activated, the dynamic fraction field also has to be provided as a `pyoasis.asarray` object and sent together with the field with the `put` function.

---

```
frac = pyoasis.asarray(np.zeros((local_size), dtype=np.float64))
... [fill frac with the dynamic fraction] ...
variable.put(date, field, fracwgt=frac)
```

---

Finally, on the target side, the field is received with the `get` function, which fills also a `pyoasis.asarray` object.

---

```
variable.get(date, field)
```

---

- Termination

Finally, the coupling is terminated with the destruction of the component:

---

```
del comp
```

---

- Exceptions and aborting

When an error occurs in `OASIS3_MCT`, the code coupler returns an error code and an `OasisException` is raised. In practice, `OASIS3_MCT` will internally handle the error, write an error message in its debug log files and to the screen, and abort before the exception is raised. It may also happen that the code aborts before the error message appears on the screen.

When an error is caught by the `pyOASIS` wrapper, such as an incorrect parameter or a wrong argument type, a `PyOasisException` is raised.

In the following example, where we attempt to initialise a component, a `PyOasisException` will be raised as the user supplies an empty name :

---

```
try:
 comp = pyoasis.Component("")
except (pyoasis.PyOasisException) as exception:
 pyoasis.pyoasis_abort(exception)
```

---

The function `pyoasis.pyoasis_abort` takes an exception as argument. It stops the execution of all the processes after having displayed an error message and written information in the log files about the error and the context in which it took place.

Another function is available, `pyoasis.oasis_abort`, for the cases where a voluntary abort is needed in the code when an exception has been raised. Its interface mimics the corresponding `OASIS3_MCT` function `oasis_abort`.

### 2.4.1 Fortran python API correspondence

Figures 2.6 , 2.7 and 2.8 show the Fortran python API correspondence for different parts of the API. Different examples implementing the different parts of the API with the Fortran, C and python interfaces are also provided as practical illustrations in directory `pyoasis/examples` and are described in section 6.3.4.

| Fortran                                                                        | python                                                                                                           | Notes                                                                                                                                                                                                         |
|--------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| use mod_oasis                                                                  | import pyoasis<br>from pyoasis import OASIS                                                                      | The second line is optional and gives access to the short constant syntax, e.g. OASIS.OUT instead of pyoasis.OasisParameters.OASIS_OUT                                                                        |
| CALL oasis_init_comp (compid, comp_name, ierror)                               | comp = pyoasis.Component(name)                                                                                   | Standard form. Notice that <i>comp</i> is a user chosen variable name for an object of the Component class. In the following you can replace <i>comp</i> by the name of the object of your choice             |
| CALL oasis_init_comp (compid, comp_name, ierror, coupled)                      | comp = pyoasis.Component(name, coupled=[True/False])                                                             | Optional argument for actual coupling, default to True                                                                                                                                                        |
| CALL oasis_init_comp (compid, comp_name, ierror, commworld=mycomm)             | comp = pyoasis.Component(name, communicator=mycomm)                                                              | Optional argument for global communicator, default to None, i.e. communicator provided by OASIS. The two optional arguments can be combined                                                                   |
| CALL oasis_get_localcomm (local_comm, ierror )                                 | comp.localcomm                                                                                                   | Local comm accessed as a property of the comp object<br>Notice that upon creation the comp.couplcomm is a duplicate of comp.localcomm                                                                         |
| CALL oasis_create_couplcomm(icpl, local_comm, coupl_comm, kinfo)               | comp.create_couplcomm(icpl)                                                                                      | Process not involved in the coupling set icpl to MPI.UNDEFINED<br>It returns an optional error code, but it sets the comp.couplcomm property.<br>Cf example 4                                                 |
| CALL oasis_set_couplcomm(coupl_comm, kinfo)                                    | comp.set_couplcomm(couplcomm)                                                                                    | Sets the comp.couplcomm property.<br>Cf example 6                                                                                                                                                             |
| CALL oasis def partition (il_part_id, [0,ig_paral(2:)], ierror, ig_size, name) | part = pyoasis.SerialPartition(n_points[, global_size, name])                                                    | ig_paral(1) = 0<br>Optional arguments global_size and name. Cf. sect 2.2.3 of the OASIS3-MCT user guide.<br>Notice that <i>part</i> is a user chosen variable name for an object of the Partition superclass. |
| CALL oasis def partition (il_part_id, [1,ig_paral(2:)], ierror, ig_size, name) | part = pyoasis.ApplePartition(offset, size[, global_size, name])                                                 | ig_paral(1) = 1<br>Optional arguments global_size and name                                                                                                                                                    |
| CALL oasis def partition (il_part_id, [2,ig_paral(2:)], ierror, ig_size, name) | part = pyoasis.BoxPartition(global_offset, local_extent_x, local_extent_y, global_extent_x[, global_size, name]) | ig_paral(1) = 2<br>Optional arguments global_size and name                                                                                                                                                    |
| CALL oasis def partition (il_part_id, [3,ig_paral(2:)], ierror, ig_size, name) | part = pyoasis.OrangePartition(offsets, extents[, global_size, name])                                            | ig_paral(1) = 3<br>Optional arguments global_size and name                                                                                                                                                    |
| CALL oasis def partition (il_part_id, [4,ig_paral(2:)], ierror, ig_size, name) | part = pyoasis.PointsPartition(global_indices[, global_size, name])                                              | ig_paral(1) = 4<br>Optional arguments global_size and name                                                                                                                                                    |

**Figure 2.6:** Fortran python AP correspondence for the initialisation, communication and partition definition.

## 2.5 Additional notes on coupling functionality

### 2.5.1 Coupling scalar values

OASIS3-MCT does not have a distinct feature to support scalar coupling. By scalar coupling, we mean scalar variables such as date and time, logical flags, integer or real parameters, or other scalar data that might be defined identically across all MPI tasks in a component or even just on the root task or a subset of tasks. It is often desirable to communicate scalar data between components to coordinate scientific or technical features. There isn't a feature that supports this type of coupling specifically, but it is possible to do so using available interfaces. The most robust implementation is to setup root to root coupling of scalars between components. To do so :

- Determine the number and type of scalars to couple
- Allocate an array in each component of that size
- Initialize a partition using the POINTS approach (see section 2.2.3) with the size of the array assigned to the root process in each component and no portion of the partition allocated to other processes.
- Define coupling field names for use in both the model components and the namcouple file and declare variables as usual. Create a set of namcouple inputs without any mapping.

| Fortran                                                                           | python                                                                  | Notes                                                                                                                                                                                                                                                                                                                                                                                                  |
|-----------------------------------------------------------------------------------|-------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| CALL oasis_start_grids_writing (flag)                                             |                                                                         | Implicitly in the pyoasis.Grid class constructor for the first instantiated grid                                                                                                                                                                                                                                                                                                                       |
| CALL oasis_write_grid (cgrid, nx_global, ny_global, lon, lat, il_part_id)         | grid = pyoasis.Grid(cgrid, nx_global, ny_global, lon, lat[, partition]) | Notice that <i>grid</i> is a user chosen variable name for each object of the Grid class<br>lon and lat are entered as 2D arrays - for unstructured grids the second dimension has to be 1 - of the same shape (nx_loc, ny_loc) where nx_loc, ny_loc are the number of points in each dimension on the calling process.<br>The optional partition argument is a member of the pyoasis.Partition class. |
| CALL oasis_write_corner (cgrid, nx_global, ny_global, nc, clon, clat, il_part_id) | grid.set_corners(clo, cla)                                              | clo, cla are 3d arrays: the last dimension is the max. number of corners per cell, the two others are nx_loc, ny_loc.                                                                                                                                                                                                                                                                                  |
| CALL oasis_write_mask (cgrid, nx_global, ny_global, mask, il_part_id, companion)  | grid.set_mask(mask[, companion])                                        | mask is a 2d integer array of size nx_loc, ny_loc.<br>Oasis convention: 1 for masked points, 0 elsewhere.<br>The optional argument companion is the name of a companion grid.                                                                                                                                                                                                                          |
| CALL oasis_write_frac (cgrid, nx_global, ny_global, frac, il_part_id, companion)  | grid.set_frac(frac[, companion])                                        | frac is a 2d array of size nx_loc, ny_loc.<br>The optional argument companion is the name of a companion grid.                                                                                                                                                                                                                                                                                         |
| CALL oasis_write_area (cgrid, nx_global, ny_global, area, il_part_id)             | grid.set_area(area)                                                     | area is a 2d array of size nx_loc, ny_loc.                                                                                                                                                                                                                                                                                                                                                             |
|                                                                                   | grid.write()                                                            | Terminates the declarations for the grid <i>grid</i>                                                                                                                                                                                                                                                                                                                                                   |
| CALL oasis_terminate_grids_writing ()                                             |                                                                         | Implicitly activated by the grid.write() of the last grid instance to call it                                                                                                                                                                                                                                                                                                                          |
| CALL oasis_def_var (var_id, name, il_part_id, [1,1], kinout, var_type, ierror)    | var = pyoasis.Var(name, partition, inout)                               | partition is a pyoasis.Partition object as in previous section.<br>inout is an enumerate parameter, either in the long form pyoasis.OasisParameters.OASIS_OUT/OASIS_IN or in the short form (cf Module) OASIS.OUT/OASIS.IN<br>Notice that the var_type is left to the dtype of the exchanged field.<br>Once again <i>var</i> is a user chosen variable name for an object of the Var class.            |
| CALL oasis_def_var (var_id, name, il_part_id, [1,n], kinout, var_type, ierror)    | var = pyoasis.Var(name, partition, inout[, bundle_size=n])              | The size of the bundle is set via the optional argument bundle_size                                                                                                                                                                                                                                                                                                                                    |
| CALL oasis_enddef (ierror)                                                        | comp.enddef()                                                           | Notice that it is a method of the comp object, comp being the user chosen variable name of the component                                                                                                                                                                                                                                                                                               |

**Figure 2.7:** Fortran python AP correspondence for the grid definition, variable declaration and end of definition phase.

- Use the `oasis_get` and `oasis_put` interfaces to communicate data between components.
- If required, broadcast the scalars after being received from the root process to the other processes, outside the OASIS3-MCT layer.

This can be extended as needed to send or receive from non root processes. But one cannot send the scalar data to multiple processes as this violates the halo restriction in MCT.

### 2.5.2 The lag concept

Using the OASIS3-MCT coupling library, the user has the flexibility to reproduce different coupling algorithms defining LAG values for the different coupling fields. See section 3.3 for more details on how to specify a LAG for a coupling field. In the components, the sending and receiving routines, respectively `oasis_put` and `oasis_get`, can be called at each component timestep, with the appropriate `date` argument giving the actual time (at the beginning of the timestep), expressed in number of seconds since the start of the run, or in any other time units as long as the same are used in all components and in the *namcouple* (see section 2.2.7). This `date` argument is automatically analysed by the coupling library and, depending on the coupling period and the lag chosen by the user for the coupling field in the *namcouple* (LAG), different coupling algorithms can be reproduced without modifying the component codes themselves.

The lag (LAG) must be expressed in the time unit used and can be positive or negative but should never be larger (in absolute magnitude) than the coupling period of any field due to problems with restartability and dead-locking. When a component calls a `oasis_put`, the value of the lag is automatically added to the

| Category                                                                               | Fortran                                                                                                                         | python                                                                                     | Notes                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          |
|----------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Put and Get (2.2.7)                                                                    |                                                                                                                                 | <code>field=pyoasis.asarray(field[, dtype])</code>                                         | Make sure that field is a numpy asfortran array of the required dtype (optional argument defaulting to <code>numpy.float64</code> )<br>The call can be skipped if field is already so                                                                                                                                                                                                                                                                                                                          |
|                                                                                        | <code>CALL oasis_put (var_id, date, fld1, info, fld2, fld3, fld4, fld5, write_restart, fracwgt)</code>                          | <code>var.put(time, field[, write_restart][, fracwgt])</code>                              | var is a previously defined object of the Var class<br>field, and possibly fracwgt, have been checked by <code>pyoasis.asarray</code><br>Bundle are supported (bundle index is the last index)<br>Bicubic and conservative 2nd extra fields not supported.<br>write_restart boolean optional argument for triggering a NetCDF dump of the sent field.<br>fracwgt is an optional dynamic fraction field.<br>Returned value is to be compared against the Oasis Parameters enumeration (in long or short syntax) |
|                                                                                        | <code>CALL oasis_get (var_id, date, fld, info)</code>                                                                           | <code>var.get(time, field)</code>                                                          | field has to be preallocated and checked by <code>pyoasis.asarray</code><br>Returned value is to be compared against the Oasis Parameters enumeration (in long or short syntax)                                                                                                                                                                                                                                                                                                                                |
| Termination (2.2.8)                                                                    | <code>CALL oasis_terminate (ierror)</code>                                                                                      | <code>del comp / nothing</code>                                                            | oasis terminates at the comp object destruction, either explicitly by <code>del comp</code> or automatically at garbage collection on exit                                                                                                                                                                                                                                                                                                                                                                     |
| Auxiliary routines (2.2.9)                                                             | <code>CALL oasis_get_debug(debug_value)</code>                                                                                  | <code>comp.debug_level</code><br><code>pyoasis.get_debug()</code>                          | Coded as a property of a Component object or as a standalone function                                                                                                                                                                                                                                                                                                                                                                                                                                          |
|                                                                                        | <code>CALL oasis_set_debug(debug_value)</code>                                                                                  | <code>comp.debug_level = debug_value</code><br><code>pyoasis.set_debug(debug_value)</code> | Coded as a setter for a Component object property or as a standalone function                                                                                                                                                                                                                                                                                                                                                                                                                                  |
|                                                                                        | <code>CALL oasis_put_inquire(var_id, date, kinfo)</code>                                                                        | <code>var.put_inquire(time)</code>                                                         | var is a Var object. Returned value is to be compared against the Oasis Parameters enumeration (in long or short syntax)                                                                                                                                                                                                                                                                                                                                                                                       |
|                                                                                        | <code>CALL oasis_get_ncpl(var_id, ncpl, kinfo)</code><br><code>CALL oasis_get_freqs(var_id, mop, ncpl, cpl_freqs, kinfo)</code> | <code>var.cpl_freqs</code>                                                                 | Coded as a property of a Var object. Returns an iterable list                                                                                                                                                                                                                                                                                                                                                                                                                                                  |
|                                                                                        | <code>CALL oasis_get_intercomm(new_comm, cdnam, kinfo)</code>                                                                   | <code>comp.get_intercomm(comp_name)</code>                                                 | Coded as a method of a Component object.                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |
|                                                                                        | <code>CALL oasis_get_intracomm(new_comm, cdnam, kinfo)</code>                                                                   | <code>comp.get_intracomm(comp_name)</code>                                                 | Coded as a method of a Component object.                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |
|                                                                                        | <code>CALL oasis_get_multi_intracomm(new_comm, cdnam, root_ranks, kinfo)</code>                                                 | <code>comp.get_multi_intracomm(complist)</code>                                            | Coded as a method of a Component object.<br>Returns a pair of values: the new comm and a dictionary associating root ranks (values) to component names (keys)                                                                                                                                                                                                                                                                                                                                                  |
| <code>CALL oasis_abort (compid, routine_name, abort_message, file, line, rcode)</code> | <code>raise PyOasisException(text)</code><br><code>raise OasisException(text, error)</code>                                     | Preferred use of exceptions over an abort function                                         |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |

**Figure 2.8:** Fortran python AP correspondence for the coupling field exchanges, termination and auxiliary routines.

value of the `date` argument and the “put” is actually performed when the sum `date+lag` is a coupling time. In the target component, this “put” will match a `oasis_get` for which the `date` argument is that same coupling time. So the lag only shifts the time at which the data is sent but not the time at which the data is received.

When the lag is positive, a restart file must be available to initiate the coupling. See section 5.2 for details on the structure of the coupling restart files. For a field with positive lag, the source component automatically reads the field in the restart file during the coupling initialization phase (below the `oasis_undef`) and send the data to match the `oasis_get` performed at `time=0` in the target component. The final coupling data on the source side will then be automatically written to the restart file for use in the next run<sup>4</sup>.

On figure 2.9, short black arrows correspond to `oasis_put` or `oasis_get` called in the component that do not lead to any “put” or receiving action; long black arrows correspond to `oasis_put` or `oasis_get` called in the components that do actually lead to a “put” or “get” action; long red arrows correspond to `oasis_put` or `oasis_get` called in the component models that lead to a reading or writing of the coupling field from or to a coupling restart file.

A coupling algorithm exploiting the LAG concept, is illustrated on figure 2.9.

During a coupling timestep, model A receives  $F_2$  and then sends  $F_1$ ; its timestep length is 4. During a coupling timestep, model B receives  $F_1$  and then sends  $F_2$ ; its timestep length is 6.  $F_1$  and  $F_2$  coupling periods are respectively 12 and 24. If  $F_1/F_2$  “put” action by model A/B was used at a coupling timestep to match the model B/A “get” action, a deadlock would occur as both models would be initially waiting on a “get” action. To prevent this,  $F_1$  and  $F_2$  produced at the timestep before have to be used to match respectively the model B and model A “get” actions.

<sup>4</sup>When there is a lag, the first and last instance of the source field in the debug netCDF file (EXPOUT fields, see section 3.3) always correspond respectively to the field read from and written to the restart file.

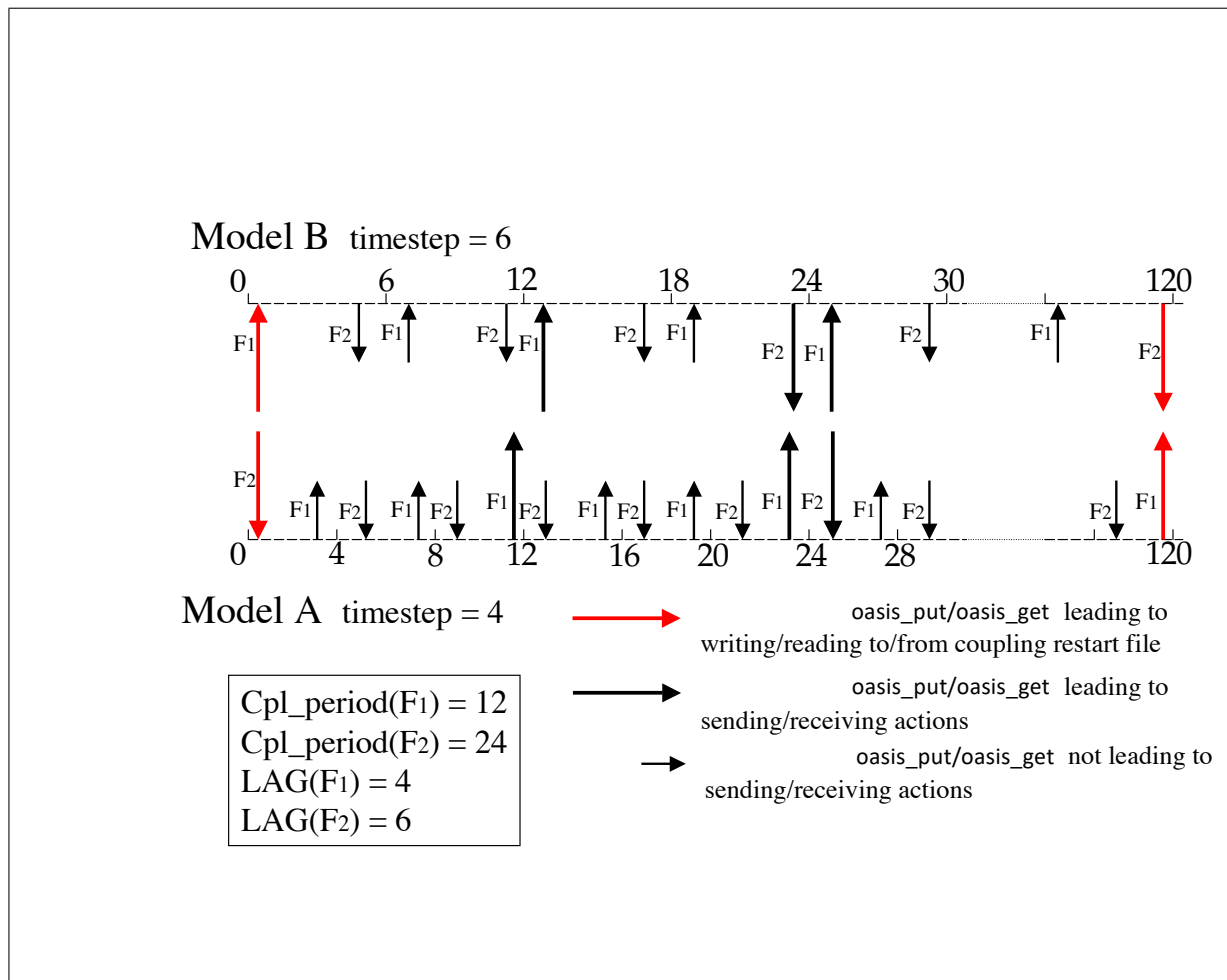


Figure 2.9: LAG concept first example

This implies that a lag of respectively 4 and 6 seconds must be defined for  $F_1$  and  $F_2$ . For  $F_1$ , the `oasis_put` performed at time 8 and 20 by model A will then lead to “put” actions (as  $8 + 4 = 12$  and  $20 + 4 = 24$  which are coupling periods) that match the “get” actions performed by `oasis_get` called by model B at times 12 and 24. For  $F_2$ , the `oasis_put` performed at time 18 by model B then leads to a “put” action (as  $18 + 6 = 24$  which is a coupling period) that matches the “get” action performed at time 24 by the `oasis_get` called by model A.

At the beginning of the run, as their LAG index is greater than 0, the first `oasis_get` of  $F_1$  and  $F_2$  will automatically be fulfilled with fields read from their respective coupling restart files. The user therefore has to create those coupling restart files before the first run in the experiment. At the end of the run,  $F_1$  having a lag greater than 0, is automatically written to its coupling restart file below the last  $F_1$  `oasis_put` as the `date+lag` equals the total run time. The analogue is true for  $F_2$ . These coupling restart fields will automatically be read in at the beginning of the next run below the respective `oasis_get`.

In conclusion, using appropriate LAG indices ensures that the target component receives the coupling fields produced by the source component at a previous time. This is, in some coupling configurations, essential to avoid deadlock situations.

### 2.5.3 The sequence concept

The order of coupling operations in the system is determined solely by the order of calls to send (`oasis_put` or “put”) and receive (`oasis_get` or “get”) data in the components in conjunction with the setting of the LAG in the *namcouple*. Receiving data is always blocking while sending data is non-blocking with respect to the component making that call. It is possible to deadlock the system if the relative orders of puts and gets in different components are not compatible.

The sequence (SEQ) index in the *namcouple* file now provides the coupling layer with an ability to detect a deadlock before it happens and exit. It does this by tracking the order of get and put calls in components compared to the SEQ specified in the *namcouple*. If there are any inconsistencies, the component will abort gracefully with a useable error message before the system deadlocks. If there are any coupling dependencies in the system, use of the SEQ index is recommended for diagnosis but has no impact on the ultimate solution and is NOT required.

In the following two examples, there are two models, each one sending a field to the other at every coupling period without any lags. In the first case, there is no dependency as each model first sends and then receives some data.

```

model1 model2

put (fld1) put (fld2)
get (fld2) get (fld1)

```

In this case, there is no sequencing dependency and specifying SEQ indices is not needed<sup>5</sup>.

In the next example, there is a dependency in the sequencing.

```

model1 model2

put (fld1) get (fld1)
 fld2=g (fld1)
get (fld2) put (fld2)

```

In model2, fld2 depends on fld1. If SEQ is not used and if, for example, model1 does not have the consistent ordering of the put and get shown above (required by model2), then the models would deadlock and hang. If this dependency is known, there is a benefit in setting SEQ=1 for fld1 and SEQ=2 for fld2;

<sup>5</sup>In that case, if set, the SEQ indices must be identical. If by mistake, SEQ is set to 1 for fld1 and 2 for fld2, then the coupled model will abort because at runtime, the coupler will detect in model 2 that fld2 was sent before fld1 was received which is out of sequence as defined by the SEQ settings.

at runtime. If the sequencing of `model1` or `model2` does not match the above diagram, then the coupling layer will detect it and will exit gracefully with an error message.

Again, the `SEQ` namecouple setting is only diagnostic and is not required.

#### 2.5.4 A brief overview of MCT

OASIS3-MCT leverages the MCT coupling library developed at Argonne National Laboratory. That library is designed to couple fields on static grids between model components. The fields can be decomposed across multiple processes, and the decomposition and number of processes involved can be arbitrary in each component.

MCT also provides the ability to map (i.e. interpolate or regrid) data between grids as long as the interpolation is linear and can be computed by a linear sparse matrix multiply. MCT does not compute interpolation weights, but it has interfaces that allow those weights to be passed into MCT. Within MCT, mapping and communication are also treated as independent features.

OASIS3-MCT supports both mapping and communication of data through MCT. A coupling field can be interpolated to the destination grid on the source processes then communicated to the destination component; or a coupling field can be communicated to the destination component on the source grid then mapped to the destination grid on the destination component (see `$MAPLOC` in section 4.3). At the present time, it is not possible to map the field as part of the communication rearrangement, although that capability should be possible to implement. The separation of mapping and communication is handled by the OASIS layer. Users only need to be aware of a few options that can be set to fine tune the performance of these operations.

OASIS3-MCT also imposes a few other constraints on the usage. For example, MCT does not support haloed communication. The partitions defined by the component that describes the field decomposition cannot reference the same global grid cell more than once.

Also, neither OASIS3-MCT nor MCT support dynamically varying grids nor dynamically varying decompositions at the present time.

Finally, MCT itself, and therefore the communication layer in OASIS, has no OpenMP support. Users are discouraged from calling OASIS3-MCT `oasis_put` and `oasis_get` from a threaded region. And it is not possible to define a partition (decomposition) across multiple active threads. The work that OASIS3-MCT does related to decomposition rearrangement for communication or mapping per se is MPI-based. There are some floating point operations in the sparse matrix multiply as well as in some of the OASIS3-MCT transformation operations; but generally, these are highly parallel and would often not benefit from use of OpenMP. The only use of OpenMP in OASIS3-MCT is within the mixed MPI+OpenMP parallel version of the SCRIP library for the calculation of the remapping weights (see section 4.3).

## Chapter 3

# The configuration file *namcouple*

The OASIS3-MCT configuration file *namcouple* contains, below pre-defined keywords, all user-defined information necessary to configure a particular coupled run.

The *namcouple* is a text file with the following characteristics:

- the keywords used to separate the information can appear in any order;
- the number of blanks between two character strings is non-significant;
- all lines beginning with # are ignored and considered as comments;
- blank lines are supported (since OASIS3-MCT\_4.0).

The first part of *namcouple* is devoted to configuration of general parameters such as the total run time or the desired debug level. The second part gathers specific information on each coupling (or I/O) field, e.g. their coupling period, the list of transformations or remapping to be performed by OASIS3-MCT and associated configuring lines (described in more details in chapter 4).

Compared to OASIS3, several *namcouple* inputs have been deprecated in OASIS3-MCT (i.e. they will not be read and not be used) but, for backwards compatibility, they can still appear in the *namcouple*. These inputs will be noted “UNUSED” in the following text and will not be fully described.

In the next sections, a *namcouple* example is given and all configuring parameters are described. Additional lines containing different parameters for each transformation are described in chapter 4. A realistic *namcouple* can be found in `oasis3-mct/examples/` sub-directories.

### 3.1 An example of a simple *namcouple*

The following simple *namcouple* configures a run into which an ocean and an atmosphere components are coupled. The ocean running on grid `toce` provides only the `SOSSTSST` field to the atmosphere (grid `atmo`), which in return provides the field `CONSFTOT` to the ocean. The field `CONSFTOT` is also explicitly written out to a file `flda3.nc` and one field `SOALBEDO` is read from a file.

```
First section
$NFIELDS
 4
#
$RUNTIME
 432000
#
$NLOGPRT
 2 1 0
#
```

```

$NCDFTYP
 cdf1
#
$NUNITNO
 901 920
#
$NMAPDEC
 decomp_wghtfile
#
$NMATXRD
 ceg
#
$NWGTOPT
 ignore_bad_index
#
$SEQMODE
$CHANNEL
$JOBNAME
$NBMODEL
$INIDATE
$MODINFO
$SCALTYPE
#
Second section
#
$STRINGS
#
Field 1
SOSSTSST SISUTESU 1 86400 5 sstoc.nc EXPOUT
182 149 128 64 toce atmo LAG=+14400 SEQ=+1
P 2 P 0
LOCTRANS CHECKIN MAPPING BLASNEW CHECKOUT
#
 AVERAGE
 INT=1
 map_toce_atmo_120315.nc src opt
 1.0 1
 CONSTANT 273.15
 INT=1
#
Field 2
CONSFTOT SOHEFLDO 6 86400 4 flxat.nc EXPORTED
atmo toce LAG=+14400 SEQ=+2
P 0 P 2
LOCTRANS CHECKIN SCRIPR CHECKOUT
#
 ACCUMUL
 INT=1
 BILINEAR LR SCALAR LATLON 1
 INT=1
#

```

```

Field 3
CONSFTOT CONSFTOT 1 86400 1 flda3.nc OUTPUT
128 64 128 64 atmo atmo
LOCTRANS
AVERAGE
#
Field 4
SOALBEDO SOALBEDO 17 86400 0 SOALBEDO.nc INPUT

```

## 3.2 First section of *namcouple* file

The first section of *namcouple* uses some predefined keywords prefixed by the \$ sign to locate the related information. The \$ sign must be the first non-blank character on the line but can be in any column. 8 keywords are used by OASIS3-MCT and 5 of these are optional :

- **\$NFIELDS:** On the line below this keyword, put a number equal (or greater) to the total number of field entries in the second part of the *namcouple*. If more than one field are described on the same line, this counts as only one entry.
- **\$RUNTIME:** On the line below this keyword, put the total simulated time of the run, expressed in seconds (or any other time units as long as they are the same in all components and in the *namcouple*, see 2.2.7). Note that by convention the first coupling of a run occurs at the beginning of the run and all other couplings occur at a time strictly smaller than \$RUNTIME.
- **\$NLOGPRT:** The first, second and third numbers on the line below this keyword refer to (i) the debug verbosity, (ii) internal timing statistics and (iii) the load balancing analysis. The information is written by OASIS3-MCT for each component and (optionnally) for each process.

The first number (that can be modified at runtime with the `oasis_set_debug` routine, see section 2.2.9) may be:

- 0 : production mode. One file `debug.root.xx` is open by the master process of each component and one file `debug_notroot.xx` is open for all the other processes of each component to write only error information.
- 1 : one file `debug.root.xx` is open by the master process of each component to write information equivalent to level 10 (see below) and memory usage information; one file `debug_notroot.xx` is open for all the other processes of each component to write error information.
- 2 : one file `debug.yy.xxxxxx` is open by each process of each component (with `yy` being the component number and `xxxxxx` the process number) to write normal production diagnostics and memory usage information
- 5 : as for 2, and some initial debug info
- 10: as for 5, and the routine calling tree
- 12: as for 10, and some routine calling notes
- 15: as for 12, and even more debug diagnostics
- 20: as for 15, and some extra runtime analysis
- 30: full debug information

The second number defines if time statistics are written out to file `comp_name.timers_xxxx` (with `comp_name` being the component name, see section 2.2.2); it can be:

- 0 : nothing is calculated or written.
- 1 : some time statistics are calculated and written in a single file by the processor 0 as well as the min and the max times over all the processors.

- 2 : some time statistics are calculated and each processor writes its own file ; processor 0 also writes the min and the max times over all the processors in its file.
- 3 : some time statistics are calculated and each processor writes its own file ; processor 0 also writes in its file the min and the max times over all processors and also writes in its file all the results for each processor.

For more information on the time statistics written out, see section 6.4.2.

The third number can be set to 1 to activate a **load balancing** diagnostic. An efficient use of the allocated computing resources in a coupled system requires the harmonisation of the components speed. This operation, called load balancing, is often neglected, either because of the apparent resource abundance and practical difficulties. To facilitate this work, OASIS3-MCT can output the full timeline of all coupling related events, for any of the allocated resources. This timeline is saved in one netCDF file per coupled component (`timeline_XXX_component.nc`). It provides the comprehensive sequence of any operations related to the coupling (field exchange through MPI, field output on disk, field interpolation and mapping, field reading on disk, restart writing, initialisation and termination phase of the OASIS3-MCT setup) so that any simulation slow down in link with the use of the coupler can be identified. The analysis of the coupling field exchanges, amongst all the coupling events, allows not only to identify the resources waste of components which are recurrently waiting for their coupling fields but it also reveals other bottlenecks such as disk access, OS interruptions or model internal load imbalance. The full picture of these events makes possible an optimum load balancing, even for the most complex configurations. For a detailed information on load balancing analysis and timeline visualisation see respectively [Maisonnavé et al 2020] and in [Piacentini and Maisonnavé 2020]. In addition to the timeline, computing information (time to solution, speed, cost) and a synthesis of the time spent on MPI routines for each coupled component can also help, in the simpler cases, to allocate resources in a balanced way ( see file `load_balancing_info.txt` ).

- `$NCDF_TYP`: Optional; on the line below this keyword is a character string that indicates the NetCDF file type for all (i.e. mapping, restart, output) new NetCDF files generated by OASIS3-MCT<sup>1</sup>. Options are `cdf1`, `cdf2`, and `cdf5`. The mode `cdf1`, which is the default, is also known as classic mode, `cdf2` as large file format or `64bit_offset` and supports larger files, `cdf5` as `64bit_data` and supports both larger files and larger variables. More information about these file types can be found in the NetCDF documentation. The file format for any NetCDF file can be diagnosed by using `ncdump -k filename'`. `$NCDF_TYP` only affects new files created by OASIS3-MCT.
- `$NUNITNO`: Optional; on the line below this keyword are two integers that indicate the minimum and maximum unit numbers to be used for input and output files in the coupling layer. The user should choose values that will NOT conflict or overlap with unit numbers in use in any of the component models. The defaults are 1024 for the minimum and 9999 for the maximum unit number if not explicitly set by the user.
- `$NMAPDEC`: Optional; on the line below this keyword is a character string that indicates the mapping decomposition value to be used during local mapping of the source/target field on the target/source component processes during the remapping. The options are `decomp_ld` and `decomp_wghtfile`. Option `decomp_ld` decomposes the field in a simple one dimensional way while `decomp_wghtfile` decomposes the field using the information in the remapping weight file to reduced mapping communication. Option `decomp_wghtfile` will take some extra time in initialization but it should result in faster mapping. The default is `decomp_ld` but it is recommended to test `decomp_wghtfile` to see if it improves performance. More details can be found in [Craig et al 2018] and in [Valcke et al 2018].
- `$NMATXRD`: Optional; on the line below this keyword is a character string that indicates the method used to read remapping weights. There are two options, `orig` and `ceg`. In both, the weights

<sup>1</sup>NetCDF will read and/or overwrite existing files of any NetCDF file type, and the file type will remain unchanged in that case.

are read in chunks by the root process. In the `orig` option, the weights are then broadcasted to all processes and each process then saves the weights needed in order to be consistent with the mapping decomposition. In the `ceg` option, the root process reads the weights and then decides which process each weight should be assigned to. A series of exchanges are then done and just the weights needed on each process are sent. The `orig` method sends much more data but is more parallel. The `ceg` method does most of the work on the root process but less data is communicated. The default option is `ceg`. More details can be found in [Craig et al 2018].

- `$NWGTOPT` : Optional; on the line below this keyword is a character string that indicates how to handle bad remapping weights. There are four options `abort_on_bad_index`, `ignore_bad_index`, `ignore_bad_index_silently`, and `use_bad_index`. Bad weights are defined as weights in the mapping file for which either the source or destination index are out of bounds relative to the number of grid cells in the grid; in that case, the weight is referencing a gridcell that does not physically exist. Note that an index equal to zero will not be considered as a bad index if the associated weight is also zero. There are other situations where the value of the actual mapping weight is scientifically incorrect, but this is not easy to detect and is not dealt with in OASIS3-MCT.
  - `abort_on_bad_index` will write error messages to the log files and abort if a bad weight index is detected. This is the default option.
  - `ignore_bad_index` will write an error message and then remove bad weights internally before continuing.
  - `ignore_bad_index_silently` will remove bad weights and continue without writing an error message.
  - `use_bad_index` will attempt to keep bad weights in the interpolation computation, but this can result in memory corruption, silent dropping of weights, and incorrect results ; this is not recommended.

Note that the ability to check mapping files at runtime in OASIS3-MCT is limited. It is always recommended that mapping files be analyzed offline before long production runs are carried out. Checks can be done to make sure the source and destination indices are valid, that weights values are reasonable (for instance, between 0 and 1, although this will depend on the mapping method), and that the sum of weights on the destination cells are reasonable (for instance, 1, in many cases). In addition, offline tests can be run with analytical functions to verify conservation, gradient preserving features and other characteristics associated with the particular mapping approach.

- `$NNOREST`: Optional; on the line below this keyword is a character string that can override the requirement that restart files must exist if they are needed. If the character string value starts with `T`, `t`, `.T`, or `.t` (as in `true`), then OASIS3-MCT will initialise with zero any variable that normally requires a restart (for instance, variables with `LAG > 0`) if the restart file does not exist. By default, or if `$NNOREST` is not `true`, missing restart files will cause the model to abort. It is strongly recommended that this keyword **NOT** be used in production runs. It exists to provide a quick shortcut for running technical tests. Note that if `$NNOREST` is `true` but the restart file nonetheless exists, it will be used.
- Keywords `$SEQMODE`, `$CHANNEL`, `$JOBNAME`, `$NBMODEL`, `$INIDATE`, `$MODINFO`, `$CALTYPE` are not used anymore.

### 3.3 Second section of *namcouple* file

The second part of the *namcouple*, starting after the keyword `$STRINGS`, contains coupling information for each coupling (or I/O) field. Its format depends on the field status given by the last entry on the field

first line (EXPORTED, IGNOUT or INPUT in the example above). The field may be<sup>2</sup> :

- EXPORTED (or IGNORED): exchanged by OASIS3-MCT
- EXPOUT (or IGNOUT): exchanged and also written to two debug NetCDF files, one before the sending action in the source component below the `oasis_put` call (after local transformations LOCTRANS and BLASOLD if present), and one after the receiving action in the target component below the `oasis_get` call (after all transformations). EXPOUT should be used only when debugging the coupled model. The name of the debug NetCDF file (one per field) is automatically defined based on the field and component names.
- INPUT: read in from the input file by the target component below the `oasis_get` call at appropriate times corresponding to the input period indicated by the user in the *namcouple*. See section 5.3 for the format of the input file.
- OUTPUT: written out to an output debug NetCDF file by the source component below the `oasis_put` call, after local transformations LOCTRANS and BLASOLD, at appropriate times corresponding to the period indicated by the user in the *namcouple*.

### 3.3.1 Second section of *namcouple* for EXPORTED and EXPOUT fields

The first 4 lines for fields with status EXPORTED and EXPOUT are as follows:

```
SOSSTSST SISUTESU 1 86400 3 sstoc.nc EXPOUT
182 149 128 64 toce atmo LAG=+14400 SEQ=+1
P 2 P 0
CHECKIN YAC CHECKOUT
```

where the different entries are:

- Field first line:
  - SOSSTSST : symbolic name for the field in the source component (80 characters maximum). It has to match the argument name of the corresponding field declaration in the source component; see `oasis_def_var` in section 2.2.5
  - SISUTESU : symbolic name for the field in the target component (80 characters maximum). It has to match the argument name of the corresponding field declaration in the target component; see `oasis_def_var` in section 2.2.5
  - 1 : UNUSED but still required for parsing
  - 86400 : coupling or I/O period for the field, in seconds or any other units as long as they are the same in all components and in the *namcouple*
  - 3 : number of transformations to be performed by OASIS3-MCT on this field, described on the fourth line
  - sstoc.nc : name of the coupling restart file for the field (32 characters maximum); mandatory even if no coupling restart file is effectively used (for more detail, see section 5.2)
  - EXPOUT : field status, see above
- Field second line:
  - 182 : number of points for the source grid first dimension (optional)
  - 149 : number of points for the source grid second dimension (optional)<sup>3</sup>
  - 128 : number of points for the target grid first dimension (optional)
  - 64 : number of points for the target grid second dimension (optional)<sup>3</sup>

<sup>2</sup>The status AUXILARY is not supported anymore.

<sup>3</sup>For 1D field, put 1 as the second dimension

These source and target grid dimensions are optional but have to be provided in order to have 2D fields written as 2D arrays in the debug files; otherwise, the fields will be written out as 1D arrays.

- `toce` : prefix of the source grid name in grid data files (see section 5.1) (80 characters maximum)
- `atmo` : prefix of the target grid name in grid data files (80 characters maximum)
- `LAG=+14400`: optional lag index for the field (see section 2.5.2)
- `SEQ=+1`: optional sequence index for the field (see section 2.5.3)
- Field third line
  - `P` : source grid first dimension characteristic (`'P'`: periodical; `'R'`: regional).
  - `2` : source grid first dimension number of overlapping grid points.
  - `P` : target grid first dimension characteristic (`'P'`: periodical; `'R'`: regional).
  - `0` : target grid first dimension number of overlapping grid points.
- The fourth line gives the list of transformations to be performed for this field. Choices are `LOCTRANS`, `CHECKIN`, `BLASOLD`, `MAPPING`, `YAC`, `SCRIPR`, `CONSERV`, `BLASNEW`, `CHECKOUT`. In addition, there is one or more configuring lines describing some parameters for each transformation. These transformations and configuring lines are described in chapter 4.

### Support to group multiple fields in a single communication

With *OASIS3-MCT*, it is possible to group multiple fields in a single communication. To activate this option, the user must list the related fields on a single entry line (with a maximum of 5000 characters on one line) through a colon delimited list in the *namcouple*, for example:

```
ATMTAUX:ATMTAUY:ATMHFLUX TAUX:TAUY:HEATFLUX 1 3600 3 rstrt.nc EXPORTED
```

All fields in this group will then use the same *namcouple* settings (source and target grids, transformations, etc.) for that entry. In the component model codes, these fields are still apparently sent or received one at a time through individual `oasis_put` and `oasis_get`. Inside *OASIS3-MCT*, the fields in this group are stored and a single mapping and send or receive instruction is executed for all fields. This is useful in cases where multiple fields have the same coupling transformations to reduce communication costs by aggregating multiple fields into a single communication.

This option does not put any constraint on the order of the related `oasis_put` and `oasis_get` in the codes. This option is compatible with the special case of 3D unbundled fields and it is particularly well suited for lowering the latency of the communications of such large objects.

As they appear in one single entry line, these fields must share the same coupling restart file but this restart file may contain other fields.

### 3.3.2 Second section of *namcouple* for OUTPUT fields

The first 3 lines for fields with status `OUTPUT` are as follows:

```
CONSFTOT CONSFTOT 1 86400 1 flda3.nc OUTPUT
128 64 128 64 atmo atmo LAG=+14400 SEQ=+1
LOCTRANS
```

where the different entries are:

- Field first line:
  - the source symbolic name must be repeated twice on the field first line
  - `1` : UNUSED but still required for parsing
  - `86400` : output period for the field, in seconds or any other units as long as they are the same in all components and in the *namcouple*
  - `1` : number of transformations to be performed by *OASIS3-MCT* on this field

- flda3.nc : name of the coupling restart file for the field (32 characters maximum); needed only if a LOCTRANS transformation is present
- OUTPUT : field status
- Field second line:
  - 128 : number of points for the source grid first dimension (optional)
  - 64 : number of points for the source grid second dimension (optional)<sup>4</sup>
  - 128 : number of points for the target grid first dimension (optional)
  - 64 : number of points for the target grid second dimension (optional)<sup>4</sup>
  - atmo : prefix of the source grid name in grid data files repeated twice (80 characters maximum)
  - LAG=+14400: optional lag index for the field (see section 2.5.2)
  - SEQ=+1: optional sequence index for the field (see section 2.5.3)
- The third line is LOCTRANS if this transformation is chosen for the field. Note that LOCTRANS is the only transformation supported for OUTPUT fields.

### 3.3.3 Second section of *namcouple* for INPUT fields

The first and only line for fields with status INPUT is:

```
SOALBEDO SOALBEDO 1 86400 0 SOALBEDO.nc INPUT
```

where the different entries are:

- SOALBEDO: symbolic name for the field in the target component (80 characters maximum, repeated twice)
- 1: UNUSED but still required for parsing (as for EXPORTED fields above)
- 86400: input period in seconds or any other units as long as they are the same in all components and in the *namcouple*
- 0: number of transformations (always 0 for INPUT fields)
- SOALBEDO.nc: the input file name (32 characters maximum) (for more detail on its format, see section 5.3)
- INPUT: field status.

---

<sup>4</sup>For 1D field, put 1 as the second dimension

## Chapter 4

# Transformations and interpolations

Different transformations and 2D interpolations are available in OASIS3-MCT to adapt the coupling fields from a source model grid to a target model grid. In the following paragraphs, a description of each transformation, with its corresponding configuration lines that the user has to write in the *namcouple* file, is given. Features that are now deprecated (non functional) compared to prior versions are noted but not described.

### 4.1 Time transformations

- **LOCTRANS:**

LOCTRANS requires one configuring line that gives a time transformation automatically performed below the call to `oasis_put`:

```
LOCTRANS operation
$TRANSFORM
```

where `$TRANSFORM` can be

- **INSTANT:** no time transformation, the instantaneous field is transferred;
- **ACCUMUL:** the arrays `field_array` provided as third argument to the `oasis_put` calls are accumulated, without any specific weighting, and exchanged at the coupling period (`field_array` should therefore not contain the field accumulated between two successive `oasis_put`);
- **AVERAGE:** the arrays `field_array` provided as third argument to the `oasis_put` calls are averaged, without any specific weighting (this means in particular that time steps of varying duration are not supported), and exchanged at the coupling period;
- **T\_MIN** or **T\_MAX:** the minimum or maximum value of the field for each source grid point over the `oasis_put` calls of the previous coupling period is transferred; cannot be used together with the `fracwgt` option (see section 2.2.7) or with higher-order remappings (**BICUBIC**, **CONSERV/SECOND**, see section 4.3);
- **ONCE:** not supported anymore in OASIS3-MCT.

When combined with **BLASOLD** (see section 4.2), **T\_MIN** or **T\_MAX** extracts the minimum or maximum of the raw field array before **BLASOLD** is applied.

When **ACCUMUL** or **AVERAGE** is specified for a field that is sent with dynamic fractions (see the `fracwgt` option in section 2.2.7), the remapping of the field and fractions is performed, the remapped fractions are applied as a normalization term on the target side and then the accumulation or averaging is done on the remapped fields. This may increase the cost compared to **ACCUMUL** or **AVERAGE** without `fracwgt`, where the remapping is done only at coupling time on the accumulated or averaged field.

If the CPP key `LOCTRANS_RESTARTS` is activated when compiling OASIS3-MCT, partial time transformations, if any, will be saved in the coupling restart file at the end of the run for exact restart of the next run. **Note that this is not backward compatible as LOCTRANS restart fields were saved by default in OASIS3-MCT 5.0.** Although we do not recommend it as it may have unexpected side effects, this allows a run to end in the middle of a coupling period. If used, it is important to note the following:

- The initial coupling restart file should not contain any LOCTRANS restart fields.
- For the following runs, it is mandatory to use the coupling restart file produced at the end of the previous run that automatically contains LOCTRANS restart fields coherent with the *namcouple* entries.
- In the restart file, LOCTRANS restart fields are named with the symbolic name of the coupling fields prefixed with `locxxxxxx_` where `xxxxxx` is the 6-digit number of the coupling field ;
- In the restart file, LOCTRANS restart fields for higher-order terms (if any, see section 2.2.7) are named with the symbolic name of the coupling fields prefixed with `av?locxxxxxx_` where `?` is 2, 3, 4, or 5 for the higher-order 2<sup>nd</sup>/3<sup>rd</sup>/4<sup>th</sup>/5<sup>th</sup> coupling field array respectively.
- If the `fracwgt` option is used (see section 2.2.7), LOCTRANS restart fields are also saved for the destination accumulated/averaged normalised scaled field and the fraction field, named with the symbolic name of the coupling fields prefixed respectively with `av1mlocxxxxxx_` and `avfwlocxxxxxx_`.
- Finally, the accumulation or averaging counter is also stored with prefix `alocxxxxxx_` and `locxxxxxx_`.

## 4.2 The pre-processing transformations

Note that previous transformations `REDGLO`, `INVERT`, `MASK`, `EXTRAP`, `CORRECT` are not supported anymore.

- **CHECKIN:**

CHECKIN calculates the global minimum, maximum, mean, and sum of the source field values, taking the mask into consideration, and writes them out. If a grid area or static fraction field is also available, (respectively in the file *areas.nc* or *masks.nc*), then the area and/or fraction weighted mean and sum are also diagnosed and written. All diagnostics are written to the master process OASIS3-MCT debug file (look for `CHECK* diags`).

This operation does not transform the field. CHECKIN operations can slow down the simulation and should not be used in production mode. For backward compatibility, CHECKIN has one generic input line that is no longer used but is still required and can contain anything. See also CHECKOUT.

The generic input line is as follows:

```
CHECKIN operation
 INT = 1
```

- **BLASOLD:**

BLASOLD allows the source field to be scaled and a scalar to be added to the field. The prior ability to perform a linear combination of the current coupling field with other coupling fields on the source side has been deprecated in OASIS3-MCT but is still supported on the target side with BLASNEW, see below. BLASOLD occurs before the interpolation but after the LOCTRANS `T_MIN` or `T_MAX` transformations, if any.

BLASOLD is not allowed with higher-order remappings `CONSERV/SECOND` or `BICUBIC` because it is not uniquely defined and different implementations may lead to different results. Additive or

multiplicative transformations might be better done in the models or with BLASNEW for higher-order remappings.

This transformation requires at least one configuring line with two parameters:

```
BLASOLD operation
 $XMULT $NBFIELDS
```

where \$XMULT is the multiplicative coefficient of the source field. \$NBFIELDS can be 0 or 1 if a scalar does not need or needs to be added. In this last case, an additional input line is required where \$AVALUE is the scalar to be added to the field, which must also be given as a REAL value :

```
 CONSTANT $AVALUE
```

### 4.3 The remapping (or interpolation or regridding)

The 3 options for remapping are MAPPING for using an existing set of weights and addresses, or YAC for using the YAC library to generate remapping weights and addresses, or SCRIPR to use the SCRIP library to generate them. Note that INTERP, MOZAIC, NOINTERP, and FILLING are not supported anymore in OASIS3-MCT.

#### 4.3.1 MAPPING

This keyword is used to specify a MAPPING input file containing the weights and addresses to be read and used for remapping.

MAPPING can be used for higher-order remapping, with up to 5 different sets of weights in the MAPPING file and applied to up to 5 different fields transferred through the oasis\_put arguments, if present (see section 2.2.7).

As for the other transformations, different mappings can be specified for the different coupling fields. Grid data files *grids.nc*, *masks.nc*, and *areas.nc* are not needed for MAPPING.

This transformation requires at least one configuring line with one filename and two optional string values:

```
$MAPNAME $MAPLOC $MAPSTRATEGY
```

- \$MAPNAME is the name of the mapping file to read; this is a NetCDF file consistent with the SCRIPR map file format (see section 5.4) but may have been generated with another tool.
- \$MAPLOC is optional and can be either *src* or *dst*. With *src*, the mapping is done in parallel on the source processors before communication to the destination model processes; this is the default. With *dst*, the mapping is done on the destination processes after the source grid data is sent from the source model. Note that higher-order remappings CONSERV/SECOND or BICUBIC can only be carried out with remapping on the source model processors,
- \$MAPSTRATEGY is optional and can be either *bf*, *sum*, or *opt*. In *bf* mode, the mapping is done using a strategy that produces bit-for-bit identical results regardless of the grid decompositions. With *sum*, the transform is done using the partial sum approach which generally introduces roundoff level changes in the results on different processor counts. Option *opt* allows the coupling layer to choose either approach based on an analysis of which strategy is likely to run faster; usually, partial sums will be used if the source grid has a higher resolution than the target grid as this should reduce the overall communication (in particular for conservative remapping). By default \$MAPSTRATEGY = *bf*.

Note that if SCRIPR or YAC (see below) is used to calculate the remapping file, MAPPING can still be listed in the namcouple, for example to use it with a specific \$MAPLOC or \$MAPSTRATEGY option.

### 4.3.2 YAC

YAC offers the parallel remapping algorithms for 2D interpolation on the sphere from the YAC coupler [Hanke et al 2016] [Hanke and Redler 2019] [Jonville et al 2024]. YAC routines are in `oasis3-mct/lib/yac` and `yaxt` directories.

An interpolation with YAC consists of one or more interpolation methods that form an interpolation stack prescribed in the *namcouple*, as described below. The name of the resulting remapping file, containing remapping weights and addresses, is composed with the character string appearing on the first YAC line in the *namcouple*. When YAC performs a remapping, it first checks if the remapping file exists; if it exists, it reads the weights and addresses from the file; if not, it calculates them and store them in a file with that name. When using an existing remapping file, one has to be sure that it was generated in exactly the same conditions than the ones it is currently used for.

YAC needs the grid data files `grids.nc` containing the longitudes and latitudes of the source and target grid centers and, if the `CONSERV` remapping is prescribed, also including the longitudes and latitudes of the source and target grid corners. If a mask is defined for a grid in the auxiliary file `masks.nc`, it will be taken into account in the interpolation (see section 5.1).

The different types of interpolations available with YAC are described in detail in [Jonville et al 2024]. Examples are also available in `oasis3-mct/pyoasis/examples/11-test-interpolation/common_data/namcouple_yac.bgnc` and `namcouple_yac.bggd`. These are :

- `NNN` : n-nearest-neighbour method ;
- `ZERO` : to set target points to 0.0 ;
- `RBF` : remapping using radial basic functions ;
- `AVG` : average of the non-masked corners of the matching cell ;
- `NCC` : average of the unmasked vertices connected to the nearest cell corner ;
- `HCSBB` : remapping using spherical Bernstein-Bézier polynomials ;
- `CONSERV` : 1<sup>st</sup> or 2<sup>nd</sup> order conservative remapping ;
- `SPMAP` : locally conservative remapping ;
- `CREEP` : creeping extrapolation.

YAC first configuration line in the *namcouple* is:

```
First configuration line for YAC in the namcouple
 $EDGS $EDGT $NINT $FILENAMEAPPEND $NIO
```

where :

- `$EDGS` defines the type of cell borders of the source grid: great circles, `$EDGS = GC`, or longitude-latitude circles, `$EDGS = LL`<sup>1</sup>.
- `$EDGT` : same as `$EDGS` for the target grid.
- `$NINT` defines the number of interpolation methods in the interpolation stack.
- `$FILENAMEAPPEND` is used for building the name of the remapping file, i.e. `rpm_${srcprefix}_to_${dstprefix}_${FILENAMEAPPEND}.nc`, where `$srcprefix` and `$dstprefix` are the prefixes of the source and target grid names, see section 3.3.1.
- `$NIO` is the number of parallel processes used by YAC to read the grid file and write the remapping file to disk. The same processes are used to calculate the intersections of the local source and target grids (while all component model processes are used to calculate the weights *per se*). This `$NIO` number may therefore have a strong impact on the overall performance of the remapping weight calculation. Other environment variables may be set to define these IO processes, see YAC documentation at

[https://dkrz-sw.gitlab-pages.dkrz.de/yac/d6/d7d/io\\_config\\_detail.html](https://dkrz-sw.gitlab-pages.dkrz.de/yac/d6/d7d/io_config_detail.html)

<sup>1</sup>This is especially important for cells from regular grids that are located close to the poles.

The following types of interpolations can then be specified in the *namcouple*, each one having a specific line of parameters described here. Note that for all parameters, the default value, if any, can be activated by setting `DEFAULT` for that parameter.

- **NNN** : n-nearest-neighbour method :

```
Second configuration line for YAC/NNN in the namcouple
NNN $weighting $n $search_distance $scale
```

where

- **NNN** : for N-Nearest-Neighbour
- `$weighting [string]` : how to weight the values of the n nearest neighbours ; can be :
  - \* **AVG** : arithmetic average ;
  - \* **DIST** : inverse distance weighting ;
  - \* **GAUSS** : calculates weights by applying a Gauss function to the distance, with an additional optional scale parameter, see below ;
  - \* **RBF** : Radial Basis Function ; has also an additional optional scale parameter, see below (NNN with RBF weighting is strictly equivalent to RBF method, see below) ;
  - \* **ZERO** : sets the weight to zero with only 1 source point per target point, and therefore sets target point values to 0.0 (strictly equivalent to the ZERO method, see below) : .
- `$n [integer]` : number of source points per target point (mandatory, except for the ZERO weight).
- `$search_distance [real]` or `DEFAULT` : Maximum search distance in degree (optional, default 0.0, valid range [0.0, 180.0]). Note: a value of 0.0 results in an unlimited search distance. Warning: if there are less than the specified number of source points within the maximum search distance, then the respective target point will not be remapped.
- `$scale [real]` or `DEFAULT` : Scaling factor; optional, only for GAUSS and RBF weighting methods; empirical tests have led to default values of 0.1 and 1.487973e+01 respectively.
- **ZERO** : is strictly equivalent to NNN with ZERO weighting. It sets the weight of one source neighbour to zero (the address of the source neighbour is irrelevant) and therefore sets target point values to 0.0 :

```
Second configuration line for YAC/ZERO in the namcouple
ZERO
```

- **RBF**: is strictly equivalent to NNN with RBF weighting. It is based on

Reinheimer, J., 2018: Vector Field Interpolation using Radial Basis Functions, pp 68. :

```
Second configuration line for YAC/RBF in the namcouple
RBF $n $scale
```

where

- **RBF** : for Radial Basis Functions
- `$n [integer]` : number of source points per target point
- `$scale [real]` or `DEFAULT` : Scaling factor; optional ; empirical tests have led to default values of 1.487973e+01
- **AVG** : generates cells based on the centers of the original source grid, then for each target grid point, searches a matching source cell and then applies the selected weighting to the non-masked corners of the matching source cell :

```
Second configuration line for YAC/AVG in the namcouple
AVG $weighting
```

where

- **AVG** : for average of the non-masked corners of the matching cell

- `$weighting [string]` : how to weight the non-masked corners of the matching source cell ; can be :
  - \* `ARITHMETIC` : arithmetic average ;
  - \* `DIST` : inverse distance weighting ;
  - \* `BARY` : triangulates the source cells and then finds the matching source triangle for each target point; computes the weights based on the barycentric coordinates of the target point within this triangle.
- `NCC` : for each target grid point, searches for a matching source cell, as defined above for the `AVG` method; determines the closest vertex in that cell ; selects all unmasked vertices of the source cells connected to this vertex; applies a weighting method to the selected vertices. This method produces results that are very similar to `AVG` method, the major differences occurring close to the mask edges, where the `AVG` method fails. This property makes the `NCC` method a good fallback option, e.g. for `AVG` or `HCSBB`.

```
Second configuration line for YAC/NCC in the namcouple
NCC $weighting
```

where

- `NCC` : for nearest cell corner
- `$weighting [string]` : how to weight the values of the vertices connected to the source nearest cell corner ; can be :
  - \* `AVG` : arithmetic average ;
  - \* `DIST` : inverse distance weighting.
- `HCSBB` : is based on [Alfeld et al 1996]; triangulates the source grid points and estimates the derivatives of the source field across the edges of the triangles; from these derivatives, constructs triangular patches from a blend of spherical Bernstein-Bézier polynomials then used for the interpolation of the target points. With this method, that is computationally expensive and produces a quite large interpolation stencil, the resulting target field always has a continuous first derivative.

```
Second configuration line for YAC/HCSBB in the namcouple
HCSBB
```

where

- `HCSBB` : for Hybrid Cubic Spherical Bernstein-Bézier
- `CONSERV` : implements a  $1^{st}$  order conservative remapping following [Jones 1999] and a  $2^{nd}$  order conservative remapping following [Kritsikis et al 2017] . The source and target cells are defined by the source and target corners (OASIS3-MCT standard). For each target cell, the weight of each source cell is defined by the overlap between the source cell and target cell. When computing the overlaps, YAC differentiates between edges defined on great circles (GC) and longitude-latitude circles (LL), as specified by the user on the first configuration line for YAC (see above). Therefore, in contrast to the `SCRIP` that also provides this interpolation method, YAC does not require coordinate transformation close to the poles.

```
Second configuration line for YAC/CONSERV in the namcouple
CONSERV $order $normalization
```

where

- `CONSERV` : for conservative remapping
- `$order [string]` ; can be :
  - \* `FIRST` : for  $1^{st}$  order conservative remapping.
  - \* `SECOND` : for  $2^{nd}$  order conservative remapping. Compared to the  $1^{st}$  order, this option computes and uses the gradient of the source field and generates a much bigger stencil; it results in a much smoother field, especially when going from a low- to a high-resolution

grid. In contrast to SCRIPR 2<sup>nd</sup> order conservative remapping (see section 4.3.3), the gradient is computed by YAC for each source cell based on its neighbouring cells and does not have to be provided as additional field array arguments in the `oasis_put` call (see section 2.2.7). If, for any reason, the gradient computation fails, a zero-gradient is used for the respective source cell, which is basically a fallback to 1<sup>st</sup> order conservative remapping.

- `$normalization [string]` ; specifies the area normalization method as for SCRIPR, see section 4.3.3; different options produce different results only for target cells that are partially covered by non-masked source cells.
  - \* `FRACAREA` : The sum of the area of the non-masked source cells overlapping with the target cell is used to normalize the respective target field value; this option gives reasonable flux values but may not be locally conservative.
  - \* `DESTAREA` : The target cell area is used to normalize the target field value; this option might generate unreasonable flux values but it ensures local conservation of the source flux.
- `SPMAP` : For each source cell, first searches for the nearest target cell within a maximum search distance (`$search_distance`); the source cell data is then distributed to all non-masked target cells that are within the user-provided maximum spread distance (`$spread`) around the nearest target cell, respecting the grid connectivity around the sea-land mask. If multiple source cells contribute to a target cell, their contributions are summed up. Target cells not associated with any source cell will not get any value. Multiple normalization options (`$scale`) are available based on the cell areas of the associated source/target cells. This method has been implemented in particular to cover the mapping of the hydrological runoff, which is provided on selected cells at the coastline. It is close to the `LOCCUNIF`, `LOCCDIST` and `LOCCGAUS` methods of SCRIPR, see below. In contrast with all other interpolation schemes, which goal is to generate a value for all non-masked target cells, this method aims to be locally mass-conserving and not to lose any flux from the donor (source) cells.

```
Second configuration line for YAC/SPMAP in the namcouple
SPMAP $weighting $spread $search_distance $scale $src_radius $tgt_radius
```

where :

- `SPMAP` : for locally conservative remapping
- `$weighting [string]`, can be :
  - \* `AVG` : for arithmetic average
  - \* `DIST` : for inverse distance weighting
- `$spread [real]` or `DEFAULT` : spread distance (optional, default: 0.0, valid range is [0.0, 90.0]). Note: a value of 0.0 results in the source field being assigned to the single closest target point.
- `$search_distance [real]` or `DEFAULT` : Maximum search distance (optional, default: 0.0, valid range is [0.0, 180.0]). Note: a value of 0.0 results in an unlimited search distance.
- `$scale [string]` or `DEFAULT` : Scaling factor (optional, default: as `NONE`), can be :
  - \* `NONE` : No scaling ; the sum of the weights for each source cell is 1.0 ; to be used in case the source and target field are not provided as a flux (for example, both source and target runoff field in m3/s).
  - \* `SRCAREA`: All weights are scaled by the area of the associated source cell; therefore, the weights for each source cell sum to its area ; to be used in case the source field is provided as a flux while the target field is not (for example, runoff source field units in m/s and runoff target field units in m3/s).

- \* INVTGTAREA : All weights are scaled by the inverse area of the associated target cell ; to be used in case the target field is provided as a flux while the source field is not (for example, runoff source field in m<sup>3</sup>/s and runoff target field in m/s).
- \* FRACAREA : All weights are scaled by the area of the associated source cell and the inverse area of the associated target cell ; to be used in case the source and target fields are provided as a flux (for example, both source and target runoff field in m/s).
- \$src\_radius [real] or [string] : if [real], gives value of the sphere radius for the computation of the source cell areas (optional, default: 1.0); if [string] , can be :
  - \* DEFAULT : default value of 1.0
  - \* EARTH : hard-coded value of 6371229 m.
  - \* FILE or USER : the source cell areas is not computed but read in the auxiliary file areas.nc
- \$tgt\_radius [real] or [string]: same as \$src\_radius but for the target cell areas
- CREEP : is an extrapolation method described in [Kara et al 2007]. For each unmapped target point, the method checks the status of its neighbouring target points. If successfully-interpolated target neighbour points are available, the respective unmapped point value is calculated based on their results. This check is repeated until all target points are remapped or for a specified number of iterations.
 

```
Second configuration line for YAC/CREEP in the namcouple
CREEP $iter
```

 where
  - CREEP : for creeping extrapolation
  - \$iter [integer] or DEFAULT: number of iterations (optional, default: -1.0, valid range is [-1, ∞ [) ; a value of -1 results in the execution of the algorithm until no additional target points can be remapped by this method.

### 4.3.3 SCRIPR

SCRIPR gathers the interpolation techniques offered by Los Alamos National Laboratory SCRIP 1.4 library [Jones 1999]<sup>2</sup>. SCRIPR routines are in `oasis3-mct/lib/scrip`. See the SCRIP 1.4 documentation in `oasis3/doc/SCRIPusers.pdf` for more details on the interpolation algorithms.

Since OASIS3-MCT 4.0 release, a hybrid MPI+OpenMP parallel version of the SCRIP library is available. It relies on the MPI parallel layout of the calling model but only uses one MPI process per node. To benefit from the OpenMP parallelisation of the SCRIP, one has, of course, to compile OASIS3-MCT with options activating OpenMP, see also section 6.1. In The number of OpenMP threads per MPI process used by SCRIP is set by a dedicated environment variable `OASIS_OMP_NUM_THREADS`, which can be different from the default number of threads per MPI process used elsewhere in the application, set by the environment variable `OMP_NUM_THREADS` ; in practice, `OASIS_OMP_NUM_THREADS` has to be smaller or equal to `OMP_NUM_THREADS`. For optimum performance, it is recommended to set `OASIS_OMP_NUM_THREADS` to the number of cores on the node. The `regridenvironment` directory (see section 6.3.3) in `oasis3-mct/examples/regridenvironment` gives a practical example on how to use the SCRIP library (or ESMF or XIOS) in parallel to calculate regridding weight-and-address files, see section 6.3.3. The details of the SCRIP parallelisation can be found in [Piacentini et al 2018] and [Valcke et al 2018].

When the SCRIP library performs a remapping, it first checks if the file containing the corresponding remapping weights and addresses exists; if it exists, it reads them from the file; if not, it calculates them

<sup>2</sup>See also <http://climate.lanl.gov/Software/SCRIP/> and the copyright statement in appendix 1.3.3.

and store them in a file. In the later case, SCRIP needs the grid data files `grids.nc` and `masks.nc` (see section 5.1).

The weight-and-address file is created in the working directory and is by default called `rmp_src_to_tgt_INTTYPE_NORMAOPT.nc`, `rmp_src_to_tgt_INTTYPE_NBR.nc`, or `rmp_src_to_tgt_INTTYPE.nc`, where:

- *src* and *tgt* are the acronyms of the source and the target grids respectively;
- *INTTYPE* is the interpolation type (i.e. `DISTWGT`, `GAUSWGT`, `GAUSWGTF`, `DISTWGTF`, `LOCCUNIF`, `LOCCDIST`, `LOCCGAUS`, `BILINEAR`, - not `BILINEA` as in OASIS3.3, `BILINEARNF`, `BICUBIC`, `BICUBICNF`, or `CONSERV`);
- *NORMAOPT* is the normalization option, for SCRIPR/CONSERV only (i.e. `DESTAREA`, `DESTARTR`, `DESTNNEI`, `DESTNNTR`, `FRACAREA`, `FRACARTR`, `FRACNNEI` or `FRACNNTR`, see below);
- *NBR* is the number of neighbors used for `DISTWGT`, `DISTWGTF`, `GAUSWGT`, `GAUSWGTF`, `LOCCUNIF`, `LOCCDIST` or `LOCCGAUS` only.

One has to take care that the remapping file will have the same name even if other details, like the grid masks or the `$MAPLOC` or `$MAPSTRATEGY` options, are changed. When reusing a remapping file, one has to be sure that it was generated in exactly the same conditions than the ones it is used for.

The following types of interpolations are available:

- `DISTWGT` performs a distance weighted nearest-neighbour interpolation (N neighbours) with a nearest neighbor fill for non-masked target points that do not receive a value. All types of grids are supported.
- `DISTWGTF` performs a distance weighted nearest-neighbour interpolation (N neighbours) without the nearest neighbor fill for non-masked target points. All types of grids are supported.

The configuring line is:

```
SCRIPR (for DISTWGT or DISTWGTF)
 $CMETH $CGRS $CFTYP $REST $NBIN $NV $ASSCMP $PROJCART
```

where:

- `$CMETH` = `DISTWGT` or `DISTWGTF`.
- `$CGRS` is the source grid type (LR, D or U):
  - \* LR: The longitudes and the latitudes of 2D Logically-Rectangular (LR) grid points can be described by two arrays `longitude(i, j)` and `latitude(i, j)`. Stretched or/and rotated grids are LR grids.
  - \* U: Unstructured (U) grids do have any particular structure. The longitudes and the latitudes of 2D Unstructured grid points must be described by two arrays `longitude(nbr_pts, 1)` and `latitude(nbr_pts, 1)`, where `nbr_pts` is the total grid size.
  - \* D: The Gaussian Reduced (D) grid is composed of a certain number of latitude circles, each one being divided into a varying number of longitudinal segments. The grid data (longitudes, latitudes, etc.) must be described by arrays dimensioned `(nbr_pts, 1)`, where `nbr_pts` is the total number of grid points. There are grid points on the Greenwich meridian, but no grid point at the equator nor at the poles, and no overlap of the grid.
- `$CFTYP` is the field type: `SCALAR`. The option `VECTOR`, which in fact leads to a scalar treatment of the field (as in the previous versions), is still accepted. `VECTOR_I` or `VECTOR_J` (i.e. vector fields) are not supported anymore in OASIS3-MCT. See “Support of vector fields with the SCRIPR remappings” below.
- `$REST` is a bin search restriction type: `LATLON` or `LATITUDE`, see SCRIP 1.4 documentation `SCRIPusers.pdf`.
- `$NBIN` the number of restriction bins that must be equal to 1, i.e. no bin restriction, for `DISTWGT`, `DISTWGTF`, `GAUSWGT`, `GAUSWGTF`, `BILINEAR`, `BILINEARNF`, `BICUBIC`

or BICUBICNF; for details, see [Piacentini et al 2018])<sup>3</sup>.

- \$NV is the number of neighbours used.
- \$ASSCMP, \$PROJCART: UNUSED; vector fields are not supported anymore in OASIS3-MCT; see “Support of vector fields with the SCRIPR remappings” below.
- GAUSWGT performs a N nearest-neighbour interpolation weighted by their distance and a gaussian function, with a nearest neighbor fill for non-masked target points that do not receive a value. All grid types are supported.
- GAUSWGTF performs a N nearest-neighbour interpolation weighted by their distance and a gaussian function, without the nearest neighbor fill for non-masked target points. All grid types are supported.

The configuring line is:

```
SCRIPR (for GAUSWGT or GAUSWGTF)
 $CMETH $CGRS $CFTYP $REST $NBIN $NV $VAR
```

where all entries are as for DISTWGT, except that:

- \$CMETH = GAUSWGT or GAUSWGTF
- \$VAR defines the weight given to a neighbour source grid point as proportional to  $\exp(-1/2 \cdot d^2/\sigma^2)$  where  $d$  is the distance between the source and target grid points, and  $\sigma^2 = $VAR \cdot \overline{D}^2$  where  $\overline{D}^2$  is, for each target grid point, the square of average distance between its source grid points (calculated automatically by OASIS3-MCT).
- LOCCUNIF, LOCCDIST and LOCCGAUS perform a locally conservative interpolation by associating N target nearest neighbours to every SOURCE grid point, and applying a weight normalization taking into account the source/target mesh area ratio. Interpolation weights can additionally be modulated by the source/target distances (LOCCDIST) or by the source/target distances and a gaussian function (LOCCGAUS). All types of grids are supported. These interpolations are convenient to transfer coupling fields from disjoints areas, such as the river runoff flux, as studied in [Voltaire 2020]. For details, see [Maisonnavé 2020].

The configuring line is:

```
SCRIPR (for LOCCUNIF or LOCCDIST)
$CMETH $CGRS $CFTYP $REST $NBIN $NV
or
SCRIPR (for LOCCGAUS)
$CMETH $CGRS $CFTYP $REST $NBIN $NV $VAR
```

where:

- \$CMETH \$CGRS \$CFTYP \$REST \$NBIN \$NV entries are as for DISTWGT and \$VAR is as for GAUSWGT. Note that for these interpolations, \$NV represents the number of *target* (and not source) neighbours.
- BILINEAR performs an interpolation based on a local bilinear approximation (see details in chapter 4 of SCRIP 1.4 documentation SCRIPUsers.pdf). Logically-Rectangular (LR) and Reduced (D) source grid types are supported. It also generates regridded values with a nearest neighbor fill for non-masked target points that do not receive any value with the original algorithm.
- BILINEARNF is identical to BILINEAR but without the nearest neighbor fill.
- BICUBIC performs an interpolation based on a local bicubic approximation for Logically-Rectangular (LR) grids (see details in chapter 5 of SCRIP 1.4 documentation SCRIPUsers.pdf), and on a 16-point

<sup>3</sup>The only exceptions are for Gaussian Reduced (D) grids for (BILINEAR, BILINEARNF, BICUBIC and BICUBICNF; in that case, if the Gaussian-reduced grid is stored from North to South the number of bins is the number of latitude circles of the grid (minus one, to be precise), independently of \$NBIN; for Gaussian-reduced grid stored from South to North to South, the bin definition will not work and the interpolation will become a 4 distance-weighted nearest-neighbour for all target points.

stencil for Gaussian Reduced (D) grids. Note that for Logically-Rectangular grids, 4 weights for each of the 4 enclosing source neighbours are required corresponding to the field value at the point, the gradients of the field with respect to  $i$  and  $j$ , and the cross gradient with respect to  $i$  and  $j$  in that order. OASIS3-MCT will calculate the remapping weights and addresses (if they are not already provided) but will not, at run time, calculate the two gradients and the cross-gradient of the source field (as was the case with OASIS3.3). These 3 extra fields need to be calculated by the source code and transferred as extra arguments of the `oasis_put` (see `fld2`, `fld3`, `fld4` in section 2.2.7). Finally, this interpolation will fill non-masked target points that do not receive a value with a nearest neighbor fill.

- BICUBICNF is identical to BICUBIC but without the nearest neighbor fill.

For BILINEAR, BILINEARNF, BICUBIC, and BICUBICNF, the configuring line is:

```
SCRIPR (for BILINEAR, BILINEARNF, BICUBIC, or BICUBICNF)
$CMETH $CGRS $CFTYP $REST $NBIN
```

where:

- \$CMETH = BILINEAR, BILINEARNF, BICUBIC, or BICUBICNF
- \$CGRS is the source grid type: LR or D.
- \$CFTYP, \$REST, \$NBIN are as for DISTWGT.

Note that for DISTWGT, GAUSWGT, BILINEAR and BICUBIC:

- Masked target grid points: the zero value is associated to masked target grid points.
- Non-masked target grid points having some of the source points normally used in the interpolation masked: a N nearest neighbour algorithm using the remaining non masked source points is applied.
- Non-masked target grid points having all source points normally used in the interpolation masked: by default, the nearest non-masked source neighbour is used (`ll_nnei` is set to `.true.` in the SCRIP run).

Note that for DISTWGTNF, GAUSWGTNF, BILINEARNF and BICUBICNF:

- Identical behavior to non-NF options except that no nearest non-masked source neighbour fill is done (`ll_nnei` is set to `.false` in the SCRIP run). Non-masked target grid points that do not receive any interpolated value are set to 0.
- CONSERV performs 1st or 2nd order conservative remapping, which means that the weight of a source cell is proportional to area intersected by the target cell (plus, for the second order, some other terms proportional to the gradient of the field in the longitudinal and latitudinal directions).

The configuring line is:

```
SCRIPR (for CONSERV)
$CMETH $CGRS $CFTYP $REST $NBIN $NORM $ORDER $NTHRESH $STHRESH
```

where:

- \$CMETH = CONSERV ;
- \$CGRS is the source grid type: LR, D and U. Note that the grid corners have to given by the user in the grid data file `grids.nc` or by the code itself in the initialisation phase by calling routine `oasis_write_corner` (see section 2.2.4) ;
- \$CFTYP, \$REST are as for DISTWGT;
- \$NBIN is the number of restriction bins (bin restriction is effectively allowed for CONSERV; for details, see [Piacentini et al 2018]) ;
- \$NORM is the normalization option:
  - \* FRACAREA or FRACARTR: The sum of the non-masked source cell intersected areas is used to normalise each target cell field value: the flux is not locally conserved, but the flux value remains in a physical range. With FRACARTR, an additional correction involving

the “true” area of the grid cells provided in the file `areas.nc` is also applied, see details in the paragraph on the “True Area” (TR) correction below.

- \* `FRACNNEI` or `FRACNNTR`: as `FRACAREA` or `FRACARTR`, except that an additional unmasked source nearest neighbour is used for unmasked target cells that intersect only masked source cells.
- \* `DESTAREA` or `DESTARTR`: The total target cell area is used to normalise each target cell field value even if it only partly intersects non-masked source grid cells: local flux conservation is ensured, but the flux can reach not physical values. With `DESTARTR`, an additional correction involving the “true” area of the grid cells provided in the file `areas.nc` is also applied, see details in the paragraph on the “True Area” (TR) correction below.
- \* `DESTNNEI` or `DESTNNTR`: as `DESTAREA` or `DESTARTR`, except that an additional unmasked source nearest neighbour is used for unmasked target cells that intersect only masked source cells.
- `$ORDER`: `FIRST` or `SECOND` for first or second order conservative remapping respectively (see `SCRIP 1.4` documentation).

For `CONSERV/SECOND`, 3 weights are needed; `OASIS3-MCT` will calculate these weights and corresponding addresses (if they are not already provided) but will not, at run time, calculate the two extra terms to which the second and third weights should be applied. These terms, respectively the gradient of the field with respect to the latitude ( $\theta$ )  $\frac{\delta f}{\delta \theta}$  and the gradient of the field with respect to the longitude ( $\phi$ )  $\frac{1}{\cos \theta} \frac{\delta f}{\delta \phi}$  need to be calculated by the source code and transferred as extra arguments of the `oasis_put` as `f1d2` and `f1d3` respectively (see section 2.2.7). Note that `CONSERV/SECOND` is not positive definite.

- `$NTHRESH` is the value of the northern latitude threshold in radians where conservative area computation switches from linear boundaries in longitude and latitude at the equator to a Lambert equivalent azimuthal projection toward the north pole. This value is optional and the default is 2.0 radians (greater than 90 degrees), which means the Lambert projection is not activated. (Note that `$NTHRESH` was set to 1.45 in previous versions prior to `OASIS3-MCT.4.0.`) If this value is set, `$STHRESH` must be set as well.
- `$STHRESH` is the value of the southern latitude threshold in radians where conservative area computation switches from linear boundaries in longitude and latitude at the equator to a Lambert equivalent azimuthal projection toward the south pole. This value is optional and the default is -2.0 radians (less than -90 degrees) which means the Lambert projection is not activated. If this value is set, `$NTHRESH` must be set as well.

### Precautions related to the use of the `SCRIPR/CONSERV` remapping (consider using `YAC` remapping instead)

1. Lambert projection above `$NTHRESH` or below `$STHRESH`

For the 1st order conservative remapping, the weight of a source cell is proportional to area of the source cell intersected by target cell. Using the divergence theorem, the `SCRIP` library evaluates this area with the line integral along the cell borders enclosing the area. As the real shape of the borders is not known (only the location of the corners of each cell is known), the library assumes that the borders are linear in latitude and longitude between two corners. This assumption becomes less and less valid closer to the pole.

To correct this inaccuracy, for latitudes above `$STHRESH` or below `$STHRESH`, the library evaluates the intersection between two cell borders using a Lambert equivalent azimuthal projection. [Valcke and Piacentini 2019] analyses the impact of the Lambert projection for specific grids.

2. Another limitation of the `SCRIP` 1st order conservative remapping algorithm is the assumption, for line integral calculation, that  $\sin(\text{latitude})$  is linear in longitude on the cell borders which again is in general not valid close to the pole.

3. For a proper conservative remapping, the corners of a cell have to coincide with the corners of its neighbour cell, with no “holes” between the cells.
4. Overlapping cells : If two cells of one same grid overlap, the one with the greater numerical index must be masked in *masks.nc* for a proper conservative remapping. For example, if the grid cells with  $i=I$  overlaps the grid cells with  $i=imax$ , the latter must be masked. If none of the overlapping cells is masked (given the original mask defined in *masks.nc*), OASIS3-MCT must be compiled with the CPP key `TREAT_OVERLAY` which will ensure that these rules are respected.
5. Remapped field values for masked (non-active) target grid cells are set to 0.
6. If a target grid cell intersects only masked source cells, or falls outside the source grid domain, it will get a zero value unless the `DESTNNEI`, `DESTNNTR`, `FRACNNEI`, or `FRACNNTR` normalisation options are used, in which case it will get the source nearest non-masked neighbour value.

#### The “True Area” (TR) correction: `DESTARTR`, `DESTNNTR`, `FRACARTR`, or `FRACNNTR`

The approximations and hypotheses adopted by SCRIP impact its estimation of the grid cell areas. Therefore, to have an exact conservation of the field surface-integrated values, a correction based on the “True” (TR) area of the cells can be applied by choosing `DESTARTR`, `DESTNNTR`, `FRACARTR` or `FRACNNTR` options. These are based respectively on `DESTAREA`, `DESTNNEI`, `FRACAREA` and `FRACNNEI` normalisations adding the so-called “TR correction”. The true area of the cell, i.e. the one considered by the component model itself, must be provided in the auxiliary file *areas.nc* in **square radians**. Equations from [Chavas et al 2013] (eqn. (27) in particular) are implemented.

Special care is taken in the implementation for “polar” cells in the SCRIP sense. The SCRIP detects cells encompassing a pole or cells with a side going through a pole as “polar” cells and a specific treatment is applied for those cells modifying its area. The resulting estimated area serves as a normalisation factor but its value is not representative of the surface of the cell anymore. For this reason, the TR correction is not activated for those “polar” cells.

More details can be found in section 6 of [Valcke and Piacentini 2019]. A full validation of the TR correction can be found in that report. It is noted there that the TR correction always improves the results even if, in the cases tested, the misfit between the true area and the area evaluated by the SCRIP is always small (except for “polar” cells of course).

#### Support of vector fields with the SCRIP remappings

Vector mapping is not supported and will not be supported by OASIS3-MCT. For proper treatment of vector fields, the source code has to send the 3 components of the vector projected in a Cartesian coordinate system as separate fields. The target code has to receive the 3 interpolated Cartesian components and recombine them to get the proper vector field.

## 4.4 The post-processing stage

Note that `SUBGRID`, `MASKP`, `REVERSE`, `GLORED` are not supported anymore in OASIS3-MCT.

- **CONSERV:**

`CONSERV` performs a global modification of the coupling field such that the area integrated fields are conserved.

This analysis requires that the source and target grid mesh areas be defined in file *areas.nc* file. **The areas must be expressed in matching units on the source and destination sides. Furthermore, these must be square radians if the TR correction is activated** (see paragraph The “True Area” (TR) correction in section 4.3).

This analysis also requires that the source and target grid mask be defined in file *masks.nc*. If grid cell static fractions are also defined in that file, the mask and static fractions are considered and

must be consistent; OASIS3-MCT will abort if they are not or if both are missing<sup>4</sup>. Note that by OASIS3-MCT conventions for the mask, a grid cell with mask=0 (active) should have fractions greater than 0 and a grid cell with mask=1 (inactive) should have fractions equal to 0.

In the *namcouple*, CONSERV requires one input line with one argument and one optional argument:

```
CONSERV operation
 $CMETH $CONSOPT
```

where:

- \$CMETH is the method desired with the following choices; a detailed analysis of these choices can be found in [Craig 2019] :
  - \* with GLOBAL, the field is integrated on both source and target grids, without considering values of masked points, and the residual (target - source) is uniformly distributed on the target grid as an additive term; this option ensures global conservation of the field;
  - \* with GLBPOS, the same operation as GLOBAL is performed except that the residual is distributed proportionally to the value of the original field as a multiplicative term; this option ensures the global conservation of the field ; if the field is well behaved, multiplication factors close to 1 are expected and the sign of the field is not changed.
  - \* with GSSPOS, the same operation as GLBPOS is performed except that the multiplicative term is computed separately for positive and negative values of the field; this option ensures the global conservation of the field and, as \$GLBPOS, does not change the sign of the field, but it is more expensive because many extra global sums are required; this should be used in cases where the area averaged field value tends to 0 as these cases can generate poor corrections, even leading to changes of sign, when carried out with the GLBPOS option.
  - \* with BASBAL, the operation is analogous to GLOBAL except that the non-masked active surface of the source and the target grids are taken into account in the calculation of the residual; this option does not ensure global conservation of the field but ensures that the energy received is proportional to the non-masked active surface of the target grid;
  - \* with BASPOS, the operation is analogous to GLBPOS except that the non-masked surface of the source and the target grids are taken into account and the residual is distributed proportionally to the value of the original field; this option does not ensure global conservation of the field but ensures that the energy received is proportional to the non-masked surface of the target grid and it does not change the sign of the field if the field is well behaved.
  - \* with BSSPOS, the same operation as BASPOS is performed except that the multiplicative term is computed separately for positive and negative values of the field; this option has the same characteristics as BASPOS except that it does not change the sign of the field, but is more expensive because many extra global sums are required; this should be used

---

<sup>4</sup>In principle, the fractions can be defined for both the source and target grids. But for ocean-atmosphere coupling, we strongly encourage the following best practice for a consistent ocean-atmosphere coupled system. Indeed, to have a well-posed coupled problem, the ocean and the atmospheric total surfaces must be the same allowing global conservation of integrated quantities. To do so, the original ocean mask should be taken as it is from the ocean model. For the atmosphere, cell fractions should be defined by the conservative remapping of [1 - ocean mask] on the atmospheric grid, retaining fractions above a certain threshold, to be fixed by the user. These atmospheric cell fractions should be used in the atmospheric model to define the % of ocean (water) subsurface to be considered. Then the atmospheric coupling mask should be adapted associating a non-masked index (i.e. 0) to all cells with a water fraction above the chosen threshold. The global water surface as seen by the atmosphere model is then the sum of its cell areas multiplied by its respective cell fractions. Note that invalid masked atmospheric cells should have null ocean fractions. If we follow this best practice, the atmospheric cell fractions and mask will be specific to the coupling with each particular ocean grid. A specific attribute named `coherent_with_grid` indicating the grid prefix of the companion grid may be defined for mask and fractions. If the OASIS API is used to define the mask and fractions, this can be done via the optional argument `companion` indicating the grid prefix of the companion grid (see section 2.2.4)

in cases where the area averaged field value tends to 0 as these cases can generate poor corrections, even leading to changes of sign, when carried out with the `BASPOS` option.

- `$CONSOPT` is an optional argument specifying the conservative algorithm. Details on the performance of the different options can also be found in [Craig et al 2017] and references there in.
  - \* `bfb`, or `reprosum`, is the default for `CONSOPT` ; it uses a fixed point method based on ordered double integer sums that requires two scalar reductions per global sum. The cost of `reprosum` will be higher than some of the other methods but it will be bit-for-bit for different processor counts or different decompositions except in extremely rare cases and the cost is significantly less than the `gather` option. See [Mirin and Worley 2012].
  - \* `gather` computes global sums by gathering a decomposed array onto the root process before doing an index ordered sum. This is guaranteed to produce identical results for different numbers of processors and decompositions but is expensive both with respect to performance and memory use. This is equivalent to the `bfb` option in previous OASIS3-MCT versions.
  - \* `lsum16` computes a local sum at quadruple precision before doing an MPI reduction on the local sums at quadruple precision. This is likely to be bit-for-bit for different numbers of processors and decompositions but that's not guaranteed. This is just like `lsum8` but at quadruple precision and a little slower.
  - \* `lsum8` computes a local sum at double precision before doing an MPI reduction on the local sums at double precision. This is NOT likely to be bit-for-bit for different numbers of processors and decompositions. This is just like `lsum16` but at double precision and faster.
  - \* `ddpdd` is a parallel double-double algorithm using a single scalar reduction. It should behave between `lsum8` and `lsum16` with respect to performance and reproducibility. See [He and Ding 2001].
  - \* `opt` carries out the global sum using the fastest algorithm generally available. Currently, this is set to `lsum8`.

- **BLASNEW:**

`BLASNEW` performs a scalar multiply or scalar add to any destination field. This is similar to `BLASOLD` on the destination side, but in addition other coupling fields on the destination side can be added with a multiplier and addition weight.

This transformation requires at least one configuring line with two parameters:

```
BLASNEW operation
 $XMULT $NBFIELDS
```

where `$XMULT` is the multiplicative coefficient of the destination field. `$NBFIELDS` will be 0 if no additional fields or scalars are needed, 1 if a single scalar needs to be added, and greater than 1 if additional fields are to be added. The number of `$NBFIELDS` indicates the number of additional lines. If `$NBFIELDS` is greater than 0, the first additional input line must be the string `CONSTANT` and then a real value, `$AVALUE`, which will be added to the field. Even if `$AVALUE` is zero, this line must still be included if `$NBFIELDS` is greater than 0. If `$NBFIELDS` is greater than 1, then additional input lines have the format `$FNAME $XMULT $AVALUE` where `$FNAME` is the name of a field received in OASIS3-MCT in the same component model and `$XMULT` and `$AVALUE` are the multipliers and additive terms to be applied to `$FNAME` :

```
CONSTANT $AVALUE
$FNAME1 $XMULT1 $AVALUE1
$FNAME2 $XMULT2 $AVALUE2
```

For example :

```

2.0 3
CONSTANT 0.0
FLD001 1.0 0.0
FLD002 5.0 -100.

```

will multiply the destination field by 2.0 and then add (FLD001 + FLD002\*5.0 - 100) to that destination field.

All combined fields must be received by the same component in OASIS3-MCT (either via coupling or input), and the field size and decomposition must be consistent across all fields being combined. The value of the field being combined is associated with the last valid coupled value. This allows fields to be combined that are not coupled at the same frequency by using valid lagged values. The order of the receive calls is also important. If a field to be combined is received after the current destination field, then the values from an earlier timestep are used. Note that a more transparent implementation might be to combine fields in the component model (not in OASIS3-MCT) after they are received independently.

- **CHECKOUT:**

CHECKOUT calculates the global minimum, maximum, mean, and sum of the destination field values taking the mask into consideration. If a grid area or fraction field is also available, (respectively in files *areas.nc* or *masks.nc*), the area and/or fraction weighted mean and sum are also diagnosed and written. All diagnostics are written to the master process OASIS3-MCT debug file (under CHECK\* diags).

Note that CHECKOUT does not provide exact weighted mean and sum when the `fracwgt` option involving dynamic fractions is activated, see section 2.2.7; indeed in that case, CHECKOUT will not use the equivalent active destination fraction and a warning will be printed in the debug files.

This operation does not transform the field. CHECKOUT operations can slow down the simulation and should not be used in production mode. For backward compatibility, CHECKOUT has one generic input line that is no longer used but is still required and can contain anything. See also CHECKIN.

The generic input line is as follows:

```

CHECKOUT operation
 INT = 1

```

## Chapter 5

# OASIS3-MCT auxiliary data files

OASIS3-MCT uses auxiliary data files, defining the grids of the models being coupled, or containing coupling field restart values or input data values or remapping weights and addresses. These files can be created by the user before the run or by OASIS3-MCT itself in the initialisation phase.

### 5.1 Grid data files

OASIS3-MCT requires grid data files *grids.nc*, *masks.nc* and *areas.nc* for certain operations:

- *grids.nc* and *masks.nc* for all SCRIPR and YAC regriddings (see section 4.3) ;
- *areas.nc* for SCRIPR conservative remapping when the normalisation by the true area of the cells is activated (i.e. `DESTARTR`, `DESTNNTR`, `FRACARTR`, or `FRACNNTR`) ; for YAC source to target mapping method SPMAP when the option to use areas defined in *areas.nc* is activated (`src_radius` or `tgt_radius = FILE`), see section 4.3;
- *masks.nc* and *areas.nc* for global CONSERV operation, see section 4.4.

These NetCDF files can be created by the user before the run or can be written directly at run time by the component model processes in the initialisation phase using the grid data definition routines (see section 2.2.4).

The arrays containing the grid information are dimensioned  $(nx, ny)$ , where  $nx$  and  $ny$  are the grid first and second dimension. Unstructured grids or other grids expressed with 1D vectors are supported by setting  $nx$  to the total number of grid points and  $ny$  to 1.

1. *grids.nc*: contains the model grid longitudes and latitudes in double precision REAL arrays. The array names must be composed of a prefix (4 characters), given by the user in the *namcouple* on the second line of each field (see section 3.3), and of a suffix, `.lon` or `.lat`, for respectively the grid point longitudes or latitudes.

If the YAC/CONSERV or SCRIPR/CONSERV remapping is specified, longitudes and latitudes for the source and target grid **corners** must also be available in the *grids.nc* file as double precision REAL arrays dimensioned  $(nx, ny, nc)$  where  $nc$  is the maximum number of corners (in the counterclockwise sense, starting by any corner) over all cells;  $nc$  can be any number. For cells that do not have the maximum number of distinct corners, we recommend to repeat the last corner as many times as needed to describe  $nc$  corners. The names of the arrays must be composed of the grid prefix and the suffix `.clo` or `.cla` for respectively the grid corner longitudes or latitudes.

Longitudes must be given in degrees East in the interval -360.0 to 720.0. Latitudes must be given in degrees North in the interval -90.0 to 90.0. Note that if some grid points overlap, it is recommended to define those points with the same number (e.g. 360.0 for both, not 450.0 for one and 90.0 for the other) to ensure automatic detection of overlap by OASIS3-MCT.

The corners of a cell cannot be defined modulo 360 degrees. For example, a cell located over Greenwich will have to be defined with corners at -1.0 deg and 1.0 deg but not with corners at 359.0 deg and 1.0 deg.

Cells larger than 180.0 degrees in longitude are not supported.

Longitudes and latitudes of grid points or corners can be defined through the `oasis_write_grid` or `oasis_write_corner` API routines respectively, see section 2.2.4.

2. *masks.nc*: contains the masks for all component model grids in INTEGER arrays. **Be careful to follow the historical OASIS3-MCT convention: 0 = not masked (i.e. active), 1 = masked (i.e. not active) for each grid point.** The array names must be composed of the grid prefix and the suffix “.msk”. Mask can be defined through the `oasis_write_mask` API routine, see section 2.2.4.

File *masks.nc* may also contain the grid cell static fractions that defines the active (i.e. not masked) part of the cells. Note that those static fractions must not be confused with *dynamic* fractions supported by the `fracwgt` option, see section 2.2.7. The static fraction fields, if present, are only used in the global CONSERV, CHECKIN and CHECKOUT operations. If both a mask and fractions are defined for a grid, they must be consistent; OASIS3-MCT will abort if they are not coherent or if both are missing. Note that given OASIS3-MCT convention for the mask, a gridcell with mask=0 (active) should have a fractions greater than 0 and a gridcell with mask=1 (inactive) should have a fractions equal to 0. The fraction array name must be composed of the grid prefix and the suffix `.frc`. Fractions can be defined through the `oasis_write_frac` API routine, see section 2.2.4.

3. *areas.nc*: this file contains mesh surfaces for the component model grids in double precision REAL arrays. The array names must be composed of the grid prefix and the suffix `.srf`. The surfaces may be given in any units but they must be the same on the source and target sides; furthermore they must be in square radians if the True Area (TR) correction is activated, see section 4.3. Surfaces can be defined through the `oasis_write_area` API routine, see section 2.2.4.

## 5.2 Coupling restart files

At the beginning of a coupled run, coupling fields having a positive LAG defined in the *namcouple* are initially read from their coupling restart file on their source grid (see the LAG concept in section 2.5.2). At the end of the run, the coupling restart files are automatically overwritten under the `oasis_put` when the `date+LAG=runtime`. **It is the user’s responsibility to ensure that the overwritten restart file is copied in the working directory of the next run.** The coupling restart files must follow the NetCDF format.

The name of the coupling restart file is given by the 6th character string on the first configuring line for each field in the *namcouple* and the restart field in that file must have the symbolic name of the coupling field in the *namcouple*, (see section 3.3). If the coupling exchange involves higher-order fields (see section 2.2.7), the higher-order restart fields in the coupling restart file must have the same name than the coupling field itself prefixed with either `av2_`, `av3_`, `av4_`, or `av5_`. For example, higher-order restart fields for the `SOSSTSST` field in section 3.3.1, if needed, would be named `av2_SOSSTSST`, `av3_SOSSTSST`, `av4_SOSSTSST`, and `av5_SOSSTSST`. These fields have to be provided on the source grid in single or double precision REAL arrays and, as the grid data files, must be dimensioned `(nx, ny)`, where `nx` and `ny` are the grid first and second dimension (see section 5.1 above). The shape and indexing of each restart field (and of the corresponding coupling fields exchanged during the simulation) must be coherent with the shape of its grid data arrays.

If the `fracwgt` option is activated for a field with a LAG (see section 3.3), special care have to be taken on the content of the coupling restart files. If no `LOCTRANS ACCUMUL` or `AVERAGE` time transformation is specified for the field, the coupling restart file may contain only the restart field, with its symbolic name, and a dynamic fraction field (see the `fracwgt` argument of the `oasis_put` call in section 2.2.7) named `av1mxxxxxx_YYYYYYYY`, where `xxxxxx` is the 6-digit number of the coupling field and `YYYYYYYY`

is the field symbolic name. But if a LOCTRANS ACCUMUL or AVERAGE time transformation is specified for the field, then the coupling restart files must contain in addition the restart coupling field remapped on the destination grid taking into account the dynamic fraction, named `avfwxxxxxx_YYYYYYYY`, where `xxxxxx` is the 6-digit number of the coupling field and `YYYYYYYY` is the field symbolic name.

Coupling fields coming from different models cannot be in the same coupling restart files, but for each model, there can be an arbitrary number of fields written in one coupling restart file. One exception is when a coupling field sent by a source component model is associated with more than one target field and model; in that case, if coupling restart files are required, it is mandatory to specify different files for the different fields.

If the CPP key `LOCTRANS_RESTARTS` is activated when compiling OASIS3-MCT, the coupling restart files are also used automatically by OASIS3-MCT to allow partial LOCTRANS time transformation to be saved at the end of a run for exact restart at the start of the next run. Although we do not recommend it as it may have unexpected side effects, this allows a run to end in the middle of a coupling period. In that case, the initial coupling restart file should not contain any LOCTRANS restart fields. For the following runs, it is mandatory that the coupling restart file contains LOCTRANS restart fields coherent with the current *namcouple* entries. For example, it will not be possible to restart a run with a multiple field entry in the *namcouple* with a coupling restart file created by a run not activating this multiple file option. See section 4.1 for more detail on the LOCTRANS restart fields in the coupling restart files.

### 5.3 Input data files

Fields with status `INPUT` in the *namcouple* are, at runtime, simply be read in from a NetCDF input file by the target model below the `oasis_get` call, at appropriate times corresponding to the input period indicated by the user in the *namcouple*.

The name of the file must be the one given on the first configuring line for that field in the *namcouple* (see section 3.3.3). There must be one input file per `INPUT` field, containing a time sequence of the field in a single or double precision REAL array named with the same symbolic name as in the *namcouple* and dimensioned `(nx, ny, time)`. The time variable has to be an array `time(time)` expressed in “seconds since beginning of run” (or any other time units as long as the same are used in all components and in the *namcouple*). The “time” dimension has to be the unlimited dimension.

### 5.4 Transformation auxiliary data files

The mapping files to be used for the MAPPING option must be consistent with the files generated by the SCRIP or YAC library to be used for the SCRIPR or YAC transformations (see also section 4.3). The NetCDF files contain the following fields:

dimensions:

```
src_grid_size = xxx ;
dst_grid_size = xxx ;
num_links = xxx ;
num_wgts = xxx ;
```

variables:

```
int src_address(num_links)
int dst_address(num_links)
double remap_matrix(num_links, num_wgts)
```

where

- `src_grid_size` is a NetCDF dimension, here a scalar integer indicating the total number of points in the source grid. This field is .

- `dst_grid_size` is a NetCDF dimension, here a scalar integer indicating the total number of points in the target grid.
- `num_links` is a NetCDF dimension, here a scalar integer indicating the total number of associated source and target grid point pairs in the file. This field is a netCDF dimension.
- `num_wgts` is a NetCDF dimension, here a scalar integer indicating the number of weights per associated grid point pair (up to 5 are supported, see sections 2.2.7 and 4.3 for BICUBIC and CONSERV/SECOND).
- `src_address` is a one dimensional array of size `num_links`. It is a NetCDF variable, containing the integer source address associated with each weight.
- `dst_address` is a one dimensional array of size `num_links`. It is a NetCDF variable containing the integer destination address associated with each weight.
- `remap_matrix` is a two dimensional array of size `(num_links, num_wgts)`. It is a NetCDF variable containing the real weight value(s) associated with the source and destination address. For each link, up to 5 weights are supported, see sections 2.2.7 and 4.3 especially for BICUBIC and CONSERV/SECOND.

## Chapter 6

# Compiling, running, debugging, load balancing

### 6.1 Compiling OASIS3-MCT

OASIS3-MCT is a mixed MPI-OpenMP parallel code. Compiling OASIS3-MCT libraries can be done in the `oasis3-MCT/util/make_dir` directory with the makefile `TopMakefileOasis3`. In the new OASIS3-MCT v6 compiling environment, two environment variables `$(OASIS_COUPLE)` and `$(OASIS_ENV)` have to be defined:

- `$(OASIS_COUPLE)` gives **the location of the coupler sources** (`/lib`, `/util`, `/examples`, etc. directories);
- `$(OASIS_ENV)` gives **the extension of the header Makefile to be used for compilation**.

`TopMakefileOasis3` includes `make.inc` which includes the header Makefile `make. $(OASIS_ENV)` specific to the hardware and compiling environment used. We recommend to compose `$(OASIS_ENV)` as `(computer_name)_(compiler_version)_(MPI_library_version)`. A file named `comp_env.$(OASIS_ENV).sh` should also exist in the compiling directory. It defines the computer environment for running coupled model, in particular the test examples from `oasis3-mct/examples`, and should to be activated in the user environment.

Several header files `make.$(OASIS_ENV)` and `comp_env.$(OASIS_ENV).sh` files are distributed with the release and can be used as a template to create a custom file for your machine. In those header files, appropriate options must be activated for compiling OASIS3-MCT in debug mode or with OpenMP, when applicable (see respectively the environment variables `$(OASIS_DEBUG)` and `$(OASIS_OPENMP)` in those files). Note that it is mandatory that the coupled model components be compiled with the same compilers and system software than the OASIS3-MCT library itself.

After successful compilation, resulting libraries are found in the directory in `$(ARCHDIR)/lib1` while object and module files are found in `$(ARCHDIR)/build-static` and `$(ARCHDIR)/build-shared`.

OASIS3-MCT has historically created static libraries for use in Fortran source codes. However, C language bindings are now available and python codes are now fully supported. Therefore, the Makefile `TopMakefileOasis3` supports compilation of both static and shared libraries.

`TopMakefileOasis3` has several targets including:

- `oasis3-psmile` = static-libs-fortran (for backwards compatibility)
- `static-libs-fortran` = static OASIS3-MCT libraries for Fortran only (default)
- `shared-libs-fortran` = shared (dynamic) OASIS3-MCT libraries for Fortran only
- `static-libs` = static OASIS3-MCT libraries including Fortran and c-bindings

---

<sup>1</sup>where `$(ARCHDIR) = $(OASIS_COUPLE)/INSTALL_OASIS.$(OASIS_ENV)`, see the header Makefiles

- `shared-libs` = shared (dynamic) OASIS3-MCT libraries including Fortran and c-bindings
- `pyoasis` = builds and installs `shared-libs` plus higher and intermediate python classes
- `realclean` = cleans and resets the build
- `help` = provides information about the different targets

Log and error messages from compilation are normally saved in the directory `/util/make_dir` in the files `COMP.log` and `COMP.err` or similar. The `TopMakefileOasis3` output will direct users to the compile output files.

To interface a component code with OASIS3-MCT and use the module `mod_oasis` (see section 2.2.1), it is required to include OASIS3-MCT modules from `$(ARCHDIR)/include` and link with appropriate libraries in `$(ARCHDIR)/lib` during the compilation and linking.

## 6.2 CPP keys

The following CPP keys can be used when compiling OASIS3-MCT, in general to be specified in the `CPPDEF` variable in the header Makefile :

- `YAC_REMAP`: to compile the YAC interpolation library, see section 4.3. Note that this CPP key can be activated by specifying `ENABLE_YAC = YES` in the header Makefile.
- `LOCTRANS_RESTARTS`: to allow partial time transformations, if any, to be saved at the end of the run in the coupling restart files for exact restart of the next run. **Note that this is not backward compatible as `LOCTRANS` restart fields were saved by default in OASIS3-MCT 5.0.** Although we do not recommend it as it may have unexpected side effects, this allows a run to end in the middle of a coupling period. Note that this CPP key can be activated by specifying `OASIS_LOCTRANS_RESTARTS = YES` in the header Makefile. See section 4.1 for important remarks regarding the use of this CPP key.
- `TREAT_OVERLAY`: to ensure, in `SCRIPR/CONSERV` remapping (see section 4.3), that if two cells of the source grid overlay and none is masked a priori, the one with the greater numerical index will not be considered (they also can be both masked); mandatory for the remapping. For example, if the grid line with  $i=1$  overlaps the grid line with  $i=imax$ , it is the latter that must be masked; when this is not the case with the mask defined in `masks.nc`, this CPP key forces this rules to be respected.
- `__NO_16BYTE_REALS`: must be specified if you compile with PGF90.

## 6.3 Examples on how to run OASIS3-MCT

The following examples of running environments are provided in the directory `oasis3-mct/examples`.

### 6.3.1 tutorial.communication

The directory `oasis3-mct/examples/tutorial.communication` contains the files of a tutorial on how to instrument a code with calls to the OASIS3-MCT library in order to couple it with other components. The tutorial involves two toy model codes, `ocean.F90` and `atmos.F90`, which give you a practical example of the use of the OASIS3-MCT API (Application Program Interface) routines. Toy models are skeleton programs that do not contain any real physics or dynamics but that can reproduce real exchanges of coupling fields.

All information about this tutorial is provided in the document `tutorial.communication.pdf` therein.

### 6.3.2 spoc

This directory contains the sources used in the Short Online Private Course (SPOC) on “Code Coupling with OASIS3-MCT” developed in the framework of the ESiWACE Centre of Excellence. This SPOC is composed of videos, quizzes and hands-on. The goal is to instrument two toy models to set-up a real coupled model exchanging coupling fields (directory /spoc\_communication) and to learn more about OASIS3-MCT regridding functionality (directory /spoc\_regridding). If you are interested in attending the SPOC, please visit the online training section of CERFACS web site at <https://cerfacs.fr/en/online-training-sessions/> . Videos and quizzes extracted from the SPOC are also available as Open Education Resources (OER) material at <https://www.oercommons.org/courseware/lesson/85340> .

### 6.3.3 regrid\_environment

The `regrid_environment` directory offers a scripting environment to calculate the regridding weights and the regridding error for specific couple of grids and specific regridding algorithms with either the SCRIP library, ESMF or XIOS. The document `regrid_environment_documentation.pdf` therein contains all instructions on how to run this tutorial.

### 6.3.4 Fortran, C and python equivalent (pyoasis/examples)

Different examples implementing the Fortran, C and python APIs are provided as practical illustrations in directory `/pyoasis/examples` :

- `1-serial`: one coupling exchange between a serial sender and a serial receiver.
- `2-apple`: one coupling exchange between an Apple-parallel sender and a serial receiver; an additional component, not part of the coupling, is also started and the example shows how to use the `commworld` argument, in Fortran and C, and the communicator optional argument when setting the component in python.
- `3-box`: one coupling exchange between an Box-parallel sender and a serial receiver; it shows also how to check if a coupling field declared in the code is activated in the configuration file `namcouple`.
- `4-orange`: one coupling exchange between an Orange-parallel sender and a serial receiver; not all processes of the sender participate in the coupling and this example shows how to use `create_couple_comm`.
- `5-points`: one coupling exchange between a Point-parallel sender and a serial receiver.
- `6-apple_and_orange`: one coupling exchange between an Apple-parallel sender and an Orange-parallel receiver; not all processes of the sender participate in the coupling and this example shows how to use `set_couple_comm`.
- `7-multiple-puts`: two coupling fields are both sent from a serial sender to two different serial receivers; this example also sets up an intra communicator between the sender and one receiver and an inter communicator between the sender and the other receiver.
- `8-interopability/fortran_and_C`: implements a coupling of a bundle field, with two bundle elements, between a Fortran Apple-parallel sender and a C component. This C component is Orange-parallel for the reception of the bundle field; it also defines another partition of type Box onto which a second coupling field is defined and sent to a third Fortran serial receiver. The sum of the Box partitions in the C component does not cover the global grid, hence the fourth argument `ig_size` is used to specify the grid global size. The C component also illustrates how the order of the partition definition does not need to be the same for the different processes but that, in that case, a meaningful name fifth argument must be used.
- `8-interopability/fortran_and_python`: implements the same coupling exchanges than `8-interopability/fortran_and_C` but with the C component replaced by a python component.

- `9-python_fortran_C-multi_intracomm`: illustrates the set-up of an intracommunicator between a Fortran, a C and a python components using OASIS3-MCT; a bcst is then realised to share some data. In this example, an additional component is also launched at start but does not participate in the coupling and hence uses the `coupled` third argument of `oasis_init_comp`.
- `10-grid`: a single Box-parallel component defines and writes two grids `pyoa` and `mono`, the first one with distributed calls from all the processes, the second one from the master process only.
- `11-test-interpolation`: one exchange of a coupling bundle field defined on real grids involving a first-order conservative regridding with either `SCRIPR` or `YAC` between an Apple-parallel sender and a serial receiver. In the Fortran and C examples, the grids are fixed, while in the python example, the user chooses the source and target grids interactively, among the ones available in the files available in the `common_data` directory. This example produces graphical output of the received fields if the following packages are installed
  - `pip3 install matplotlib`
  - `pip3 install scipy`
  - `pip3 install cartopy`
  - `pip3 uninstall shapely`
  - `pip3 install shapely --no-binary shapely`
- `12-grid-functions`: Graphical version of `10-grid` (i.e. the `pyoa` grid layout is displayed if the same graphical packages than for `11-test-interpolation` are installed).
- `13-test-fracwgt` : illustrates the `fracwgt` option, see appendix A ; exchange of 3 coupling fields: the basic coupling field with its dynamic fractions as `fracwgt` argument, the dynamic fraction field *per se*, the scaled coupling field (i.e. the basic coupling field multiplied by its dynamic fractions).
- `14-test-inter3d` : illustrates the use of a cube partition, see section 2.2.3 .

The different examples can be launched with the Makefile from directory `/pyoasis` using targets `examples`, `examples.f` or `examples.c` to run respectively python, Fortran and C examples.

## 6.4 Debugging

### 6.4.1 Debug files

If you experience problems while running your coupled model, you can obtain more information in OASIS3-MCT debug files, by increasing the value of the first number on the line below `$NLOGPRT` in your `namcouple`, see section 3.2 for details.

### 6.4.2 Time statistics files

The level of time statistics written out in files named `*.timers_XXXX` depends on the value of the second number on the line below the `$NLOGPRT` keyword in the `namcouple` :

- 0 : timers are not activated, nothing is calculated, nothing is written.
- 1 : timers are activated and written in a single file by the process 0 as well as the min and the max times over all the processes.
- 2 : timers are activated and each process writes its own file ; process 0 also writes the min and the max times over all the processes in its file.
- 3 : timers are activated and each process writes its own file ; process 0 also writes in its file the min and the max times over all processes and also writes in its file all the results for each process.

The time given for each timer is calculated by the difference between calls to `oasis_timer_start()` and `oasis_timer_stop()` and is the accumulated time over the entire run. Here is an overview of the meaning of the different timers implemented by default.<sup>2</sup>

- `'total'` : total time of the simulation, implemented in `mod_oasis_method` (i.e. between the end of `oasis_init_comp` and the `mpi_finalize` in routine `oasis_terminate`).
- `'init_thru_enddef'` : time between the end of `oasis_init_comp` and the end of `oasis_enddef`, implemented in `mod_oasis_method`.
- `'part_definition'` : time spent in routine `oasis_def_partition`.
- `'oasis_enddef'` : time spent in routine `oasis_enddef`; this routine performs basically all the important steps to initialize the coupling exchanges, e.g. the internal management of the partition and variable definition, the definition of the patterns of communication between the source and target processes, the reading of the remapping weight-and-address file and the initialisation of the sparse matrix vector multiplication.
- `'grcv_00x'` : time spent in the reception of field `x` in `mct_recv` (including communication and possible waiting time linked to unbalance of components).
- `'wout_00x'` : time spent in the I/O for field `x` in routine `oasis_advance_run`.
- `'gcpy_00x'` : time spent in routine `oasis_advance_run` in copying the field `x` just received in a local array.
- `'pcpy_00x'` : time spent in routine `oasis_advance_run` in copying the local field `x` in the array to send (i.e. with local transformation besides division for averaging).
- `'pavg_00x'` : time spent in routine `oasis_advance_run` to calculate the average of field `x` (if done).
- `'pmap_00x'/'gmap_00x'` : time spent in routine `oasis_advance_run` for the matrix vector multiplication for field `x` on the source/target processes.
- `'psnd_00x'` : time spent in routine `oasis_advance_run` for sending field `x` (i.e. including call to `mct_waitsend` and `mct_isend`).
- `'wtrn_00x'` : time spent in routine `oasis_advance_run` to write fields associated with non-instant `loctrans` operations to restart files (see section 5.2 for details).
- `'wrst_00x'` : time spent in routine `oasis_advance_run` to write fields to restart files (see section 5.2 for details).

## 6.5 Load balancing analysis of coupled model components

An efficient use of the allocated computing resources in a coupled system requires the harmonisation of the component execution speed. This operation, called load balancing, is often neglected, either because of the apparent resource abundance or practical difficulties. To facilitate this work, a load balancing analysis functionality is included in OASIS3-MCT and can be activated by setting to 1 the third number under `$NLOGPRT` in the `namcouple` configuration file (see section 3.2). Some details on this functionality are provided here and more information can be found in the `balancing_documentation.pdf` file in `oasis3-mct/util/load_balancing` directory.

When activated, the load balancing analysis functionality provides general computing information (simulation time, speed, waiting time, etc.) in a text file `load_balancing_info.txt` for the coupled model and for each component; this global information can help to allocate resources in a balanced way.

This functionality also outputs the full timeline of all OASIS3-MCT related events, for any of the allocated resources. This timeline is saved in one NetCDF file per coupled component, `timeline_XXX.component.nc`

<sup>2</sup>Other measures can be obtained by defining the logical `local_timers_on` as `.true.` in different routines or by implementing other timers. Of course, OASIS3\_MCT and the model code then have to be recompiled.

where XXX is the component name. It provides the comprehensive sequence of all operations related to the coupling (field put and get through MPI, field output on disk, field interpolation and mapping, field reading on disk, restart writing, initialisation and termination phase of the OASIS3-MCT setup) so that any simulation slow down in link with the use of the OASIS3-MCT library can be identified. The analysis of the coupling field exchanges, amongst all coupling events, allows to not only identify the waste of resources by components which are recurrently waiting for their coupling fields but it also reveals other bottlenecks such as disk access or model internal load imbalance. The full picture of these events makes possible an optimal load balancing, even for the most complex configurations.

## Appendix A

# Conservative remapping with the `fracwgt` option

When a coupling field is valid only on a fraction of the source cells (fractional source field), there are different interpretations about the destination field that should be obtained on the destination grid after conservative remapping.

A **destination scaled field**, i.e. the remapping of the original field multiplied by its valid fraction on each cell, ensures conservation when the full destination cell area of the target grid is considered. However, because of the source fractions, it can become nonphysically small.

Another way to consider the problem is to calculate a **destination normalized scaled field** that will be in the physical range of the original field. This **destination normalized scaled field** will be valid only on a fraction of the target, the **active destination surface** that can be calculated with the **active destination surface coefficient**.

The new option `fracwgt` in OASIS3-MCT v6 allows to pass, in the source component as argument of the `oasis_put`, the active fraction of the source cells together with the source coupling field and to obtain, in the target component in the `oasis_get`, the **destination normalized scaled field**. It also allows to calculate the **active destination surface coefficient**.

For instantaneous fields, option `fracwgt` is not essential as the user could get the **active destination surface coefficient** simply with the conservative remapping of the source fraction field. He/she could then obtain the **destination normalized scaled field** by applying the element-wise division of the **destination scaled field** by the **active destination surface coefficient**.

However, for accumulated or averaged fields, option `fracwgt` is essential to obtain the **destination accumulated normalized scaled field** and the **accumulated active destination surface coefficients**, as they cannot be calculated in the models because of the non permutation of the time and remapping operations.

An example of how to use the `fracwgt` option is given in `/pyoasis/examples/13-test-fracwgt`.

### A.1 Introduction

Conservative remapping is relevant for physical fields representing a flux or a concentration.

We can think of energy fluxes ( $W/m^2$ ), precipitation rates ( $mm/m^2$ ), tracer concentrations (ppb or similar).

The remapping is required to:

1. preserve the total energy or mass
2. avoid abnormal local values of fluxes or concentrations

In models with no sub-grid fractions, the surface integral quantity to be preserved is the sum of the area of the unmasked source cells times the field on the cell

$$\sum_i A_{Si} F_{Si}$$

where  $A_{Si}$  and  $F_{Si}$  are respectively the area and the field of the source cell  $i$ . A conservative remapping will grant that

$$\sum_i A_{Si} F_{Si} = \sum_j A_{Dj} F_{Dj}$$

where  $A_{Dj}$  and  $F_{Dj}$ , the area and the field of the target cell  $j$ , can be expressed as:

$$F_{Dj} = \sum_i w_{ij} F_{Si}$$

$$A_{Dj} = \sum_i \omega_{ij} A_{Si}$$

$\omega_{ij}$ , the fraction of the source cell  $i$  overlapping the destination cell  $j$ , is related to the conservative remapping weight  $w_{ij}$  representing the area normalized fraction of the destination cell  $j$  intersected by the source cell  $i$  by, as in [Taylor 2024],:

$$w_{ij} = \omega_{ij} \frac{A_{Si}}{A_{Dj}}$$

Notice that, because of some assumptions on the representation of the cell edges on a sphere and to rounding errors, the  $A_{Dj}$  surface can slightly differ from the surface  $A_{Dj}^d$  prescribed in the destination model, but we will neglect this difference here.

## A.2 What is a fractional field?

For some models, a process represented on the model grid actually takes place only on a subset of it, whose boundaries are described at a finer resolution than the grid. The numerical translation of this choice is an extra mask-like field, providing the active (or unmasked) fraction  $f_i \in [0, 1]$  for every cell.

The field is therefore described with values ranging in the usual physical valid interval (*cf.* point 2. in the introduction), yet the integral quantity is scaled down to

$$\sum_i A_i f_i F_i$$

This scaling can be interpreted in two ways

1. On cells with usual area  $A_i$ , the flux is lowered to the **scaled field**  $F_i^f = f_i F_i$
2. The flux  $F_i$ , with values in the usual physical range, is active on a smaller **active source surface**  $A_i^a = A_i f_i$

## A.3 Instantaneous remapping

When applying the conservative remapping on an instantaneous fractional source field (i.e. in OASIS jargon, no `LOCTRANS` or `LOCTRANS/INSTANT` time transformation), we grant the conservation of the scaled integral source quantity

$$\sum_i A_{Si} f_{Si} F_{Si}$$

Once again, we can use different interpretations.

If we consider that we are interpolating a **scaled field**  $F_{S_i}^f = f_{S_i} F_{S_i}$  with a standard conservative remapping, and we are receiving the resulting **destination scaled field**

$$F_{D_j}^f = \sum_i w_{ij} f_{S_i} F_{S_i} \quad (\text{A.1})$$

then we will grant, as defined by the conservative remapping

$$\sum_i A_{S_i} f_{S_i} F_{S_i} = \sum_j A_{D_j} F_{D_j}^f$$

where  $A_{D_j}$  is the full destination cell area. Yet the local values  $F_{D_j}^f$  could locally become nonphysically small.

It is now essential to define the **active destination surface coefficients**, i.e. the active fraction of every destination cell,  $f_{D_j}$ , with the conservative remapping of the source grid fractions:

$$f_{D_j} = \sum_i w_{ij} f_{S_i} \quad (\text{A.2})$$

Using this coefficient  $f_{D_j}$ , we can define a **destination normalized scaled field** that will be in the physical range of the original field as

$$F_{D_j}^{fn} = \frac{F_{D_j}^f}{f_{D_j}} \quad (\text{A.3})$$

Note that if the active fraction of all source cells associated to one target cell  $f_{S_i}$  are zero then both  $F_{D_j}^f$  and  $f_{D_j}$  are zero and equation A.3 becomes undefined. In that case, in OASIS3-MCT v6, we set  $F_{D_j}^{fn}$  equal to zero.

In order to preserve the integral quantity we have to scale back the field:

$$\sum_i A_{S_i} f_{S_i} F_{S_i} = \sum_j A_{D_j} f_{D_j} F_{D_j}^{fn}$$

For computing a conservative integral quantity on the destination side, the active destination surface coefficient  $f_{D_j}$  can be used to define the **active destination surface** :

$$A_{D_j}^a = A_{D_j} f_{D_j} \quad (\text{A.4})$$

so that

$$\sum_i A_{S_i} f_{S_i} F_{S_i} = \sum_i A_{S_i}^a F_{S_i} = \sum_j A_{D_j}^a F_{D_j}^{fn}$$

Since OASIS3-MCT v6, the `oasis_put` routine comes with an extra optional argument `fracwgt`. If the argument is present, it has to correspond to the active source fraction field  $f_S$ , and the matching `oasis_get` will receive the normalized scaled field  $F_D^{fn}$ .

Notice that for technical reasons, the active destination surface coefficient  $f_D$  cannot be recovered as an extra output of the `oasis_get` call. The user can recover it in two ways:

1. send the active source fraction field  $f_S$  as a separate coupling field through a standard conservative remapping and directly get  $f_D$  (equation A.2) ; or
2. compute the ratio between the destination scaled field and the destination normalized scaled field
  - (a) in the source model compute the scaled source field  $F_S^f = f_S F_S$
  - (b) send this scaled field with a standard `oasis_put` call (i.e. without a `fracwgt` argument) and on the destination side store the corresponding (non normalized) scaled field  $F_D^f$

- (c) send the (non scaled) source field  $F_S$  passing `fracwgt=f_S` to the `oasis_put` call and on the destination side store the corresponding normalized scaled field  $F_D^{fn}$  (equation A.3)
- (d) compute the active destination surface coefficient as an element-wise division  $f_D = F_D^f \odot \frac{1}{F_D^{fn}}$

Once  $f_D$  is computed, the active destination surface to be used for computing integral quantities is finally obtained by scaling the destination cells surfaces  $A_D^a = A_D \odot f_D$

Notice that it is mandatory that the treatments described in the `namcouple` for the two remappings b. and c. are strictly the same.

Even if the implementation of the first method is easier for instantaneous fields, we suggest using the second one that relies on the `fracwgt` option because, as we will show in the following sections, it is compatible with time accumulated or time averaged remapping.

## A.4 Time accumulation

Let's now consider the case in which the destination field has to be computed as the accumulation of remapped source fields at different times (i.e. in OASIS jargon, `LOCTRANS/ACCUMUL` time transformation)

$$F_{Dj}^+ = \sum_t \alpha F_{Dj}^t + \beta = \sum_t \alpha \left( \sum_i w_{ij} F_{Si}^t \right) + \beta \quad (\text{A.5})$$

where  $\alpha$  and  $\beta$  are the coefficient of an affine transform that can be indicated with the `BLASOLD` method. Because of the linearity of the operators we can neglect them in the following for the sake of readability. We therefore define the **destination accumulated field** as

$$F_{Dj}^+ = \sum_t F_{Dj}^t = \sum_t \sum_i w_{ij} F_{Si}^t \quad (\text{A.6})$$

In the case where no subgrid fractions are used, the accumulating sum and the linear mapping can permute and we can easily obtain the **destination accumulated field** as the remapping of the **accumulated source fields**

$$F_{Dj}^+ = \sum_t \sum_i w_{ij} F_{Si}^t = \sum_i w_{ij} \sum_t F_{Si}^t \quad (\text{A.7})$$

In that case, we can rewrite the conservative remapping as

$$\sum_t \sum_i A_{Si} F_{Si}^t = \sum_t \sum_j A_{Dj} F_{Dj}^t = \sum_j A_{Dj} \sum_t F_{Dj}^t = \sum_j A_{Dj} F_{Dj}^+ \quad (\text{A.8})$$

If we introduce a time varying fractional coefficient in the time accumulation equations, we can obtain the corresponding **destination accumulated scaled field** by sending and remapping  $f_{Si}^t F_{Si}^t$ :

$$F_{Dj}^{+f} = \sum_i w_{ij} \sum_t f_{Si}^t F_{Si}^t \quad (\text{A.9})$$

In that case, the conservation equation is satisfied when the full destination area is used

$$\sum_t \sum_i A_{Si} f_{Si}^t F_{Si}^t = \sum_i A_{Si} \sum_t f_{Si}^t F_{Si}^t = \sum_j A_{Dj} F_{Dj}^{+f} \quad (\text{A.10})$$

However,  $F_{Dj}^{+f}$  will not be physically consistent with the source field because it has been scaled by the source fractions. If we want to preserve the physical range of the destination field, we should compute the

**destination accumulated normalized scaled field** that, given (A.1) and (A.2) for each timestep, can be written as:

$$F_{Dj}^{+fn} = \sum_t \frac{\sum_i w_{ij} f_{Si}^t F_{Si}^t}{\sum_i w_{ij} f_{Si}^t} = \sum_t \frac{F_{Dj}^{ft}}{f_{Dj}^t} \quad (\text{A.11})$$

Note that, as for equation A.3, if for any timestep the active fraction of all source cells associated to one target cell  $f_{Si}^t$  are zero then both  $F_{Dj}^{ft}$  and  $f_{Dj}^t$  are zero and equation A.11 becomes undefined. In that case, in OASIS3-MCT v6, we set  $F_{Dj}^{+fn}$  equal to zero.

Notice that in equation A.11, the summation on the source grid cells does not permute with the summation in time and the summation of the  $N_t$  time contribution has to be carried on on the destination grid after remapping: it requires  $2N_t$  sparse multiplications by the weight matrix.

Once again we can derive the expression of the **active surface for every destination cell**  $A_{Dj}^a$  by imposing

$$A_{Dj}^a F_{Dj}^{+fn} = A_{Dj} F_{Dj}^{+f}$$

hence

$$A_{Dj}^a = A_{Dj} \frac{F_{Dj}^{+f}}{F_{Dj}^{+fn}} \quad (\text{A.12})$$

which corresponds to the definition of **accumulated active destination surface coefficients**:

$$f_{Dj}^+ = \frac{F_{Dj}^{+f}}{F_{Dj}^{+fn}} \quad (\text{A.13})$$

Since the user cannot recover it directly, the computation of  $f_D^+$  requires the following steps:

1. at every put time, in the source model compute the scaled source field  $F_S^f = f_S F_S$  and send this scaled field through a standard `oasis_put` call (i.e. without a `fracwgt` argument) ;
2. send the (non scaled) source field  $F_S$  passing `fracwgt=f_S` to the `oasis_put` call;
3. on the destination side receive and store the (non normalized) accumulated scaled field  $F_D^{+f}$  (equation A.9) corresponding to the fields sent in 1 ;
4. on the destination side receive and store the accumulated normalized scaled field  $F_D^{+fn}$  (equation A.11) corresponding to the fields sent in 2 ;
5. compute the accumulated active destination surface coefficients as an element-wise division  $f_D^+ = F_D^{+f} \odot \frac{1}{F_D^{+fn}}$  (equation A.13).

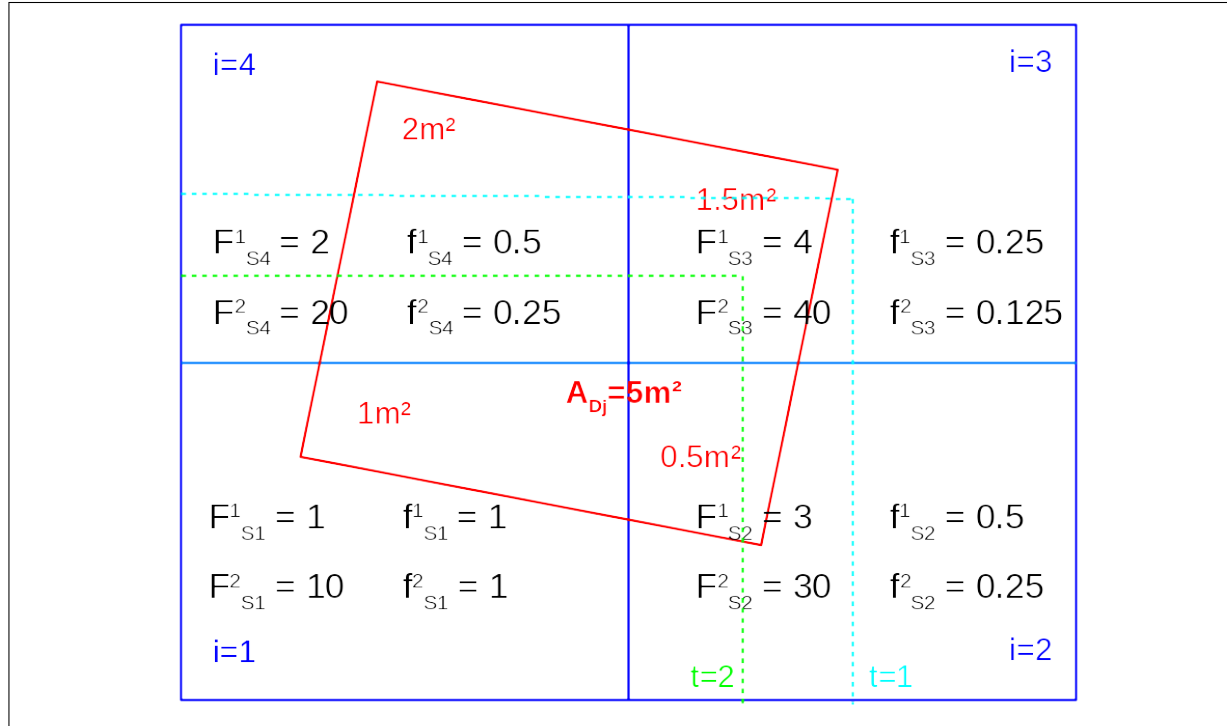
Notice that it is mandatory that the treatments described in the `namcouple` for the two remappings 1.→3. and 2.→4. are strictly the same.

## A.5 Time average

In the case of a time average with individual contributions at equal intervals, (i.e. in OASIS jargon, LOCTRANS/AVERAGE time transformation) the average is obtained by dividing the time accumulated fields by the number of contributions  $N_t$ . It's easy to see that the coefficient  $1/N_t$  can permute with all the time summations and that the active fraction on the destination grid is the same as in the time accumulation case.

$$\bar{f}_{Dj} = \frac{\bar{F}_{Dj}^f}{\bar{F}_{Dj}^{fn}} = \frac{F_{Dj}^{+f}/N_t}{F_{Dj}^{+fn}/N_t} = \frac{F_{Dj}^{+f}}{F_{Dj}^{+fn}} = f_{Dj}^+$$

The case with unevenly spaced time contributions would be more complicated, but it is not implemented in OASIS3-MCT v6.



**Figure A.1:** Source and destination grids for the test case

## A.6 An example

Let's consider a destination cell with total surface of 5 sq. mt. intersecting four source cells by, respectively, 1, 0.5, 1.5 and 2 sq. mt as in Fig.A.1

The conservative remapping weights are therefore:

$$w_1 = 0.2; w_2 = 0.1; w_3 = 0.3; w_4 = 0.4$$

At time 1, the active source surface covers the following fractions:

$$f_{S1}^1 = 1.0; f_{S2}^1 = 0.5; f_{S3}^1 = 0.25; f_{S4}^1 = 0.5$$

and the source fields is:

$$F_{S1}^1 = 1.0; F_{S2}^1 = 3.0; F_{S3}^1 = 4.0; F_{S4}^1 = 2.0$$

If we neglected the sub-grid fractions, the remapped field at time 1 would be

$$F_{Dj}^1 = \sum_i w_i F_{Si}^1 = 2.5$$

and it's contribution to the integral quantity on the destination grid would be

$$A_{Dj} F_{Dj}^1 = 12.5$$

If we take into consideration the source fractions, the destination scaled field as in eqn. (A.1) is

$$F_{Dj}^{1f} = \sum_i w_i f_{Si}^1 F_{Si}^1 = 1.05$$

It's contribution to the integral quantity on the destination grid is reduced to:

$$A_{Dj} F_{Dj}^{1f} = 5.25$$

Taking into consideration the active destination surface coefficients as in eqn. (A.2)

$$f_{Dj}^1 = \sum_i w_i f_{Si}^1 = 0.525$$

We can get the destination normalized scaled field as in eqn. (A.3)

$$F_{Dj}^{1fn} = \frac{F_{Dj}^{1f}}{f_{Dj}^1} = 2.0$$

and the active destination surface as in eqn. (A.4)

$$A_{Dj}^{1a} = A_{Dj} f_{Dj}^1 = 2.625$$

and find again the appropriate contribution to the integral quantity on the destination grid

$$A_{Dj}^{1a} F_{Dj}^{1fn} = 5.25$$

Let's now suppose that at time 2, the active source surface covers the following fractions:

$$f_{S1}^2 = 1.0; f_{S2}^2 = 0.25; f_{S3}^2 = 0.125; f_{S4}^2 = 0.25$$

and the source fields is:

$$F_{S1}^2 = 10.0; F_{S2}^2 = 30.0; F_{S3}^2 = 40.0; F_{S4}^2 = 20.0$$

The instantaneous destination scaled field will then be:

$$F_{Dj}^{2f} = \sum_i w_i f_{Si}^2 F_{Si}^2 = 6.25$$

It's contribution to the integral quantity:

$$A_{Dj} F_{Dj}^{2f} = 31.25$$

The active destination surface coefficient:

$$f_{Dj}^2 = \sum_i w_i f_{Si}^2 = 0.3625$$

The destination normalized scaled field:

$$F_{Dj}^{2fn} = \frac{F_{Dj}^{2f}}{f_{Dj}^2} = 17.2414$$

and the corresponding active destination surface :

$$A_{Dj}^{2a} = A_{Dj} f_{Dj}^2 = 1.8125$$

which confirms the appropriate contribution:

$$A_{Dj}^{2a} F_{Dj}^{2fn} = 31.25$$

If we accumulate the field on the two time steps, neglecting the sub-grid fractions we obtain

$$F_{S1}^+ = 11.0; F_{S2}^+ = 33.0; F_{S3}^+ = 44.0; F_{S4}^+ = 22.0$$

and the destination accumulated field would be (equation A.7)

$$F_{Dj}^+ = \sum_i w_i F_{Si}^+ = 27.5$$

Meanwhile, if we accumulate the scaled source fields we have

$$F_{S1}^{+f} = 11.0; F_{S2}^{+f} = 9.0; F_{S3}^{+f} = 6.0; F_{S4}^{+f} = 6.0$$

and the destination accumulated scaled field (equation A.9) has a much lower value:

$$F_{Dj}^{+f} = \sum_i w_i F_{Si}^{+f} = 7.3$$

It's contribution to the integral quantity on the destination grid is:

$$A_{Dj}F_{Dj}^{+f} = 36.5$$

We can obtain a more representative value by accumulating the instantaneous normalized scaled fields as in eqn. (A.11)

$$F_{Dj}^{+fn} = \sum_t \frac{F_{Dj}^{ft}}{f_{Dj}^t} = 19.24137931$$

and derive its equivalent active destination cell fraction from eqn. (A.13)

$$f_{Dj}^+ = \frac{F_{Dj}^{+f}}{F_{Dj}^{+fn}} = 0.379390681$$

and an active surface:

$$A_{Dj}^a = A_{Dj}f_{Dj}^+ = 1.896953405$$

Since the average values are just the half sum of the instantaneous values, the fractions are the same as in the accumulation case.

The averaged scaled source fields on the four cells are

$$\bar{F}_{S1}^f = 5.5; \bar{F}_{S2}^f = 4.5; \bar{F}_{S3}^f = 3.0; \bar{F}_{S4}^f = 3.0$$

The scaled remapped field is:

$$\bar{F}_{Dj}^f = \sum_i w_i \bar{F}_{Si}^f = 3.65$$

It's contribution to the integral quantity on the destination grid is:

$$A_{Dj}\bar{F}_{Dj}^f = 18.25$$

By averaging the normalized remapped values we obtain the more representative averaged field

$$\bar{F}_{Dj}^{fn} = 9.620689655$$

and derive its equivalent active destination cell fraction

$$\bar{f}_{Dj} = \frac{\bar{F}_{Dj}^f}{\bar{F}_{Dj}^{fn}} = 0.379390681$$

which is the same as in the accumulation case and implies an active surface:

$$A_{Dj}^a = A_{Dj}\bar{f}_{Dj} = 1.896953405$$

## Appendix B

# Changes between the different versions of OASIS3-MCT

The evolution between the different versions of OASIS3-MCT can be followed in real-time by registering on the Redmine project management site at <https://inle.cerfacs.fr/> (see "Register" at the right top of the page). On this site, registered users can browse the sources and consult tickets describing bug fixes and developments. To follow day to day evolution of the OASIS3-MCT sources, it is also possible to have your e-mail added to the list of addresses to which the log files of the SVN checkins are automatically sent; contact [oasishelp@cerfacs.fr](mailto:oasishelp@cerfacs.fr) if you wish to be added to that list.

### B.1 Changes between OASIS3-MCT v6 and OASIS3-MCT\_5.0

The last version of the coupler, OASIS3-MCT v6, comes with the following new functionalities and improvements :

- **YAC interpolation library** : interfacing with the parallel remapping algorithms for 2D interpolation on the sphere from version 3.10 of the YAC coupler [Hanke et al 2016] [Hanke and Redler 2019] all details are provided in section 4.3.2 and in [Jonville et al 2024].
- **Support of dynamic fractions**, see `fracwgt` option in section 2.2.7 and in appendix A.
- An **upgraded compiling environment** based on **two environment variables** `$(OASIS_COUPLE)` and `$(OASIS_ENV)`, see section 6.1.
- **Support of a new "Cube" partition**, see section 2.2.3, adapted to specific couplings, e.g. horizontal coarsening in the exchanges between a high-resolution ocean model and a lower-resolution bio-geo-chemistry model.
- Suppression of big NetCDF files in the git repository and its history in order to drastically reduce its size, see Redmine issue #2907
- Migration of the OASIS3-MCT web site from DKRZ (content manager Plone) to Cerfacs (content manager Wordpress) in July 2022.
- Migration of the CI/CD (Continuous Integration and Continuous Deployment) environment from Buildbot to gitlab .
- Different bug fixes, see the following Redmine closed issues :
  - #3098, an array bound mismatch in `mod_oasis_load_balancing.F90` ;
  - #2900, a bug with partition duplication in case the grid dimensions were not declared in the `namcouple` ;
  - #2841, the support of multiple grids for one same partition (linked with #2900) ;

- #2816, a freeze in the load balancing analysis when a component was not at all involved in the coupling ;
- #2812, an uninitialized array in grid writing routines ;
- #2793, wrong *namcouple* processing for BLASNEW ;
- #2725, a problem with BICUBIC/BILINEAR sequence in weights generation ;
- #2704, a problem with `coupled = .false.` in `oasis_init_comp`.

## B.2 Changes between OASIS3-MCT\_5.0 and OASIS3-MCT\_4.0

The last version of the coupler, OASIS3-MCT\_5.0, comes with the following novelties:

- Python, C and C++ bindings, see sections 2.3, 2.4 and 6.3.4
- A new load balancing analysis tool, see section 6.5
- An environment to use either SCRIP, ESMF or XIOS to generate regridding weights and to analyse the quality of the regridding, see directory `examples/regrid_environment` and section 6.3.3
- A new locally conservative remapping, to be used in particular for runoffs, see LOCCUNIF, LOCCDIST and LOCCGAUS in section 4.3
- Extension of BLASNEW operation to support combination of coupling fields, see section 4.4
- Improved and additional diagnostics in CHECKIN and CHECKOUT, see sections 4.2 and 4.4
- SCRIPR/CONSERV option for normalisation by the true area of the grid cells, see DESTARTR, FRACARTR or FRACNNTR in section 4.3
- New GSSPOS and BSSPOS options for global CONSERV (as GLBPOS and BASPOS options respectively except that the multiplicative term is computed separately for positive and negative values of the field), see section 4.4
- Extension of `oasis_get_intracomm` to support multiple components, see `oasis_get_multi_intracomm` in section 2.2.9
- Communication/exchange of simple scalars, see section 2.5.1
- Update examples in `oasis3-mct/examples` directory, see section 6.3
- Update to MCT 2.11
- Update of compiling environment, see 6.1
- Migration from SVN to GIT for source management
- Migration of the OASIS3-MCT web site from DKRZ (content manager Plone) to Cerfacs (content manager Wordpress), see the new site at <https://oasis.cerfacs.fr/en/>
- For DISTWGT, DISTWGTF, GAUSWGT, GAUSWGTF LOCCUNIF, LOCCDIST or LOCCGAUS, inclusion of the number of neighbours used in the remapping file name
- For SCRIPR/CONSERV, specification through the *namcouple* of North thresh `$NTHRESH` and South thresh `$STHRESH` values, above which a Lambert projection is activated
- New `$NCDFTYP` NetCDF file format setting through the *namcouple*, see section 3.2
- Overload `oasis_def_var` interface to support excluding the argument `id_var_shape` from the argument list
- Systematic tests of NetCDF returned error code
- Update `oasis_abort` to also write to unit 0 (stderr)
- Bugfixes
  1. Component name argument in `oasis_get_intercomm` (ticket #2776)
  2. BICUBIC/BILINEAR sequence in weights generation (ticket # 2725)
  3. Array bounds check for `cplfind` (ticket # 2655)

4. Fix error in maxloops/kfac computation in `oasis_mpi_reducelists` (ticket # 2654)
5. Exact calculation of local distance in `GAUSWGT` interpolation (ticket #2500)
6. Compatibility of load balancing analysis tool with CRAY compiler (ticket # 2473)
7. Problem with `SCRIPR/BILINEAR` periodicity (ticket # 2419)
8. Problem in `m_MCTWorld.F90` in the routine `initm` (ticket # 1321)
9. Argument `id_var_nodim` defined as `IN` in `mod_oasis_var.F90` to compile with `NEMO_4.0` and `NEMO` trunk (git commit 28f4fe59)
10. In `SCRIPR/CONSERV`, condition on coincidence of segments is reinitialised to false for each cell (bugfix “lcoinc”; see [Valcke and Piacentini 2019])

### B.3 Changes between OASIS3-MCT\_4.0 and OASIS3-MCT\_3.0

Different developments were realised to improve the parallel performance of OASIS3-MCT\_4.0. These developments are detailed in [Valcke et al 2018].

- A new communication method, using the remapping weights to define the intermediate mapping decomposition, offers a significant gain at run time, especially for high-resolution cases running on a high number of tasks, thanks to reduced communication. However, as expected, the new method takes longer to initialize, partly due to the fact that the mapping weight file has to be read twice but also due to the extra cost for the initialization of the different MCT routers. That initialization cost is largely mitigated by an upgrade to MCT 2.10.beta1 which reduces the penalty to few seconds. Generally, it should be worth the extra initial cost to speed up the run time. See `$NMAPDEC` in section 3.2.
- The hybrid MPI+OpenMP parallelisation of the `SCRIP` library (previously fully sequential) leads to great improvement in the calculation of the remapping weights. The results obtained here show a reduction in the weight calculation time of 2 to 3 orders of magnitude with the new parallel `SCRIP` library for high-resolution grids. Details are available in [Piacentini et al 2018]. The `test_interpolation` environment gives a practical example on how to use OASIS3-MCT\_4.0 to pre-calculate (i.e. in a separate job prior to the “real” simulation) the remapping weight and address file.

Thanks to some preliminary work, few bugs were fixed, in particular in the bounding box definition of the grid cells. This solves an important bug observed in the Pacific near the equator for the bilinear and bicubic interpolations for Cartesian grids.

However, given these modifications, one cannot expect to get exactly the same results for the interpolation weight-and-address remapping files with this new parallel `SCRIP` version as compared to the previous `SCRIP` version in OASIS3-MCT\_3.0. We checked in many different cases that the interpolation error is smaller or of the same order than before. We also observed that the parallelisation does not ensure bit reproducible results when varying the number of processes or threads.

- The new methods introduced in the global `CONSERV` operation reduce its calculation costs by one order of magnitude while still ensuring an appropriate level of reproducibility. This removes the bottleneck foreseen at high resolution with this important, and in few cases still unavoidable, global operation. These new methods are detailed in section 4.4.

The other new features offered by OASIS3-MCT\_4.0 are the following:

- Support for bundled coupling fields.  
Bundled fields is now supported in the `oasis_put` and `oasis_get` interfaces to allow easier coupling of multi-level or other related fields via a single `namcouple` coupling definition and a single call to `oasis_put` or `oasis_get`. Further details are provided in sec 2.2.7
- Automatic coupling restart writing

An optional argument `write_restart` was added to the `oasis_put` routine. This argument is false by default but if it is explicitly set to true in the code, a coupling restart file will be written for that field only for that coupling timestep, saving the data that exists at the time of the call (see section 2.2.7).

- Exact consistency between the number of weights and fields

Exact consistency is now required between number of weights fields in the coupling restart file and the arrays passed as arguments to the `oasis_put` routine. For example, for a 2nd order conservative remapping (CONSERV SECOND), 3 weights are needed and 3 fields must be provided as arguments: the value of the field, its gradient with respect to the longitude and its gradient with respect to the latitude. For a first order conservative remapping (CONSERV FIRST), only one weight and one field are needed. Using a weight file with 3 weights for a first order conservative remapping is no longer allowed.

- Upgrade in the `namcouple` configuration file

The `namcouple` reading routine was cleaned up including a refactoring of the `gotos` and `continue` statements, addition of few reusable routines including an abort routine, removal of some dead code, addition of support for blank lines (which are now considered comments), removal of requirement that keywords start at character 2 on a line, removal of requirement for `$END` in the `namcouple`, and updates to some error messages.

- Other new functionalities with corresponding new `namcouple` keywords (see section 3.2)
  - `$NUNITNO`: specifies the minimum and maximum unit numbers to be used for input and output files in the coupling layer.
  - `$NMATXRD`: indicates the method used to read mapping weights, either `orig` or `ceg`. In both methods, the weights are read in chunks by the model master task. With the `orig` option, the weights are then broadcast to all other tasks and each task then saves the weights that will be applied to its grid points. With the `ceg` option, the master task reads the weights and then identifies to which other task each weight should be sent. A series of exchanges are then done with each other task involving just the weights needed by that other task. The `orig` method sends much more data but is more parallel, while the `ceg` method does most of the work on the master task but less data is communicated.
  - `$NWGTOPT`: indicates how to handle bad interpolation weights.
  - `$NNOREST` : if true, OASIS3-MCT will initialise any variable that normally requires a coupling restart file with zeros if that file does not exist.

## B.4 Changes between OASIS3-MCT\_3.0 and OASIS3-MCT\_2.0

The main evolution of OASIS3-MCT\_3.0 with respect to OASIS3-MCT\_2.0 is the support of coupling exchanges between parallel components deployed in much more diverse configurations than before, for example, within one same executable between components running concurrently on separate sets of tasks or between components running sequentially on overlapping sets of tasks. All details are provided in section 2.1.

This new version also includes:

- memory and performance upgrades
- a new LUCIA tool for load balancing analysis
- new memory tracking tool (gilt)
- improved error checking and error messages
- doxygen documentation
- expanded test cases and testing automation

- testing at high resolution ( $\geq 1M$  gridpoints), high processor counts (32k pes), and with large variable counts ( $\geq 1k$  coupling fields)
- many bug fixes

## B.5 Changes between OASIS3-MCT\_2.0 and OASIS3-MCT\_1.0

The main changes and bug fixes new in OASIS3-MCT\_2.0 are the following:

- Support of BICUBIC interpolation, see paragraph BICUBIC in section 4.3. If the source grid is not a gaussian reduced grid (D), the gradient in the first dimension, the gradient in the second dimension, and the cross-gradient of the coupling field must be calculated by the model and given as arguments to `oasis_put`, as explained in section 2.2.7. If the source grid is a gaussian reduced grid (D), OASIS3-MCT\_2.0 can calculate the interpolated field using only the values of the source field points.
- Support of CONSERV/SECOND remapping, see paragraph CONSERV/SECOND in section 4.3.
- Support of components exchanging data on only a subdomain of the global grid: a new optional argument, `ig_size` was added to `oasis_def_partition`, that provides the user with the ability to define the total number of grid cells on the grid (see section 2.2.3).
- The variable `TIMER_Debug` controlling the amount of time statistics written out is now an optional argument read in the *namcouple*; see the `NLOGPRT` line in 3.2 and all details about time statistics in section 6.4.2.
- Specific problems in writing out the time statistics when all the processors are not coupling were solved (see Redmine issue #497)
- The problem with restart files when one coupling field is sent to 2 target components was solved (see Redmine ticket #522)
- A memory leak in `mod_oasis_getput_interface.F90` was fixed thanks to R. Hill from the MetOffice (see Redmine ticket #437)
- A bug fix was provided to ensure that the nearest neighbour option is activated when the option `FRACNNEI` is defined in the *namcouple* for the conservative interpolation .
- The behaviour of OASIS3-MCT was changed in the case a component model tries to send with `oasis_put` a field declared with a `oasis_def_var` but not defined in the configuration file *namcouple*; this will now lead to an abort. In this case, the field ID returned by the `oasis_def_var` is equal to -1 and the corresponding `oasis_put` should not be called. Conversely, all coupling fields appearing in the *namcouple* must be defined with a call to `oasis_def_var`; this constraint is imposed to avoid that a typo in the *namcouple* would lead to coupling exchanges not corresponding to what the user intends to activate.
- OASIS3-MCT developments are now continuously tested and validated on different computers with a test suite under Buildbot, which is a software written in Python to automate compile and test cycles required in software project (see <https://inle.cerfacs.fr/projects/oasis3-mct/wiki/Buildbot> on the Redmine site).

## B.6 Changes between OASIS3-MCT\_1.0 and OASIS3.3

### B.6.1 General architecture

- OASIS3-MCT is (only) a coupling library

Much of the underlying implementation of OASIS3 was refactored to leverage the Model Coupling Toolkit (MCT). OASIS3-MCT is a coupling library to be linked to the component models and that carries out the coupling field transformations (e.g. remappings/interpolations) in parallel on either

the source or target processes and that performs all communication in parallel directly between the component models; there is no central coupler executable anymore<sup>1</sup>.

- MAPPING transformation to use a pre-defined mapping file

With MAPPING, OASIS3-MCT has the ability to read a predefined set of weights and addresses (mapping file) specified in the *namcouple* to perform the interpolation/remapping. The user also has the flexibility to choose the location and the parallelization strategy of the remapping with specific MAPPING options (see section 4.3).

- Mono-process mapping file generation with the SCRIP library

But as before, OASIS3-MCT\_1.0 can also generate the mapping file using the SCRIP library [Jones 1999]. When this is the case, the mapping file generation is done on one process of the model components; all previous SCRIPR remapping schemes available in OASIS3.3 are still supported besides BICUBIC and CONSERV/SECOND. (Note: these remapping schemes, not available in OASIS3-MCT\_1.0 were reactivated in OASIS3-MCT\_2.0, see B.5.)

- MPI2 job launching is NOT supported.

Only MPI1 start mode is allowed. As before with the MPI1 mode, all component models must be started by the user in the job script in a pseudo-MPMD mode; in this case, they will automatically share the same MPI\_COMM\_WORLD communicator and an internal communicator created by OASIS3-MCT needs to be used for internal parallelization (see section 2.2.2).

## B.6.2 Changes in the coupling interface in the component models

- Use statement

The different OASIS3.3 USE statements were gathered into one USE *mod\_oasis* (or one USE *mod\_prism*), therefore much simpler to use.

- Support of previous *prism\_xxx* and new *oasis\_xxx* interfaces

OASIS3-MCT retains prior interface names of OASIS3.3 (e.g. *prism\_put\_proto*) to ensure full backward compatibility. However, new interface names such as *oasis\_put* are also available and should be preferred.

- Auxiliary routines not supported yet

Auxiliary routines *prism\_put\_inquire*, *prism\_put\_restart\_proto*, *prism\_get\_freq* are not supported yet. (Note: *prism\_put\_inquire* and *prism\_get\_freqs* were reintroduced in OASIS3-MCT\_3.0 and equivalent of *prism\_put\_restart\_proto* in OASIS3-MCT\_4.0.)

- Support of components for which only a subset of processes participate in the coupling

New routines *oasis\_create\_couplcomm* and *oasis\_set\_couplcomm* are now available to create or set a coupling communicator in the case only a subset of the component processes participate in the coupling. But even in this case, all OASIS3-MCT interface routines, besides the grid definition (see section 2.2.4) and the “put” and “get” call per se (see section 2.2.7), are collective and must be called by all processes. (Note: this has changed with OASIS3-MCT\_3.0.)

- New routines *oasis\_get\_debug* and *oasis\_set\_debug*

New routines *oasis\_get\_debug* and *oasis\_set\_debug* are now available to respectively retrieve the current OASIS3-MCT internal debug level (set by *\$NLOGPRT* in the *namcouple*) or to change it (see section 2.2.9).

---

<sup>1</sup>As with OASIS3.3, the “put” calls are non-blocking but the “get” calls are blocking. As before, the user has to take care of implementing a coupling algorithm that will result in matching “put” and “get” calls to avoid deadlocks (see section 2.2.7). The lag (LAG) index works as before in OASIS3.3 (see section 2.5.2)

### B.6.3 Functionality not offered anymore

- SCRIPR/BICUBIC and SCRIPR/CONSERV/SECOND remappings

As in OASIS3.3, the SCRIP library can be used to generate the remapping/interpolation weights and addresses and write them to a mapping file. All previous SCRIPR remapping schemes available in OASIS3.3 are still supported in OASIS3-MCT\_1.0 besides BICUBIC and CONSERV/SECOND because these remapping involve at each source grid point the value of the field but also the value of the gradients of the field (which are not known or calculated). (Note: these remapping schemes, not available in OASIS3-MCT\_1.0 were reactivated in OASIS3-MCT\_2.0, see B.5.)

- Vector field remapping

Vector field remapping is not and will not be supported (see “Support of vector fields with the SCRIPR remappings” in section 4.3).

- Automatic calculation of grid mesh corners in SCRIPR/CONSERV

For SCRIPR/CONSERV remapping, grid mesh corners will not be compute automatically if they are needed but not provided.

- Other transformations not supported

- The following transformations are not available in OASIS3.3 and will most probably not be implemented as it should be not too difficult to implement the equivalent operations in the component models themselves: CORRECT, FILLING, SUBGRID, MASKP
- LOCTRANS/ONCE is not explicitly offered as it is equivalent to defining a coupling period equal to the total runtime.
- The following transformations are not available as they were already deprecated in OASIS3.3 : REDGLO, INVERT, REVERSE, GLORED
- MASK and EXTRAP are not available but the corresponding linear extrapolation can be replaced by the more efficient option using the nearest non-masked source neighbour for target points having their original neighbours all masked. This is now the default option for SCRIPR/DISTWGT, GAUSWGT and BILINEAR interpolations. It is also included in SCRIPR/CONSERV if FRACNNEI normalization option is chosen (see section 4.3).
- INTERP interpolations are not available; SCRIPR should be used instead.
- MOZAIC is not available as MAPPING should be used instead.
- NOINTERP does not need to be specified anymore if no interpolation is required.
- Field combination with BLASOLD and BLASNEW; these transformations only support multiplication and addition terms to the fields (see section 4.2).

- Using the coupler in interpolator-only mode

This is not possible anymore as OASIS3-MCT is now only a coupling library. However, it is planned, in a further release, to provide a toy coupled model that could be use to check the quality of the remapping for any specific couple of grids. (Note: this was done in OASIS3-MCT\_2.0.)

- Coupling field CF standard names

The file cf\_name\_table.txt is not needed or used anymore. The CF standard names of the coupling fields are not written to the debug files.

- Binary auxiliay files

All auxiliary files, besides the *namcouple* must be NetCDF; binary files are not supported anymore.

### B.6.4 New functionality offered

- Better support of components for which only a subset of processes participate in the coupling

In OASIS3.3, components for which only a subset of processes participated in the coupling were supported in a very restricted way. In fact, this subset had to be composed of the N first processes and N had to be specified in the *namcouple*. Now, the subset of processes can be composed of any of the component processes and does not have to be pre-defined in the *namcouple*. New routines `oasis_create_couplcomm` and `oasis_set_couplcomm` are now available to create or set a coupling communicator gathering only these processes (see section 2.2.2). (Note: this was further improved in OASIS3-MCT\_3.0.)

- Exact restart for LOCTRANS transformations

If needed, LOCTRANS transformations write partially transformed fields in the coupling restart file at the end of a run to ensure an exact restart of the next run (see section 4.1). For that reason, coupling restart filenames are now required for all *namcouple* entries that use LOCTRANS (with non INSTANT values). This is the reason an optional restart file is now provided on the OUTPUT *namcouple* input line. If the coupling periods of two (or more) coupling fields are different, it is necessary to define two (or more) restart files, one for each field.

- Support to group multiple fields in a single communication.

This is supported through colon delimited field lists in the *namcouple*, for example

```
ATMTAUX:ATMTAUY:ATMHFLUX TAUX:TAUY:HEATFLUX 1 3600 3 rstrt.nc EXPORTED
```

in a single *namcouple* entry. All fields in this group will use the *namcouple* settings for that entry. In the component model codes, these fields are still sent (“put”) or received (“get”) one at a time. Inside OASIS3-MCT, the fields are stored and a single mapping and send or receive instruction is executed for all fields. This is useful in cases where multiple fields have the same coupling transformations and to reduce communication costs by aggregating multiple fields into a single communication. If a LOCTRAN transformation is needed for these multiple fields, it is necessary to define a restart file for these multiple fields. The coupling fields must be sent and received in the same order as they are defined in the corresponding single entry of the *namcouple* (not relevant in further versions of OASIS3-MCT).

- Matching one source field with multiple targets

A coupling field sent by a source component model can be associated with more than one target field and model (get). In that case, the source model needs to send (“put”) the field only once and the corresponding data will arrive at multiple targets as specified in the *namcouple* configuration file. Different coupling frequencies and transformations are allowed for different coupling exchanges of the same field. If coupling restart files are required (either if a LAG or if a LOCTRANS transformation is specified), it is mandatory to specify different files for the different fields.

The inverse feature is not allowed: a single target (get) field CANNOT be associated with multiple source (put) fields.

- The debug files

The debug mode was greatly improved as compared to OASIS3.3. The level of debug information written out to the OASIS3-MCT debug files for each model process is defined by the \$NLOGPRT value in the *namcouple*. All details are provided in section 3.2.

### B.6.5 Changes in the configuration file *namcouple*

- The *namcouple* configuration file of OASIS3-MCT is fully backward compatible with OASIS3.3. However, several *namcouple* keywords have been deprecated even if they are still allowed. These keywords are noted “UNUSED” in sections 3.2 and 3.3 and are not fully described. Information below these keywords will not be read and will not be used: \$SEQMODE, \$CHANNEL, \$JOB-NAME, \$INIDATE, \$MODINFO, \$SCALTYPE.
- Also the following inputs should not appear in the *namcouple* anymore as the related functionality are not supported anymore in OASIS3-MCT (see above): field status AUXILARY, time transforma-

tion ONCE, REDGLO, INVERT, MASK, EXTRAP, CORRECT, INTERP, MOZAIC, FILLING, SUBGRID, MASKP, REVERSE, GLORED.

- To get 2D fields in the debug output NetCDF files, the 2D dimensions of the grids must be provided in the *namcouple* (except if the field has the status OUTPUT); otherwise, the fields in the debug output files will be 1D.

### B.6.6 Other differences

- IGNORED and IGNOUT fields are converted to EXPORTED and EXPOUT respectively.
- The file `areas.nc` is not needed anymore to calculate some statistics with options CHECKIN and/or CHECKOUT.
- SEQ index is no longer needed to ensure correct coupling sequencing within the coupler. Use of SEQ allows the coupling layer to detect potential deadlocks before they happen and to exit gracefully (see section 2.5.3).
- The I/O library `mpp.io` is no longer used to write the restart and output files.

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