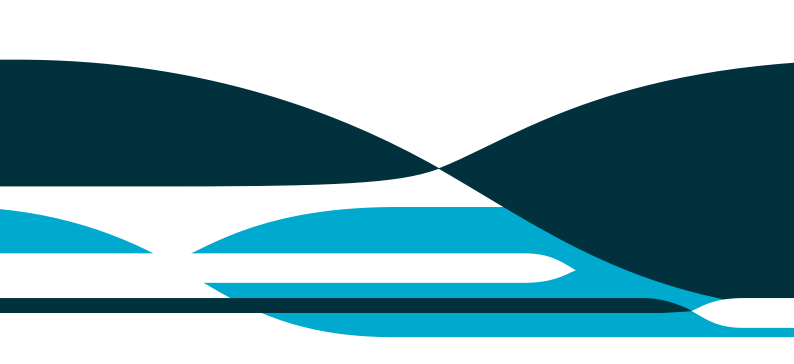


# User guide for the CSIRO Environmental Modelling Suite (EMS): Optical, biogeochemical, and carbon chemistry models.

Mark Baird, Mathieu Mongin, Karen Wild-Allen, Barbara Robson, Jennifer Skerratt, Farhan Rizwi, Monika Soja-Wozniak, Anna Maggiorano, Lachlan Phillips

> GitHub Version 1.3.1 Local SVN 7520

1st November 2024



## **Original citation**

Baird ME, Mongin M Wild-Allen KA Robson B Skerratt J. Rizwi F. 2018. User guide for the CSIRO Environmental Modelling Suite (EMS): Optical, biogeochemical, and carbon chemistry models. CSIRO Hobart.

## Copyright

© Commonwealth Scientific and Industrial Research Organisation 2018. To the extent permitted by law, all rights are reserved and no part of this publication covered by copyright may be reproduced or copied in any form or by any means except with the written permission of CSIRO.

## Further information, acknowledgments and code access

The code is available from: https://github.com/csiro-coasts/EMS. All conditions for the use of the code can be found on the GitHub archive, as well as acknowledgement of funding sources and authors of the CSIRO EMS code. Thank you

# **Contents**

1	Introdu	oduction1		
2	Code a	ode availability3		
3	Model configuration files			
	3.1	Master configuration file	4	
	3.2	The process list	7	
	3.3	The parameter file	0	
4	Tempe	Temperature dependence		
5	Sedime	Sediment – water column exchange rates		
6	Optical	model configuration	5	
	6.1	Original spectrally-resolved model	5	
	6.2	Column-based light calculations	8	
	6.3	Outputs of optical variables	6	
	6.4	Moonlight	7	
	6.5	Colour visualisation	8	
7	Carbon	Carbon chemistry model		
8	Ecologi	cological integration options		
9	Pre-pro	Pre-processing tools for BGC		
10	Runnin	g simulations3	3	
	10.1	Fully-coupled mode	3	
	10.2	Off-line transport mode	3	
11	Compu	tational diagnostics	5	
	11.1	Mass balance	5	
12	RECOM	1 / test case	7	

# **Figures**

Figure 1 The EMS model suite used for the eReefs project	2	
Figure 2 Schematic illustrating the vertical grid of the model and the locations on this grid the optical properties have been calculated.		
Tables		
Table 1 Standard process lists	0	
Table 1 Standard set of parameter values	2	
Table 2 Simulated observations	7	

#### Introduction 1

The CSIRO Environmental Modelling System (EMS) is a modelling tool that allows for the coupling of hydrodynamic / sediment / optical / biogeochemical (BGC) processes. It has been designed to represent shallow water systems, where processes at the sediment interface and below impact the water column, and vice versa, but also works well in deep waters.

One feature of EMS is the ability to include / exclude biogeochemical processes using the same model executable (or version), but different input configuration files. For example, the same executable can run a model of a tropical region with coral reefs, as well as a temperature region with multiple seagrass species or to just model the optical properties of the water column. The inclusion of model processes is therefore an important component of the model configuration, and makes up a significant portion of this User Guide.

This documentation is designed to be used only for user options that are not detailed in the much more extensive SHOC / COMPAS user guide (https://research.csiro.au/cem/software/ems/ems-documentation/). In particular, details regarding model running, model grids, advection and diffusion schemes, boundary conditions, as well as input / output formats are, are all documented in the SHOC / COMPAS user guide.

For developers considering adding new ecological processes, the original description of the ecological module (The "new" ecological model: user guide by Pavel Sakov) is available on the documentation page.

For more information on the model equations, please see the Biogeochemical Scientific Description for version B3p0.

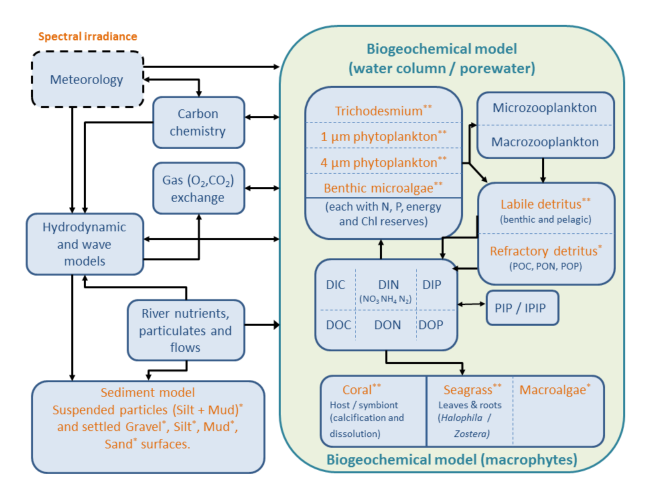


Figure 1 The EMS model suite used for the eReefs project.

## **Code availability** 2

The source code, and conditions of use, is available from a GitHub archive:

https://github.com/csiro-coasts/EMS.

Model documentation (including the latest version of this document) are available at:

https://research.csiro.au/cem/software/ems/ems-documentation/

# 3 Model configuration files

EMS is run from a compiled executable (see Section 10) that points to a number of configuration files, all of which are text files. The "Master" configuration is called in the execution line, and contains the names of any other files that are called.

For convenience, instead of using the filename suffix .txt, we use the following (noting they are still text files):

- .prm files are text files containing the model parameters.
- .tran files are text files used to run the off line transport model.
- .pss files are text files that list the location of the tracer point source.
- .ts files are text which contains time series of data of model input or outputs.
- .mnc files are text file listing source of multiple netcdf files.

Within these files any line starting with # is a comment line.

# 3.1 Master configuration file

This is the main file to run the model it contains:

- o model run time
- o model output and restart files
- hydrodynamic flags and parameters
- o grid & bathymetry definition
- model tracer list definition
- model forcing
- model time series outputs
- o model boundaries
- vector of bathymetry
- o x grid
- y grid

#### 3.1.1 Tracer list

The tracer list contains all the prognostic and diagnostic tracers that are available for output from the model. Most of the flags are self-explanatory, but the "diagn" is subtle.

A prognostic tracer has a tracer flag TRACER.diagn of 0, while a diagnostic tracer can take a value of 1 or 2. Diagnostic variables not altered by advection, diffusion or sediment processes.

In the ECOLOGY model, if diagn = 1 (flux diagnostic), then the diagnostic is the mean flux for the ecological time period (obtained by taking the change in the tracer from the initial value to the end and dividing by the timestep.). If diagn = 2 (mass diagnostic), then the diagnostic is the value of the tracer at the end of the time period.

In the rest of the code (hydro, sediments) diagn 1 and 2 simply the value at the output time, but are not advected of diffused. Therefore, tracers which are never given an increment by the ecology, such as TSS, the use of diagn = 1 by the sediment model is not a problem.

#### Example prognostic hydrodynamic tracer:

```
TRACERO.name
                               salt # SHOC variable name.
TRACERO.long name
                               Salinity # long name in netcdf
TRACERO.units
                               PSU
                             WATER SEDIMENT
TRACERO.type
TRACERO.fill_value 35.0 # default value initial value.

TRACERO.valid_range 0.0 40.0 # gives warning if outside range

# this tracer is advected
                              1
                                        # this tracer is diffused
TRACERO.diffuse
                              0
TRACERO.diagn
                                       # this tracer is not a diagnostic
tracer
                                             # if present, overwrite the
TRACERO.data
                        data/temp.nc
                         fill value, the data in temp.nc are interpolated onto
                         the model grid (see SHOC user guide for specification
                         of temp.nc)
```

#### Example prognostic dissolved BGC tracer:

```
TRACER23.name DIP
TRACER23.long name Dissolved Inorganic Phosphorus
TRACER23.units mg P m-3
TRACER23.valid range wc 0 1e+35
TRACER23.valid range sed 0 1e+35
TRACER23.diagn 0
TRACER23.type
                    WATER SEDIMENT
TRACER23.fill value wc 0.5
TRACER23.fill value sed 100 # fill value in the sediment, if present,
overwrite the value wc in the sediment
TRACER23.inwc 1
TRACER23.insed 1
TRACER23.dissol 1 # dissolved tracer
TRACER23.partic 0
TRACER23.advect 1
TRACER23.diffuse 1
TRACER23.decay 0
TRACER23.svel 0 # sinking rate
TRACER23.resusp rate 0
```

#### Example prognostic particulate BGC tracer:

```
TRACER33.name DetR_P
TRACER33.long_name Refractory Detrital Phosphorus
TRACER33.units mg P m-3
TRACER33.valid_range_wc 0 1e+35
TRACER33.valid_range_sed 0 1e+35
```

```
TRACER33.diagn 0

TRACER33.type WATER SEDIMENT

TRACER33.fill_value_wc 0.01

TRACER33.fill_value_sed 1.1e+03

TRACER33.inwc 1

TRACER33.insed 1

TRACER33.dissol 0

TRACER33.partic 1 # particulate tracer altered by sediment code

TRACER33.advect 1

TRACER33.diffuse 1

TRACER33.decay 0
```

# the following parameter are used in the sediment code to do sinking and resuspension

```
TRACER33.psize 1e-05 # size
TRACER33.b_dens 1e+09 # density
TRACER33.i_conc 2e+08
TRACER33.f_conc 2e+08
TRACER33.svel -2.31e-05
TRACER33.crit_stress 0.2
TRACER33.resusp_rate 0
```

#### TRACERO.TYPE #which type of variable:

```
WATER # 3D water column only
SEDIMENT # 3D sediment only
WATER SEDIMENT# 3D water column and SEDIMENT
WC2D# 2D water column
BENTHIC# 2D BENTHIC LAYER (sees both the last layer of the water column and top layer of the sediment)
```

An option available is the automatic generation of the ecological tracers in the tracer list based on the choice of processes. This development is the key to ecological implementations of RECOM, as the user only specifies the process list and parameter values, not the tracers. For this option, see standard process list (Section 3.2).

NOTE: A number of tracer name formats have special uses in EMS:

- TRACERNAME WC, TRACERNAME EPI, TRACERNAME SED should not be used.
- R\_XXX signifies reflectance at wavelength XXX. No tracer should have 'R\_' beginning unless it is specifying a reflectance.
- Ed\_XXX signifies downwelling irradiance at wavelength XXX. No tracer should have 'Ed\_' at the beginning unless it is specifying downwelling irradiance at a particular wavelength.

#### 3.1.2 Tracer statistics

To calculate a diagnostic tracer from other tracer output, or obtain a time-mean of the tracer of the, the tracerstat option can be used:

```
TRACER50.name Chl_a_sum

TRACER50.long_name Chlorophyll_sum

TRACER50.units mg Chla m-3

TRACER50.type WATER SEDIMENT

TRACER50.valid_range_wc 0 1e+35

TRACER50.valid_range_sed 0 1e+35

TRACER50.diagn 2

TRACER50.tracerstat sum(PhyL_Chl, PhyS_Chl, MPB_Chl, Tricho_Chl)

TRACER50.fill_value_wc 0

TRACER50.fill value sed 0
```

The time-means are especially useful when calculating a daily mean of a variable that changes cyclically through the day, such as chlorophyll concentration.

# 3.2 The process list

EMS contains many processes (i.e. phytoplankton growth, gas exchange etc.) that have been develop for projects around Australia and in Chile. To understand how these processes work the reader is encouraged to look at the top of the \*.c process file. This section discusses how to implement a sub-set of the EMS processes in a particular model configuration.

The process list file contains the listing of all the BGC processes that are to be run in the model, broken into the 3 vertical zones: water column, epibenthic and sediment. Each of these processes has 2 associated files (a .h and .c file), located in the source code folder at:

```
https://github.com/csiro-
coasts/EMS/tree/master/model/lib/ecology/process library
```

The process list use is invoked in the configuration file with the statement:

```
PROCESSFNAME process list.prm
```

All processes in the source codes (listed in allprocessess.c) are included in the model executable when it is compiled. But processes are only implemented in the model run if they are in the process list. This allows the same executable to be used for multiple applications. Some processes have arguments. In this case, the comments at the top of the \*.c file explain the options.

In general, processes with a  $_{wc}$ ,  $_{sed}$ , or  $_{epi}$  suffix can only be applied in the water column, sediment or epibenthic respectively.

The processes values\_common and values\_common\_epi are used define variables either in precalc, such that they can be used in the other processes, or in post-calc, so they can be outputted. Examples of variables defined include DIN.

Occasionally conflicts can occur with combinations of processes, and the user needs to be careful. For example, if a process is listed twice, the code will attempt to execute the rate process twice. Also, where two alternate versions of a process exists, such as trichodesmium\_grow\_wc and trichodesmium\_spectral\_grow\_wc, applying both will have unintended consequences.

Furthermore, a peculiarity of the code is that it checks for the mass balance of model through adding state variable to the sums TN, TP, TC and TO. It does these at the beginning and end of the process. It uses the fact that it is in a particular process to know to add a variable to this sum. So for example, if it is in the process trichodesmium spectral grow wc, it adds the nitrogen associated with the

trichodesmium structural material and the reserves of nitrogen to TN. However, if trichodesmium\_spectral\_grow\_wc is not specified, but there is trichodesmium in the tracer list, and zooplankton\_large\_spectral\_grow\_wc is called, then the mass balance will fail because TN does not include structural material of trichodesmium, and it is altered by grazing. Usually these conflicts are due to genuine incompatibility of processes. But occasionally it may be necessary to turn off the mass balance (see Section 11.1).

The configuration used in this example contains the following processes:

#### in the water column:

```
tfactor
             # Q10 temperature dependence of some parameters
viscosity
            # calculate the viscosity from T and S
          # calculate the molecular diffusivity from T and S
moldiff
remineralization # contain all the remineralization processes.
microphytobenthos spectral grow wc
phytoplankton spectral grow wc(small)
phytoplankton spectral grow wc(large)
trichodesmium mortality wc
trichodesmium grow wc
phytoplankton spectral mortality wc(small)
phytoplankton spectral mortality wc(large)
zooplankton mortality wc(small)
zooplankton mortality wc(large)
zooplankton large spectral grow wc
zooplankton small spectral grow wc
nitrification wc
p adsorption wc
carbon chemistry wc # OCMIP routines
gas exchange wc(oxygen,carbon) # if just one, (gas,dummy)
carbon chemistry wc
values common # calculate value across the different processes
massbalance wc
light spectral wc
age wc # perform age tracer experiment from a source
```

#### Notes:

- phytoplankton\_spectral\_grow\_wc, phytoplankton\_spectral\_mortality\_wc and zooplankton\_mortality\_wc contain an argument of either (small) or (large). For these three processes the code is identical, but the parameters vary between small and large.
- 2. gas\_exchange\_wc(oxygen,carbon) Gas exchange is calculated once, and applied to the gases in the argument. If carbon specified, carbon\_chemistry must be called above.
- 3. Trichodesmium processes can only be used with the spectrally-resolved processes.
- 4. Mass balance processes—this is a test within a cell and an ecological time step that the model conserves mass. It's most appropriate use is to ensure the code is consistent. It does not assess whether the model losses mass across boundaries or during advection or diffusion.
- in the epibenthic layer (~ benthic layer):

```
tfactor_epi
macroalgae_spectral_grow_epi
seagrass_spectral_grow_epi(Zostera)
coral_spectral_grow_epi
macroalgae_mortality_epi
seagrass_spectral_mortality_epi(Zostera)
values_common_epi
massbalance_epi
light_spectral_epi
diffusion_epi
```

#### Notes.

- 1. diffusion\_epi calculates a diffusive exchange between the sediment and water column based on a constant diffusion coefficient specified by parameters EpiDiffCoeff and EpiDiffDz. It is in additional to the diffusive flux specified in the sediment model by
- in the sediment layers:

```
tfactor
viscosity
moldiff
remineralization
light_sed
microphytobenthos_spectral_grow_sed
microphytobenthos_spectral_mortality_sed
phytoplankton_spectral_mortality_sed(small)
phytoplankton_spectral_mortality_sed(large)
trichodesmium_mortality_sed
nitrification_denitrification_sed
p_adsorption_sed
values_common
massbalance_sed
```

## • for the entire column:

From SHOC version 6773 onwards, a fourth process type became available, "column"

```
column {
    light_spectral_col()
}
```

This has been developed for the purposes of radiative transfer solution for the light field, but will no doubt be used in more applications in the future.

#### 3.2.1 Arguments to processes and multiple process calling.

Some of the processes contain arguments. These are to allow the user to pursue different options (such as calculating moonlight in <code>light\_spectral\_wc</code>) or to invoke a processes using a different set of state variables and parameter values (such as large and small phytoplankton). In the second case, processes can be called more than once to allow for both varieties. So, for <code>example</code>, <code>seagrass\_spectral\_grow\_epi</code> can be called with "Zostera" or "Halophila". The

equations are the same, but "Zostera" uses SG and SGROOT tracers and parameters starting with SG, while "Halophila" uses SGH and SGHROOT tracers and SGH parameters.

#### 3.2.2 Standard process lists

Instead of specifying a list of processes from a file as described above, a number of key words are reserved for standard process lists. For example,

PROCESSFNAME BCG2p0

invokes a standard list of processes. When a standard process list is used, the model also generates the tracers that is required by the processes in this list. The list it uses is also written into the execution folder with a filename processes \*.prm, where \* is the key word. A list of the most commonly used lists is given below (Table 1).

**Table 1 Standard process lists** 

Keyword	Description of process list	
BGC2p0	Biogeochemical model used in the eReefs implementation B2p0 with 4 microalgae, 2 zooplankton, 3 seagrass, 1 macroalgae and a simple coral host symbiont model.	
BGC3p1	Biogeochemical model used in the eReefs implementation B3p1 with 4 microalgae, 2 zooplankton, 3 seagrass, 1 macroalgae and the coral host symbiont model with coral bleaching.	
gas_only	Air-sea gas exchange of carbon and oxygen only.	
optics_only	The spectrally-resolved optical model calculations only.	
dead_only	All processes in BGC3p1, except those involving living processes, and also including anammox. This includes the spectrally-resolved optical model.	
porewater_age	Aging process in sediment porewaters only.	
TASSE1p0	Processes for eTas model.	

Most of the standard process lists are available through the RECOM interface.

#### 3.3 The parameter file

This file contains a list of the BGC model parameters and their metadata.

NPARAMETERS 141

PARAMETERO.name Tref
PARAMETERO.desc Reference temperature

PARAMETERO.units Deg C PARAMETERO.value 20.000000 PARAMETERO.stderr 0.000000

Set by parameter library options. PARAMETERO.ref

#### Notes:

- 1. EMS does read the "units", and if a time unit other than "seconds" is specified, it will convert to seconds for use in the code.
- 2. As a matter of principle, all biological parameters are read into the code during the initialisation phase.
- 3. The top of the bio\_spectral.prm file contains a line: "NPARAMETERS 98" which must be manually changed to the number of parameters (remember the count starts at 0).
- 4. Just because a variable appears in the bio.prm files does not imply it is used in the code. If it is never called by any processes, or a typo exists in the "name", the code will not use it. Often this results in a default value in the code being used. To ensure the value of the parameter that is implemented is known, **ecology\_setup.txt** prints out each time the code uses a parameter value. The self-explanatory statements, are one of the following three:

```
Ecol. parameter forced read, <NAME> = XXX
Ecol. parameter tried to read <NAME> and found XXX
Ecol. parameter tried to read <NAME>, but not in parameter file.
Code default of <NAME>, XXX
```

5. The variables that are read from the parameter file and are used in an ecological process are output to a netcdf file named ecology\_parameter\_setup.nc which has a format like:

```
netcdf ecology parameter setup {
         variables:
                  double Q10 ;
                  010:name = "010";
                  Q10:description = "Temperature coefficient for rate
parameters";
                  Q10:units = "none";
                  Q10:function call = "try parameter";
            // global attributes:
                 :title = "CSIRO Environmental Modelling Suite (EMS) biological
parameter setup.";
                  :description = "All variables specified in the biological
parameter file" ;
                  :date created = "Mon Apr 1 09:35:09 2024";
           data:
           Q10 = 2;
         }
```

Apart from providing a providence of parameter values, this file can be used to automate recalculation of model rates (e.g. umax = umax\_t0 \* Q10^((T-Tref)/10), where Tref, Q10 and umax\_t0 are read (in matlab, python etc.) from the <code>ecology\_parameter\_setup.nc</code> and T comes from an output file). The attribute function call informs whether the parameter needed to be specified in the bio.prm file ("try\_parameter"), or whether it is optional ("get\_parameter") in the first instance that it is read.

#### 3.3.1 Standard parameter file

Instead of specifying parameter values from a file as described above, a number of key words are reserved for a set of standard parameter values. For example,

```
BIOFNAME BCG2p0
```

invokes a standard set of parameter values used in the B2p0 configuration. The list used is also written into the execution folder with a filename bio \*.prm, where \* is the key word. A list of the most commonly used lists is given below (Table 1).

**Table 2 Standard set of parameter values** 

Keyword	Description of set of parameter values
BGC2p0	Biogeochemical model used in the eReefs implementation B2p0 with 4 microalgae, 2 zooplankton, 3 seagrass, 1 macroalgae and a simple coral host symbiont model.
BGC3p1	Biogeochemical model used in the eReefs implementation B3p1 with 4 microalgae, 2 zooplankton, 3 seagrass, 1 macroalgae and the coral host symbiont model with coral bleaching.
porewater	Just parameter values for porewater aging.
TASSE1p0	Biogeochemical model used in the eTas.

# 4 Temperature dependence

The functional form of the temperature-dependence of the vast majority of ecological rates is set by choosing a process, either tfactor or tfactor\_clim. In both cases a power law relationship is applied, of the form  $k_T = k_{ref} \, Q10^{(T-T_{ref})/10)$ , where  $k_T$  is the rate at temperature T,  $k_{ref}$  is the rate at  $T_{ref}$ , and Q10 is a unitless coefficient, often 2.0 to represent a doubling in the rate over 10 degrees.

If tfactor is called, Tref is calculated based Tref from the biological parameter file.

If tfactor clim is called, T<sub>ref</sub> is spatially- and temporally changing, and is specified by:

```
TRACER142.name
                          temp clim
TRACER142.long name
                          Climatological temperature
TRACER142.units
                          deg C
TRACER142.valid range
                        0.00e+00 1.00e+35
TRACER142.fill value
                         26.0
TRACER142.advect
                          0
TRACER142.diffuse
TRACER142.diagn
TRACER142.partic
                          0
TRACER142.dissol
                          0
TRACER142.type
                         WC3D SEDIMENT ECOLOGY DIAGNOSTIC
TRACER142.eco_flag
                        NORESET
STRACER142.reset file forcing bgc/TS CARSa.nc4(temp clim=temp CARS)
TRACER142.reset dt 1 day
```

where the value of Tref is updated from file reset dt every days.

#### **Sediment – water column exchange rates** 5

The exchange of dissolved tracers between the water column and sediment occurs because of:

- Deposition of sediments, entraining the porewater from the bottom layer of the water column (a flux into the sediment).
- 2. Erosion of sediments, releasing porewater into the water column (a flux out of the sediment).
- 3. Diffusion in the sediment model (which is slow).
- 4. Diffusion in the ecology model (implemented in the process diffusion epi).

The sum of processes 1-4 is difficult to calculate from the state variables in the model output, so inline diagnostic calculations are available.

The sediment model calculates the sum of the instantaneous rates of 1-3 of variable XXX if the tracer XXX fluxsedi inst is specified:

```
TRACER196.name DIC_fluxsedi_inst
TRACER196.long_name Sed-water sed. DIC
TRACER196.units mg C m-2 s-1
TRACER196.fill_value 0.00e+00
TRACER196.valid_range -1.00e+35 1.00e+35
                                          Sed-water sed. DIC flux
TRACER196.diagn
TRACER196.advect
TRACER196.diffuse
TRACER196.type
                                              WC2D DIAGNOSTIC
```

The ecological model calculates diffusion in the ecological model if the process diffusion epi is evoked, and the diagnostic tracer XXX sedflux is specified.

```
TRACER195.name DIC_sedflux
TRACER195.long_name Sed-water ecol. DIC flux
mg C m-2 s-1
TRACER195.tall_value 0.00e+00
TRACER195.valid_range 0.00e+00 1.00e+35
TRACER195.diagn
                                   WC2D ECOLOGY DIAGNOSTIC
TRACER195.type
```

At present the only variables available for XXX sedflux are DIC, DIP, NO3, NH4, alk, and Oxygen. Unless a large resuspension or deposition event has occurred, XXX sedflux will generally be an order of magnitude more than XXX fluxsedi inst. The total instantaneous sediment-water exchange (positive in the sediment to water direction) is the sum of the XXX sedflux and XXX fluxsedi inst.

# 6 Optical model configuration

# 6.1 Original spectrally-resolved model

From 2010-2021 (up until B3p3), the spectrally-resolved optical was implemented using the processes light\_spectral\_wc, light\_spectral\_uq\_epi and light\_spectral\_sed. This is being replaced with a column-based calculation (light\_spectral\_col) described in Sec. 6.2.

The spectrally-resolved optical model calculates the inherent optical properties (IOPs) from the optically-active state variables, and then uses the IOPs to calculate the apparent optical properties (AOPs) such as the light field and remote-sensing reflectance.

The optical model is configured using parameter variables specified in **bio.prm**, through option calls to light processes in the process list models, and through tracer list specification. The relationships used to relate state variables to IOPs (such as the pigment-specific absorption coefficient of each of the pigments) are specified in the file bio opt.c:

https://github.com/csiro-coasts/EMS/blob/master/model/lib/ecology/bio opt.c.

The wavelengths (centre and bandwidths) used are specified in the parameter file using:

```
PARAMETER4.name Light_lambda
PARAMETER4.desc Wavelengths of light
PARAMETER4.units nm
PARAMETER4.value 290.000000 310.000000 330.000000 350.000000 370.000000
390.000000 410.000000 430.000000 440.000000 450.000000 470.000000 490.000000
510.000000 530.000000 550.000000 570.000000 590.000000 610.000000 630.000000
650.000000 670.000000 690.000000 710.000000 800.000000
PARAMETER4.stderr 0.000000
PARAMETER4.ref Approx. 20 nm resolution with 10 nm about 440 nm. PAR (400-700) is integral of bands 2-22.
```

The 'hard' coded shortest wavelength edge is 140 nm, and the longest edge at 8000 nm. The spectral 'grid' that results from this specification is output into **ecology\_setup.txt**:

```
Centre of waveband (edge):
290.00 (140.00 - 300.00)
310.00 (300.00 - 320.00)
330.00 (320.00 - 340.00)
350.00 (340.00 - 360.00)
370.00 (360.00 - 380.00)
390.00 (380.00 - 400.00)
410.00 (400.00 - 420.00)
430.00 (420.00 - 435.00)
440.00 (435.00 - 445.00)
450.00 (445.00 - 460.00)
470.00 (460.00 - 480.00)
490.00 (480.00 - 500.00)
510.00 (500.00 - 520.00)
530.00 (520.00 - 540.00)
550.00 (540.00 - 560.00)
570.00 (560.00 - 580.00)
590.00 (580.00 - 600.00)
```

```
610.00 (600.00 - 620.00)
630.00 (620.00 - 640.00)
650.00 (640.00 - 660.00)
670.00 (660.00 - 680.00)
690.00 (680.00 - 700.00)
710.00 (700.00 - 755.00)
800.00 (755.00 - 8000.00)
```

For this particular optical grid, and the clear sky spectrally-resolved solar radiation, approximately 40 % of the energy of the sun is between 400 and 700 nm. Note that in this configuration we have used a 20 nm resolution, but added an extra wavelength at 440 nm to better resolve absorption by photosynthetic pigments.

In the tracer list, the inclusion of tracers with tracer names of  $R_*^{***}$  results in the optical model calculating remote-sensing reflectance,  $R_{rs}$ , at wavelength \*\*\* nm:

```
TRACER93.name R_640

TRACER93.long_name Remote-sensing reflectance @ 640 nm

TRACER93.units sr-1

TRACER93.fill_value 0.00e+00

TRACER93.valid_range 0.00e+00 1.00e+35

TRACER93.diagn 2

TRACER93.type WC2D ECOLOGY DIAGNOSTIC
```

Warning: Do not include a tracer with R\_ as its first two characters unless the next characters are specifying a wavelength.

In the tracer list, the inclusion of tracers with tracer names of Ed $_****$  results in the optical model calculating downwelling irradiance,  $E_d$ , at wavelength \*\*\* nm:

Warning: Do not include a tracer with Ed\_ as its first three characters unless the next characters are specifying a wavelength.

In the case that the wavelength specified in R\_\*\*\* or Ed\_\*\*\* is not one of the centre wavelengths the output will be a linear interpolation of centre wavelength.

Further, a number of simulated satellite products can be outputted if the required remote-sensing reflectance at the required wavelengths, and the key tracer name are specified (with similar options to the Remote-sensing reflectance above) (Table 3). If using the light\_spectral\_epi process, the code requires remote-sensing wavelengths (e.g. OC3M would require R\_443, R\_488 and R\_547). If using the light\_spectral\_col process that uses a spectral response curve, the code requires bands (e.g. OC3M\_SR would require MODIS\_B2, MODIS\_B4, MODIS\_B6].

Table 3 Simulated observations. [variables in brackets refer to spectral response curves used in light\_spectral\_col].

Tracer name	Description of simulated variable	wavelength (nm)	Source
OC3M [OC3M_SR]	Three-band chlorophyll algorithm calibrated for the MODIS sensor.	443, 488, 547; [MODIS_B2, MODIS_B4, MODIS_B6]	
OC3V [OC3V_SR]	Three-band chlorophyll algorithm calibrated for the VIIRS sensor.	443, 486, 551; [VIIRS_B2, VIIRS_B3, VIIRS_B4]	
OC4Me [OC4Me_SR]	Four-band chlorophyll algorithm calibrated for the MERIS and Sentinel3A sensors	443, 488, 510, 560; [Sentinel_B3, Sentinel_B4, Sentinel_B5, Sentinel_B6]	
Secchi	Secchi depth calculated from vertical integral of attenuation of light at 488 nm.	488	Lee et. al (2016).
TSSM [TSSM_SR]	Turbidity based on red reflectance – local GBR algorithm.	645 [MODIS_B9]	Petus
KD490M [KD490M_SR]	Two band vertical attenuation at 490 nm.	488, 547; [MODIS_B4, MODIS_B6]	
nFLH	Normalized Fluorescence Line Height based on MODIS wavelengths.	667,678, 748 [MODIS_B9, MODIS_B10, MODIS_B11]	
Turbidity	Turbidity calculated from total scattering at 590 nm.	590	
Fluorescence	Chlorophyll concentration 'scaled' by processes that reduce fluorescence such as self-shading and photochemical quenching.	Integral of photosynthetic wavelengths.	

Finally, the default calculation of the zenith angle is on the time zone (in below example +10) specified at the top of the master configuration file:

TIMEUNIT seconds since 1990-01-01 00:00:00 +10

#### However, if the option:

```
PROJECTION geographic
```

is invoked, the zenith angle is calculated based on the longitude (as well as the hour, day, and latitude).

# 6.2 Column-based light calculations

A major upgrade to the optical model occurred through 2021/22, with improved configuration and calculation of the light field.

# 6.2.1 Tracer attribute specification of spectrally-resolved mass-specific optical parameters.

In light\_spectral\_col, the bulk of the spectrally-resolved optical parameters (e.g. mass-specific absorption by inorganic particles) are provided to model in a netcdf file. In order to assign a mass-specific optical property to a tracer, the file name and variable name are given in the tracer list (here for a particulate substance).

```
TRACER4.name Mud-mineral

TRACER4.units kg m-3

TRACER4.optical_file csiro_siop_library.nc

TRACER4.type WC3D SED SEDIMENT PROGNOSTIC OPTICAL

TRACER4.absorp_name ap_terr

TRACER4.scatter_name btp_terr

TRACER4.backscat_name B_terr

TRACER4.benreflt_name rho_mud
```

On initialisation, the process <code>light\_spectral\_col</code> finds all tracers that are optically-active (as specified by type <code>OPTICAL</code>). It then reads their mass-specific optical properties from the file specified. The values are then linearly interpolated onto the optical grid. Not all the attributes for each tracer must come from the same file.

During the integration the component of absorption due to the tracer X is calculated from ap\_X\*X. Similarly scattering, the backscattering ratio and benthic reflectance can be specified.

For benthic variables, the optical properties are the absorbance, reflectance and transmission of the surface:

```
TRACER59.name MA_N

TRACER59.long_name Macroalgae N

TRACER59.units g N m-2

TRACER59.fill_value 0.2

TRACER59.valid_range 0.00e+00 1.00e+35

TRACER59.diagn 0
```

```
TRACER59.type WC2D ECOLOGY PROGNOSTIC OPTICAL
TRACER59.optical_file project_siop_library.nc
TRACER59.abtance_name Ahalophila
TRACER59.refltce_name rho_macroalgae
TRACER59.trnmiss name Thalophila
```

For suspended macroalgae (implemented in macroalgae\_spectral\_grow\_wc), which use the 2D benthic optical properties for a 3D tracer, the two options available are:

```
TRACER59.abtance3d_name Ahalophila
TRACER59.refltce3d name rho macroalgae
```

At initialisation the model produces a file <code>optical\_setup.nc</code> that contains the spectrally-resolved optical parameters and the optical grid (wave band specification etc.), as well as useful optical attributes of the tracer. An example is absorption of FineSed:

```
double ap_FineSed(wave_centre) ;
    ap_FineSed:description = "Mass-specific absorption by non-CaCO3 particles" ;
    ap_FineSed:codename = "ap_FineSed" ;
    ap_FineSed:sourcefile = "project_siop_library.nc" ;
    ap_FineSed:sourcename = "ap_terr" ;
    ap_FineSed:symbol = "a_{non-CaCO3}" ;
    ap_FineSed:units = "m-1 (mg m^{-3})-1" ;
    ap_FineSed:interpolation_percent_error_in_data_range = 7.86551155453676 ;
    ap_FineSed:source_data_range_in_nm = 401.6, 750.3 ;
    ap_FineSed:model_range_used_for_interpolation_error_calculation = 410., 690. ;
    ap_FineSed:data_range_used_for_interpolation_error_calculation = 410., 686.3 ;
    ap_FineSed:rgb_colour_space = 164.85628351858, 151.793433426242, 116.99795659274 ;
    ap FineSed:mass concentration used for rgb colour space in clear water = 1. ;
```

Attributes include the percent area generated by interpolating onto the optical grid and the MODIS true colour produced (given as RGB) of suspending the particles in clear water.

## 6.2.2 Tracer attribute specification of spectral response curves.

Tracer attributes can also be used to specify the spectral response of a sensor. For the tracer attribute of specresp2d, the response is interpreted as a sensor to remote-sensing reflectance and is ideally used for an airborne or satellite based sensor with measurements of both downwelling irradiance and upwelling radiance (i.e.  $R_{rs} = E_d/L_u$ ). For example:

```
TRACER218.name Himawari8_Band1_SR
```

TRACER218.long\_name Col:Himawari8 Band 1

TRACER218.units 
TRACER218.fill\_value 0.00e+00

TRACER218.valid\_range -1.00e+35 1.00e+35

TRACER218.diagn 2

TRACER218.type WC2D ECOLOGY DIAGNOSTIC OPTICAL

TRACER218.optical\_file csiro\_siop\_library.nc

The tracer name does not need to be the same as the specresp2d name.

TRACER218.specresp2d name Himawari8 Band1 SR

Alternatively, the tracer attribute of specresp3d is used for a downwelling radiometer:

TRACER224.name Seabird SR 490 TRACER224.long\_name Col: Downwelling light at 490 nm TRACER224.units W m-2 nm-1TRACER224.fill value 0.00e+00 TRACER224.valid range 0.00e+00 1.00e+35 TRACER224.fill value sed 0.00e+00 TRACER224.valid range sed 0.00e+00 1.00e+35 TRACER224.advect 0 TRACER224.diffuse TRACER224.diagn 2 TRACER224.partic TRACER224.dissol 0 WC3D SED ECOLOGY DIAGNOSTIC OPTICAL TRACER224.type 0.00e+00 TRACER224.psize TRACER224.b\_dens 0.00e+00 TRACER224.i\_conc 0.00e+00 TRACER224.svel 0.0 TRACER224.cohesive 1 TRACER224.calcvol TRACER224.floc TRACER224.resuspend 1 TRACER224.deposit TRACER224.eco flag TRACER224.optical file seabird.nc TRACER224.specresp3d name Seabird SR 490

In this case the spectral response curve is multiplied by the downwelling irradiance, and then normalised by the area under the spectral response curve, in order to give a measure of the downwelling irradiance that would be measured by that sensor by nm.

Spectral response curves are often quite narrow (say 10 nm for Sentinel ocean colour bands), so using the optical model grid for the response curves is unreliable. As a result, the remote-sensing reflectance is interpolated onto a 1 nm grid that is then used against the spectral response curve.

The file optical\_file can be a netcdf file of any name. Five libraries that can be requested from CSIRO are:

- Spectral response curves csiro spectral response library.nc
- General optical properties csiro optical properties library.nc
- Benthic plant reflectance csiro benthic plant reflectance library.nc
- Mass-specific inherent optical properties csiro mass specific iops library.nc
- Generic coloured particles and tracers csiro coloured dyes particles library.nc

A sixth library, csiro\_iops\_standards\_library.nc, contain spectrally-constant values of 0, 0.001, 0.01, 0.1, 1, 10, 100 and 1000 that are useful for testing the code and model configuration.

# 6.2.3 Column-based light calculations.

Light\_spectral\_col is a column-based process meaning it has access to state variables in the entire water column. This allows the calculation to be solved via the ray tracing technique used in the earlier optical model, or alternatively using radiative transfer models. The code provides two schemes do the conversion from IOPs to AOPs, a geometric optics scheme (default and only available until August 2024), and a three directional radiative transfer model (RT3) based on Gregg and Carder (1990).

To undertake the RT3 calculations, a diagnostic tracer for the PAR-integrated direct downwelling light, PAR\_d\_3RT, must be specified. The remaining outputs of light field can then be specified such as the diffuse downwelling (PAR\_s\_3RT), scalar light (PAR\_sc\_3RT), and diffuse upwelling (PAR u 3RT).

The calculated light field is used to drive photosynthesis and to calculate remote-sensing reflectance. To choose which light model (geometric optics or directional radiative transfer model), an argument is given in the process list.

Process specification: light spectral col(G|R)

The three directional radiative transfer model (RT3) requires a diffuse and direct component of the surface radiative forcing. By default this is set to be 50% direct and 50% diffuse. Alternatively, if the 2D tracers SWR\_diffuse and SWR\_direct appear in the tracer list containing the diffuse and direct Short Wave Radiation (< 6000 nm) above the sea surface, then these will be used in the RT3 calculation. Further, with SWR\_diffuse and SWR\_direct specified in the tracer list, the surface albedo will be calculated separately for the direct and diffuse streams using the OASIM model formulation.

### 6.2.4 Radiative transfer calculation (HydroLight library).

HydroLight is a radiative transfer numerical model computes radiance distributions and related quantities (irradiances, reflectances, diffuse attenuation functions, etc.) in any water body. After Curtis Mobley, the developer of HydroLight, retired from Sequoia in December 2016, HydroLight became a product of Numerical Optics, Ltd. and is managed by Dr. John Hedley.

EMS contains an interface to hydrolight that allows calling to their routines. The HydroLight Code itself is not part of the EMS distribution, and must be obtained from Numerical Optics, Ltd. For users without an EMS licence, light spectral col needs to comment out the following line:

```
// #define INCLUDE HYDROLIGHT 1 // Compile with Hydrolight calls.
```

Light\_spectral\_col is archived in SVN and GitHub with this line commented out, so no action is required by users not intending to use the HydroLight interface.

## 6.2.5 Generation of a Matlab / python script to plot optical setup.

During model initialisation, EMS generates a Matlab script (optical\_setup\_matlab\_check.m) for plotting all the optical properties that have been interpolated onto the optical grid. For each optically property of each optically-active variable, a segment of code is produced:

```
Figure; set (gca, 'FontSize', 14);
var = ncread(file, 'ap_FineSed');
sourcefile = 'project_siop_library.nc';
var1 = ncread(sourcefile, 'ap_terr');
rgb = ncreadatt(file, 'ap_FineSed', 'rgb_colour_space');
pp = plot(wave, var, '+k'); set (pp, 'LineWidth', 2); hold on;
pp = plot(var1(:,1), var1(:,2), '-k'); set (pp, 'LineWidth', 4.4); hold on;
pp = plot(var1(:,1), var1(:,2), '-k'); set (pp, 'LineWidth', 4.4); hold on;
xlabel('wavelength [nm]');
ylabel('ap_FineSed [m-1 (mg m^{-3})-1]', 'Interpreter', 'none');
title('Interpolation error: 7.865512e+00');
set(gca, 'xlim', [200 1000]);
```

The Matlab script creates a text file (references.txt) that lists all the spectrally-resolved optical properties and the original source of the data, which must be an attribute in the source netcdf file listed in the tracer list.

EMS also generates a similar python script (optical setup matlab check.py).

# 6.2.6 Output of optical properties in a column

The optical properties of single columns are outputted into a separate netcdf files named optical\_column\_out\_0.nc at the ECOLOGY\_DT timestep. The columns used are specified in the biological parameter file, using the format:

```
PARAMETER144.name opt_output_i_index

PARAMETER144.desc Output of optical column at i_index

PARAMETER144.value 266 266 266 266 266

PARAMETER145.name opt_output_j_index

PARAMETER145.desc Output of optical column at j_index

PARAMETER145.value 23 24 25 26 27 28
```

The first column also becomes the column on which HYDROLIGHT calculations are undertaken. The variables are plotted in a so for reference the variable:

```
double depth(record, k_grid) ;
    depth:description = "metres from sea surface of k_grid" ;
```

So the depth on the EMS grid can be calculated using the surface elevation.

## 6.2.7 Other aspects of light\_spectral\_col

Points to note:

- When using light\_spectral\_col a water column, epibenthic and sediment process must be implemented or otherwise the variable list for the column is not generate properly.
- As the light field is calculated at the output time (or postcalc), the rate of light absorption and the incremental benefit of adding chlorophyll must be stored as tracers
  - o For each microalgae added, two new tracers:

```
TRACER195.name PhyL_kI

TRACER195.long_name Col: Large Phytoplankton absorption

TRACER195.units mol cell-1

TRACER195.fill_value 0.00e+00

TRACER195.valid_range 0.00e+00 1.00e+35

TRACER195.fill_value_sed 0.00e+00

TRACER195.valid_range_sed 0.00e+00 1.00e+35

TRACER195.advect 0

TRACER195.diffuse 0

TRACER195.diagn 0
```

```
TRACER195.partic
TRACER195.dissol
                     WC3D SED ECOLOGY DIAGNOSTIC
TRACER195.type
TRACER195.psize
                     0.00e+00
TRACER195.b dens
                      0.00e+00
TRACER195.i conc 0.00e+00
TRACER195.css erosion
                           -1.00e+00
TRACER195.css_deposition -1.00e+00
TRACER195.svel
                     0.000000
TRACER195.cohesive
                      1
TRACER195.calcvol
TRACER195.floc
TRACER195.resuspend
TRACER195.deposit
TRACER195.adsorb
TRACER195.adsorbkd 0.00e+00
TRACER195.adsorbrate
                      0.00e+00
TRACER195.eco_flag
                     NORESET
TRACER196.name
              PhyL_yCfac
TRACER196.long name
                    Col: Large Phytoplankton opaqueness
TRACER196.units
TRACER196.fill_value 0.00e+00
TRACER196.valid range
                      0.00e+00 1.00e+35
TRACER196.fill value sed 0.00e+00
TRACER196.valid range_sed 0.00e+00 1.00e+35
TRACER196.advect
TRACER196.diffuse
                       \cap
TRACER196.diagn
TRACER196.partic
TRACER196.dissol
                WC3D SED ECOLOGY DIAGNOSTIC
TRACER196.type
TRACER196.psize
                      0.00e+00
TRACER196.b dens
                      0.00e+00
TRACER196.i_conc 0.00e+00
TRACER196.css erosion
                          -1.00e+00
TRACER196.css_deposition
                            -1.00e+00
TRACER196.svel
                      0.0
TRACER196.cohesive
                      1
TRACER196.calcvol
```

TRACER196.floc 0
TRACER196.resuspend 1
TRACER196.deposit 1
TRACER196.adsorb 0

TRACER196.adsorbkd 0.00e+00
TRACER196.adsorbrate 0.00e+00
TRACER196.eco\_flag NORESET

#### o For each benthic plant added, one new tracer:

TRACER201.name SGH\_kI

TRACER201.long\_name Col: SGH absorption

TRACER201.units mol m-2
TRACER201.fill\_value 0.00e+00

TRACER201.valid\_range 0.00e+00 1.00e+35

TRACER201.diagn 0

TRACER201.type WC2D ECOLOGY PROGNOSTIC

# o For suspended macroalgae, one new tracer:

TRACER211.name SMA kI

TRACER211.long\_name Floating MA absorption

TRACER211.units mol m-2
TRACER211.fill\_value 0.00e+00

TRACER211.valid\_range 0.00e+00 1.00e+35

TRACER211.fill value sed 0.00e+00

TRACER211.valid range sed 0.00e+00 1.00e+35

TRACER211.advect 0

TRACER211.diffuse 0

TRACER211.diagn 0

TRACER211.partic 0

TRACER211.dissol 0

TRACER211.type WC3D SED ECOLOGY DIAGNOSTIC

TRACER211.psize 0.00e+00
TRACER211.b\_dens 0.00e+00
TRACER211.i\_conc 0.00e+00

TRACER211.css\_erosion -1.00e+00

TRACER211.css\_deposition -1.00e+00

TRACER211.svel 0.000000

TRACER211.cohesive 1
TRACER211.calcvol 0

```
TRACER211.floc
TRACER211.resuspend
                          1
TRACER211.deposit
TRACER211.adsorb
                          0
TRACER211.adsorbkd
                          0.00e+00
TRACER211.adsorbrate
                          0.00e+00
TRACER211.eco flag
                          NORESET
```

#### The most important improvements are:

- Specification of reflectance, absorptance (photosynthetic and non-photosynthetic) and transmission for benthic plants.
- CDOM absorption only uses CDOM tracers.
- Reflectance calculated for satellite sensor bands.
- Spectrally-resolved bioluminescence emission spectra.
- Wavelength dependence of air-sea water interface refraction.
- The fraction of light into the reddist band is the remainder of the light not allocated into bluer bands.

#### 6.3 Outputs of optical variables

The output of optical variables is complicated by the calculation of downwelling irradiance (PAR z) at the cell faces, but PAR at the cell centres. Further, cell centres of the partially-filled surface and bottom layers deviate from the nominal cell centre (z centre in output files). The schematic below (Figure 2) illustrates these complications and gives equations that can be calculated from the output. None of these complications affect the output of 2D variables (such as remote-sensing reflectance) which are calculated on the moving free surface at the output time.

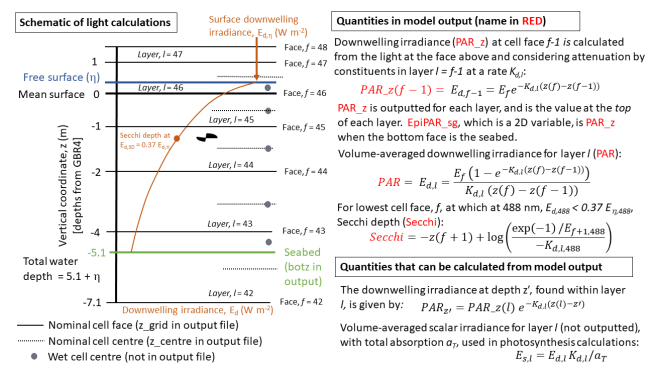


Figure 2 Schematic illustrating the vertical grid of the model and the locations on this grid that the optical properties have been calculated.

# 6.4 Moonlight

The optical model includes the calculation of moonlight parameters using orbital calculations in the hydrodynamic model, and the USGS ROLO spectral irradiance model (https://astrogeology.usgs.gov/moon-cal/modeling/irradiance.php, Kieffer, H. H and T. C. Stone (2005) The spectral irradiance of the moon. The Ast. J. 129: 2887-2901).

In order to invoke moonlight calculations with <code>light\_spectral\_wc</code>, the third argument to the process must be 'M'. For <code>light\_spectral\_col</code>, moonlight is always calculated.

In order for moonlight calculations to be undertaken, PROJECTION must be geographic.

The output variables for moonlight are:

Moonlight	Moonlight (downwelling PAR-integrated irradiance)
Lunar_zenith	Moon zenith (0 overhead; $\pi/2$ at horizon)
Lunar_phase	Moon phase (0 new moon; $\pi/4$ first quarter; $\pi/2$ full moon;
	3π/4 last quarter; 0 new moon)
Moon_fulldisk	Moonlight brightness (PAR-integrated) if looked at directly.
Lunar_azimuth	Moon azimuth angle, following a north clockwise-wise convention (0 north; $\pi/4$ west; $\pi/2$ south; $3\pi/4$ east)

The full disk reflectance (i.e. looking straight at the moon without including the reduction to the zenith angle,  $1/\cos\theta$ ) includes the effect of moon phase on irradiance.

The USGS ROLO model include the effect of Earth-Sun distance, Earth Moon distance, spatially varying albedo of the moon (i.e. the craters are darker than the plains), the point of the Moon facing the Earth, phase of the moon, time of day, point on the Earth (latitude).

The Moonlight calculations in light spectral col include the same atmospheric attenuation calculations as used for Sunlight in the EMS hydrodynamic model (although using the lunar instead of solar zenith). For these to be undertaken the .tran file must include inputs of CLOUD, PRESSURE and HUMIDITY:

```
CLOUD /home/eReefs/forcing/access/gbr 2016 03 access.nc
CLOUD INPUT DT 1 hour
PRESSURE /home/eReefs/forcing/access/gbr 2016 03 access.nc(pressure=mslp)
PRESSURE INPUT DT 1 hour
HUMIDITY /home/eReefs/forcing/access/gbr 2016 03 access.nc(hr=hr2m)
HUMIDITY INPUT DT 1 hour
```

In addition to Moonlight, a small starlight term is given (~1000 x less than the full moon overhead) that allows calculations to be undertaken with a non-zero light field. For simplicity, the galactic centre of the Milky Way is assumed to be at a zero zenith.

#### 6.5 Colour visualisation.

For production of true colour a red, green and blue wavelength of remote-sensing reflectance needs to be specified. In the CSIRO visualisation software DIVE, MODIS true colour uses tracers R\_470, R\_555 and R\_667, and MODIS SR true colour uses tracers MODIS\_B9, MODIS\_B6 and MODIS\_B3. If these tracers aren't available in the output files then no option to visual true colour is given by DIVE.

# 7 Carbon chemistry model

To implement the carbon chemistry model requires that the tracers temperature (temp), salinity (salt), total alkalinity (alk), dissolved inorganic carbon (DIC) and pH (PH) be specified. Additional diagnostic variables that can be produced by the carbon chemistry model are calcite saturation, aragonite saturation, surface partial pressure of carbon dioxide, bicarbonate ion concentration, carbonate ion concentration and atmospheric  $CO_2$  concentration. The atmospheric  $CO_2$  concentration can either be set to a constant using the parameter file:

```
PARAMETER3.name xco2_in_air
PARAMETER3.desc Atmospheric CO2
PARAMETER3.units ppmv
PARAMETER3.value 396.480000
PARAMETER3.stderr 0.000000
PARAMETER3.ref Mean '13, Mauna Loa: http://co2now.org/current-co2/co2-now/
```

#### Or by using a resetting spatially-resolved tracer:

```
TRACER77.name
                          xco2_in_air
TRACER77.long name
                         Atmospheric CO2
TRACER77.units
                          ppmv
#TRACER77.fill_value 400.0
TRACER77.valid_range 0.00e+00 1.00e+35
TRACER77.advect
                          0
TRACER77.diffuse
                          0
TRACER77.diagn
                          \cap
TRACER77.partic
TRACER77.dissol
                           0
                                   SEDIMENT ECOLOGY PROGNOSTIC
TRACER77.type
                           WC3D
TRACER77.eco flag
TRACER77.reset file forcing bgc/global co2 merged 2016.ts
TRACER77.reset dt 30 days
```

The Ocean-Carbon Cycle Model Intercomparison Project (OCMIP) numerical methods are used to quantify air-sea carbon fluxes and the carbon dioxide system equilibria in seawater. The OCMIP procedures quantify the state of the  $CO_2$  system using two prognostic variables, the concentration of dissolved inorganic carbon,  $C_T$ , and total alkalinity,  $A_T$ . The value of these prognostic variables, along with salinity and temperature, are used to calculate the partial pressure of carbon dioxide,  $pCO_2$ , in the surface waters using a set of governing chemical equations that are solved using a Newton-Raphson method with literature-derived constants. One alteration from the typical implementation of the OCMIP algorithm is that we increased the search space for the iterative scheme from  $\pm 0.5$  pH units (appropriate for global models) to  $\pm 2.5$ . The air-sea flux of carbon dioxide is calculated based on wind speed data (requiring process gas\_exchange\_wc(carbon)), and the air-sea gradient in  $pCO_2$ , using an empirical net flux relationship.

# **8** Ecological integration options

In the model configuration file .prm or .tran file: ECOLOGY options are:

```
DO_ECOLOGY 1  ## DO ECOLOGY FLAG SET TO 1 TO run ecology ECOLOGY DT 1200  # ECOLOGY time step
```

Note: make sure that the ecology timestep is the same length or less than the shortest timestep in the physics or forcing data. If this is not true, the model will change <code>ECOLOGY\_DT</code> from what is specified in the file, which will be noted in the **runlog**.

#### Specify the other configuration files:

```
biofname in/bio_spectral.prm # file that contain model parameters processfname in/processes_spectral.prm # file that contains the list of which processes.
```

#### **Specify integration checks:**

```
check_nans 0 \,\, \, \, \, test to check is there is any NaN in the state variable array , computationally expensive so only used during debugging.
```

check\_negs 0 # test to check is there is any <0 in the model , expensive so only used during debugging.

Both check\_nans and check\_negs cause the model to stop, and output diagnostic information to the runlog. check\_negs is will often fail with dissolved oxygen, that is allowed to go slightly negative.

#### Miscellaneous options:

```
internal_tracers 1  # legacy
mandatory_water 0  # legacy
mandatory sediment 0  # legacy
```

## **Light forcings:**

The short wave solar radiation (i.e up to about 5000 nm, or which approximately 40% is PAR, see Section 4) can be specified from a file:

```
LIGHT in/light.ts # get the light from a .ts text file LIGHT_INPUT_DT 1 days
ALBEDO LIGHT 0.2# legacy
```

#### Or from an internal tracer:

```
LIGHT swr # get the light from the internal swr hydro tracer (form meteorological forcing)
```

If it is coming from an internal tracer, the tracer can either be from the fully-coupled model, or in the transport model. For convenience, if the following diagnostic tracer is specified:

```
TRACER120.name PAR_z
TRACER120.long_name Downwelling PAR at top of layer
TRACER120.units mol photon m-2 s-1
```

```
TRACER120.fill value sed 0.00e+00
TRACER120.valid range sed 0.00e+00 1.00e+35
TRACER120.advect
TRACER120.diffuse
                     2
TRACER120.diagn
TRACER120.partic
TRACER120.dissol
                    0
WC3D SE
0.00e+00
TRACER120.type
                      WC3D SED ECOLOGY DIAGNOSTIC
TRACER120.psize
TRACER120.b dens
                    0.00e+00
                    0.00e+00
TRACER120.i conc
TRACER120.svel
                     0.0
                     1
TRACER120.cohesive
TRACER120.calcvol
TRACER120.floc
TRACER120.resuspend
                     1
TRACER120.deposit
TRACER120.eco flag
                        0
```

then the surface value of PAR z will be the surface PAR radiation (see Section 4).

#### Numerical scheme for ecology integration

The model uses an adaptive time step to solve the ecological ordinary differential equations (ODEs). The model configuration file allows specification of the adaptive integrator used, and the precision to which it is applied.

where the above options are the default. Integrator schemes include (dopri8, dopri5, euler1). The adaptive schemes use an approximation of the solution through the time-step (for dopri5, four and five time-steps), and use the difference in solution at the end to determine if the precision requirement has been meet. Only the solution at the end of the time step is kept.

#### By-passing ecological integration

The ecological integration is not undertaken if:

- The cell is not wet. Note whether a cell is 'wet' or 'dry' can change in time due to surface elevation changes.
- If the adaptive integrator fails to converge after NMAX = 2000 steps (this is hardcoded), the ODEs for that timestep are set to zero, the tracer ECOLOGY\_ERROR is incremented. In this case the updating excludes the whole column (wc, epibenthic and sed).
- If the master configuration file lists the cell below "EXCLUDE\_BGCSED". For example:

```
EXCLUDE_BGCSED 2 (169,15) - (170,15) 7 11
```

# **Pre-processing tools for BGC** 9

[TO BE COMPLETED]

#### **Running simulations** 10

Simulations can be run either fully coupled to one of the two hydrodynamic models available in the Environmental Modelling Suite (SHOC - Sparse Hydrodynamic Ocean Code; COMPAS - Coastal Ocean Marine Prediction Across Scales), or in an off-line mode where the current, temperature and salinity fields from an earlier hydrodynamic run are used to force the BGC model.

#### 10.1 Fully-coupled mode

By convention, model configuration files that are run in the fully-coupled mode have endings .prm.

To execute the model in the fully coupled SHOC model:

```
./shoc -p test estuary spectral.prm
```

To execute the model in the fully coupled COMPAS model:

```
./compas -p test_estuary_spectral.prm
```

#### 10.2 Off-line transport mode

There are two transport models available, one in SHOC and one in COMPAS.

This is just a quick description of the off-line transport model, more details are available in the SHOC user guide.

The following files has been saved when running the hydrodynamic model:

```
out/test estuary tran.nc
file8.name
file8.filetype
                     sparse
                     1 hour
file8.tinc
file8.bytespervalue 4
file8.vars
                     eta salt temp ulmean u2mean wmean Kzmean swr
```

The transport model then reads test\_estuary\_tran.nc and runs the BGC and sediment in an off line version. This file need to be copied across to the trans folder (so it does not get deleted).

Instead of using a test estuary spectral.prm as a main parameter file, we use a stripped version called test estuary spectral.tran (grid and model forcing are removed only replaced by transport model options, all the ecology tracer list sediment ocean boundary stay the same, expect the flag advection and diffusion for temp and salt need to be set to 0 as they are read from files).

If the model source code has been configured with the --enable-omp option during configuration before compiling (see SHOC user guide):

```
./conf/configure --enable-omp
```

Then the BGC model can be run using more than 1 CPU using the following option specified in the test\_estuary\_spectral.tran

```
eco_omp_num_threads 12 # 12 cpus in this case
```

To run the transport model in the two models:

```
./shoc -t trans/test_estuary_spectral.tran
./compas -t trans/test_estuary_spectral.tran
```

#### **Computational diagnostics** 11

The model outputs a few files during execution that are informative. Files written during the initialization

- 1. **setup.txt** describes the model configuration, including tracers, boundary conditions etc. This is from the transport model.
- 2. sedlog.txt outputs at the beginning the sediment configuration, plus any errors in the sediment model (requires VERBOSE SED to be activated in the in.tran or in.prm file).
- 3. ecology\_setup.txt describes the optical model characteristics, and lists the biological parameters read and used, as well as the processes. Arguments given in the process list, such as gases calculated in gas exchange wc etc. are also listed.
- 4. **optical setup.nc** netcdf file containing the spectrally-resolved parameters on the optical
- 5. **optical setup matlab check.m** a matlab script for plotting the optical configuration. In particular, because this file is generated during model integration it only displays optical properties that have been implemented in the simulation.
- 6. **optical\_setup\_python\_check.py** a python script for plotting the optical configuration (see above).

Files written during the initialization and model integration:

- 7. diag.txt this file is written to each ecological time step, and updates computing performance and expected finish time.
- 8. **runlog** outputs during runtime warnings and errors.

The files ecology setup.txt and setup.txt are automatically copied to the OUTPUT folder at the completion of the model initialization.

#### Mass balance 11.1

Like all biogeochemical models, maintaining mass balance is an important requirement of the biogeochemical model. The Biogeochemical Science Guide details the sink / source equations that are conserved. The model code contains checks for the conservation of carbon, nitrogen, phosphorus and oxygen. This check involves calculating the mass before and after each ecological step. The difference between the two is calculated. If the difference exceeds a relative tolerance (which is printed in ecology\_setup.txt, and is calculated as the mass difference over the original value), then the model prints an error to the runlog. The hardcoded value for the relative tolerance is 10<sup>-9</sup> for the water column and sediment, and 10<sup>-6</sup> for the epibenthic. The exception to the above calculation is oxygen, for which the mass difference is divided by the minimum of the original mass and 8000 mg m<sup>-3</sup>.

In order to implement the mass balance the process list must contain massbalance\_wc, massbalance epi and massbalance sed in the process list.

To determine mass budgets, including external inputs, circulation etc. see the SHOC manual region files.

#### **RECOM / test case 12**

#### [TO BE COMPLETED]

time series file for tracer age concentration at the river boundary (set to 0 in this age.ts: example)

dicriver.ts: time series file for tracer DIC and alkalinity concentration at the river boundary

gbr gravel 600x180.txt: text file with gravel data to initialize the model

input.nc : initial condition file created using ./shoc -g prm/bgc/prm data/input.nc

: time series file for meteorological forcing (time varying but spatially constant across the model domain)

passive0.ts: point source data file that contain NO3 and passive tracer fluxes need to be in

data.txt : text file with geographic short wave radiation bottom absorption ~ spatially variable parameter.

flow.ts : time series file with river flow temperature and salinity condition at the river boundary.

gbr\_mud\_600x180.txt : text file with mud data to initialize the model

nutrient loads.ts: time series file with nutrient and sediment concentration at the river boundary (flow is also present but not used in fully coupled, only when used in transport mode).

dicocean.ts: time series file for tracer DIC and alk concentration at the river boundary

forcing.ts: time series file for meteorological forcing (time varying but spatially constant across the model domain)

gbr sand 600x180.txt : text file with sand data to initialize the model

met2.ts: time series file for meteorological forcing (time varying but spatially constant across the model domain)