
AUTOMATIC CODE GENERATOR (ACG) FOR CONSTRUCTING USER-DEFINED
BIOGEOCHEMICAL REACTION NETWORKS

version 1.2

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INPUT TYPES

000 : Sections that should be modified by the user

000 : Sections that should NOT be modified by the user

000 : comments

000 : Maple input

000 : Maple output (appears only after you have executed the
spreadsheet)

000 : Maple input entries that have to be specified by the user

WWW : Hyperlink to the Knowledge Book

▼ Maple specific info

```
[> restart ;  
  #with(Spread) :  
[> precision := double :
```

▼ Summary and governing equations

▼ Governing equation

The governing equation solved is of the form :

theta(x)*dC/dt = -d(Fdiff+Fadv)/dx + R,

where

- dX/dp is the partial derivative of X with respect to p [M/L³/[p]
- theta(x) = A(x)*por(x) for dissolved, and A(x)*(1-por(x)) for

solid species [L²]
 - Fdiff = -(D(x)*theta(x)*dC/dx) [M/L²/T]
 - Fadv = (v*theta(x)*C)[M/L²/T]
 - A is the cross section area [L²], por is porosity [-]
 - v is the advection velocity [L/T], the sum of the water flow velocity (vwat = water flow q in [L³/T]/theta(x), solutes only) and an advective velocity w acting upon both solids and solutes (e.g. movement due to fixed reference frame)
 - D is the effective diffusion coefficient [L²/T], the sum of the molecular diffusion coefficient (Dmol, solutes only), the bioturbation coefficient (Db) and the dispersion coefficient (Disp = aL*|vwat|, solutes only, aL in [L]).
 - R is the sum of the reaction terms [M/L³/T] as specified in the reaction network established below

▼ Caveats

- for real numbers you should add a point after the number.
 - make sure all units match.
 e.g. rates and dXdt, or flux boundary conditions and concentrations for the conversion of solid to solute units, one may define temporary variables that can be used in the rate laws below:
 s_dens := 2.5; # solid density in [g/cm_solid³]
 sd := 1000. * s_dens * (1. - por(j)) / por(j);
 the factor 1.d03 converts cm³ to liter, e.g. [g/cm³] to [g/l].
 note that you need to refer to porosity exactly as por(j).

▼ Physics - Parameters

The list of length npophys & npophys2 is given by phys_name & phys_name2; the values collected in phys_val & phys_val2.

▼ Spatial and temporal domain size

tot_time: total length of simulation (T, e.g. years)

depth_max: spatial extent, e.g. total depth of simulation (L, e.g. cm)

>

Start conditions for GSA

TOC_wt := 1.0 ;

SedRate:= 0.5;

SWI_pH := 8.1;

Log_a := 1.0;

DTime:= 0.005;

SWI_a := 10**Log_a ;

SWI_TOC:= TOC_wt*2.5/12/100 ;

depth_max := 100.4 ;

tot_time := 2*depth_max/SedRate ;

#####

```
TOC_wt := 1.0
SedRate := 0.5
SWI_pH := 8.1
Log_a := 1.0
DTime := 0.005
SWI_a := 10.
SWI_TOC := 0.002083333333
depth_max := 100.4
tot_time := 401.6000000
```

(3.1.1)

Temperature and salinity

T_C: temperature (Celsius)

S : salinity (PSU)

T and S are used to calculate the molecular diffusion coefficients

T: absolute temperature (Kelvin)

```
> T_C := 10.3 ;
T := T_C + 273.15 ;
S := 35.0 ;
Depth := 100.0;
P := Depth*9.81*1027./100000.;
R := 83.145;
```

```
T_C := 10.3
T := 283.45
S := 35.0
Depth := 100.0
P := 10.07487000
R := 83.145
```

(3.1.1.1)

Transport coefficients

val: longitudinal dispersivity (L). The dispersion coefficient is calculated as $\text{Disp} = aL \cdot |v_{\text{wat}}|$

viq: depth dependency of flow. 0: constant, else changing with depth (i.e. distance)

vq0: water flow (either constant or else at $x=0$, L^3/T)

viw: depth dependency of w. 0: constant, else changing with depth (i.e. distance)

vw0: advection velocity working on solids and solutes (either constant or else at $x=0$, L/T)

viDb: depth dependency of Db. 0: constant, else changing with depth (i.e. distance)

vDb0: bioturbation coefficient working on solids and solutes (either constant or else at $x=0$, L^2/T)

- user specified profiles of $q(x)$, $w(x)$ or $Db(x)$ must be defined in the fortran subroutines `advcoeff.f` and `diffcoeff.f`, respectively.

- the molecular diffusion coefficients for the species used are specified further below

```
> val := 0.0 ;
viq := 0 ;
```

```

vq0 := 0. ;
viw := 0 ;
vw0 := SedRate; #3.3*10.0^(-0.87478367-0.00043512*
Depth);
viDb := 1 ;
vDb0 := 5.2*(10.0^(0.7624-0.0003972*Depth));
#Middelburg (water depth)
#vDb0 := 15.7*(vw0^0.69) ; #Boudreau (sedimentation
rate)

```

```

val := 0.
viq := 0
vq0 := 0.
viw := 0
vw0 := 0.5
viDb := 1
vDb0 := 27.45891316

```

(3.2.1)

Porosity profile and cross section area

vipor: depth dependency of porosity. 0: constant, else changing with depth (i.e. distance)
vpor0: porosity value (either constant or else at x=0)
viarea: depth dependency of cross section area. 0: constant, else changing with depth (i.e. distance)
varea0: cross section area (either constant or else at x=0, L²)
- user specified profiles of por(x) or area(x) must be defined in the fortran subroutine porarea.f.

```

> vipor := 1 ;
vpor0 := 0.85 ;
viarea := 0 ;
varea0 := 1.0 ;

```

```

vipor := 1
vpor0 := 0.85
viarea := 0
varea0 := 1.0

```

(3.3.1)

Grid and discretization

Dt: time step of numerical integration (T).

note that there are options defined in drivervalues.f that allow automatical selection of timestep

nnodes: number of nodes in the spatial domain.

for a regular grid, the grid spacing of the concentration profile is then depth_max/(nnodes-1)

vigridv: type of grid.

0 is for regular, evenly spaced grid,

else the user needs to specify the grid in the fortran subroutine gridsetup.f

```

> Dt := DTime;
nnodes := 121;
vigrid := 1;

```

```

Dt := 0.005

```

```
nnodes := 121
vgrid := 1
```

(3.5.1)

Reaction Network - Size and Variables

Size of reaction network

nsolids : number of solid species

ndissolved : number of dissolved species

ncompo : total number of species

nreactions : total number of reactions (including equilibrium rxns)

negrxns : number of equilibrium reaction

```
> nsolids := 26;
   ndissolved := 29;
   ncompo := nsolids + ndissolved;
   nreactions := 60;
   negrxns := 13 ;
```

```
nsolids := 26
ndissolved := 29
ncompo := 55
nreactions := 60
negrxns := 13
```

(4.1.1)

List of variables

variables: list of variables to model. example:

listsolids: species number which is a SOLID species.

note: all other variables are temporary and are NOT parsed to the ACG

Example:

```
variables:=[O2, so4, MnOx, FeOx, hco3, co3, hplus, hs];
listsolids:= [3,4];
```

```
> variables := [ch2o, o2, no3, mno2, feoh3, so4, ch4,
nh4, po4, mn, fe, h2s, hs, ch4g, h2co3, hco3, co3,
boh4, boh3, hplus, caco3, ca, snh4, spo4, fes,
feco3, s0, fes2, sfe, sfp, mnco3, age, mno2pr,
feoh3mr, feoh3pr, arago, mgcal, mg, BSi, DSi, sfSi,
al, aloh, aloh2, aloh3, aloh4, glass, plagio, na,
kplus, olive, pyrox, illite, smect, kaoli] ;
```

```
listsolids := [1, 4, 5, 21, 23, 24, 25, 26, 28, 29,
30, 31, 32, 33, 34, 35, 36, 37, 39, 41, 47, 48, 51,
52, 53, 54, 55] ;
```

```
variables := [ch2o, o2, no3, mno2, feoh3, so4, ch4, nh4, po4, mn, fe, h2s, hs, ch4g,
h2co3, hco3, co3, boh4, boh3, hplus, caco3, ca, snh4, spo4, fes, feco3, s0, fes2, sfe,
sfp, mnco3, age, mno2pr, feoh3mr, feoh3pr, arago, mgcal, mg, BSi, DSi, sfSi, al,
aloh, aloh2, aloh3, aloh4, glass, plagio, na, kplus, olive, pyrox, illite, smect, kaoli]
```

```
listsolids := [1, 4, 5, 21, 23, 24, 25, 26, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 39, 41, 47, 48, 51, 52, 53, 54, 55] (4.2.1)
```

Biogeochemistry - Rate laws

Definition of **kinetic rate laws**

rate.i : array of rates.

- **For equilibrium rate expression, a kinetic rate MUST be specified as well.** It will be overwritten in the equilibrium section below, but you need it as space holder and stoichiometry. Furthermore, the steadystate module uses detailed balancing method with fast kinetics. Therefore, in the example below, kf will have to be defined as a rather large number and kb = kf*Keq

note: all other variables are temporary and are NOT parsed to the ACG

- **conditional statements:** if a rate law depends on a conditional statement you need to make use of the subroutine switches.f. Example: dissolution (Rd) is only to take place at undersaturation, thus $Rd = f(\text{saturation})$. If $\text{saturation} > 1$, $Rd > 0$, else $Rd = 0$. This can be implemented as $Rd = k \cdot H1 \cdot \text{saturation}$, where H1 is toggled between 0 and 1. Rather than giving the condition here in maple, for now you need to do this in "switches.f", where you program the conditions for H1, e.g. $H1 = 0$, If $(A \cdot B / K > 1)$ then $H1 = 1$.

example:

```
rate1 := 1000.*O2*hs; # rate law for 2O2 + HS -> SO4 + Hplus
rate2 := kf*hplus*co3 - kb*hco3; # kinetic rate law for HCO3 = CO3 +
Hplus (equilibrium)
```

Primary redox reactions [WWW](#)

```
> #primary redox
#attpor:=5.0e-1;
#porX:=0.70;

#age :=sw01*1e-10 + (1.0-sw01)*(((1-porX)*(x_pos-
xbiot)+1.0/attpor*(vpor0-porX)*(exp(-attpor*(x_pos-
xbiot))-1.0))/(vw0*(1-vpor0)));
fo2 :=(sw02+(1.0-sw02)*o2/kmo2);
fno3 :=sw03*(1.0-fo2)*(sw04+(1.0-sw04)*no3/kmno3);
fmno2 :=sw05*(1.0-fo2-fno3)*(sw06+(1.0-sw06)*
mno2/kmmno2);
ffeh3:=sw07*(1.0-fo2-fno3-fmno2)*(sw08+(1.0-sw08)*
feh3/kmfeh3);
fso4 :=sw09*(1.0-fo2-fno3-fmno2-ffeh3)*(sw10+(1.0-
sw10)*so4/kmso4);
fch4 :=sw11*(1.0-fo2-fno3-fmno2-ffeh3-fso4);

rate1 := 0.0; # nu/(inita+age)*ch2o;
rate2 := nu/(inita+age)*ch2o*fo2;
rate3 := nu/(inita+age)*ch2o*fno3;
rate4 := nu/(inita+age)*ch2o*fmno2;
rate5 := nu/(inita+age)*ch2o*ffeh3;
rate6 := nu/(inita+age)*ch2o*fso4;
rate7 := nu/(inita+age)*ch2o*fch4;
```

```

#secondary redox
rate8 := knit*nh4*o2;
rate9 := kmnox*mn*o2;
rate10:= kfemno2*fe*mno2;
rate37:= kfemno2pr*fe*mno2pr; # FSF PR MnO2
rate11:= kfeo2*fe*o2;
rate12:= kh2so2*(h2s+hs)*o2;
rate13:= kh2smno2*mno2*(hs+h2s);
rate38:= kh2smno2pr*mno2pr*(hs+h2s); # FSF PR MnO2
rate14:= kh2sfeoh3*feoh3*(hs+h2s);
rate40:= kh2sfeoh3mr*feoh3mr*(hs+h2s); # FSF MR
FeOH3
rate41:= kh2sfeoh3pr*feoh3pr*(hs+h2s); # FSF PR
FeOH3
rate15:= kaom*ch4*so4;
rate16:= kaomo2*ch4*o2;
rate17:= kfeso2*fes*o2;

#mineral
rate18:= kmnco3precip*sw12*((mn*co3/KsMnCO3)-1.0);
rate19:= kfesprecip*sw13*((fe*hs)/(hplus*KsFeS))-1.0);
rate20:= kfesdiss*(1.0-sw13)*fes*(1.0-((fe*hs)/(hplus*KsFeS)));
rate21:= kfeco3precip*sw14*((fe*co3/KsFeCO3)-1.0);
rate22:= kpyr*fes*(h2s+hs);
rate23:= kfess0*fes*s0;
rate24:= sw15*kcaldiss*caco3*((1.0 - (ca*(co3)/kspcal)));
rate25:= sw16*kapa*(po4-po4_eq);
rate43:= sw20*kcalppt*((ca*co3/kspcal)-5.0); #FSF
CaCO3 (calcite) precipitation
rate44:= sw21*karadis*arago*((1.0 - (ca*co3/kspara)));
rate45:= sw22*karappt*((ca*co3/kspara)-5.0);

#misc
rate26:= (1.0-sw17)*kdis*sw18*(ch4g)*(ch4eq-(ch4));
rate27:= sw17*kgas*(ch4-ch4eq);
rate28:= kdi*sw19*s0;
rate39:= kmnage*mno2; #FSF HR MnO2 ageing into
unreac MnO2
rate42:= kfeage*feoh3; #FSF HR FeOH3 ageing into MR
FeOH3

#Silicon

```

```

#Biogenic Silica dissolution
kBSidis_z := (kBSidis*bd) + ((kBSidis - kBSidis*bd)*
exp(-ad*x_pos));
rate48 := sw25*kBSidis_z*BSi*(1.0-DSi/BSisat) ;

#eq
rate29:= kf1*hplus*hco3-kb1*h2co3;
rate30:= kf2*hplus*co3-kb2*hco3;
rate31:= kf3*hplus*hs-kb3*h2s ;
rate32:= kf4*hplus*boh4-kb4*boh3;
rate33:= kf5*po4*por(j) - kb5*spo4*(1. - por(j));
rate34:= kf6*nh4*por(j) - kb6*snh4*(1. - por(j));
rate35:= kf7*(feoh3+feoh3mr+feoh3pr)*po4- kb7*sfp;
#FSF Iron fractions;
rate36:= kf8*fe*por(j) - kb8*sfe*(1. - por(j));

#eq Silicon
rate49:= kf9*(feoh3+feoh3mr+feoh3pr)*(DSi) - kb9*
(sfSi);

#eq aluminium
rate50:= kf10*aloh*hplus - kb10*al;    #reformulate
as pH eq51,52,53
rate51:= kf11*aloh2*hplus - kb11*aloh;
rate52:= kf12*aloh3*hplus - kb12*aloh2;
rate53:= kf13*aloh4*hplus - kb13*aloh3;

#Silicate weathering
rate54:= sw26*kglass*glass*(hplus**3/al)**(1/3) *
(1.0 - DSi*al**0.36/hplus**1.08/kspglass);
rate55:= sw27*kplagio*plagio*(hplus**3/al)**0.35 *
(1.0 - DSi**2.3*al**1.7*na**0.7*ca**0.3/kspplagio);
rate56:= sw28*kolive*olive * (1.0 - DSi*mg**1.6*fe**
0.4/hplus**4/kspolive);
rate57:= sw29*kpyrox*pyrox * (1.0 - DSi**2 * fe**
0.46 * mg**0.84 * ca**0.7 / hplus**4 / kspprox);

#Silicate reverse weathering
rate58:= sw30*killite * (DSi**3.5 * al**2.3 * kplus*
*0.6 * mg**0.25 / hplus**8 / kspillite - 1.0);
rate59:= sw31*ksmect * (ca**0.165 * mg**0.33 * al**
1.7 * DSi**4 /hplus**6 / kspsmect - 1.0);
rate60:= sw32*kkaoli * (al**2 * DSi**2 / hplus**6/
kspkaoli - 1.0);

```

$$fo2 := sw02 + \frac{(1.0 - sw02) \, o2}{kmo2}$$

[illegible]

$$\begin{aligned}
& - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 \right. \\
& + \left. \frac{(1.0 - sw04) no3}{kmno3} \right) - sw05 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 \right. \right. \\
& - \left. \left. sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 + \frac{(1.0 - sw04) no3}{kmno3} \right) \right) \left(sw06 \right. \\
& + \left. \frac{(1.0 - sw06) mno2}{kmmno2} \right) \left(sw08 + \frac{(1.0 - sw08) feoh3}{kmfeoh3} \right) - sw09 \left(1.0 - sw02 \right. \\
& - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 \right. \\
& + \left. \frac{(1.0 - sw04) no3}{kmno3} \right) - sw05 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 \right. \right. \\
& - \left. \left. sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 + \frac{(1.0 - sw04) no3}{kmno3} \right) \right) \left(sw06 \right. \\
& + \left. \frac{(1.0 - sw06) mno2}{kmmno2} \right) - sw07 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 \right. \right. \\
& - \left. \left. sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 + \frac{(1.0 - sw04) no3}{kmno3} \right) - sw05 \left(1.0 \right. \right. \\
& - \left. \left. sw02 - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 \right. \right. \\
& + \left. \frac{(1.0 - sw04) no3}{kmno3} \right) \left(sw06 + \frac{(1.0 - sw06) mno2}{kmmno2} \right) \left(sw08 \right. \\
& + \left. \frac{(1.0 - sw08) feoh3}{kmfeoh3} \right) \left(sw10 + \frac{(1.0 - sw10) so4}{kms04} \right)
\end{aligned}$$

$$rate1 := 0.$$

$$rate2 := \frac{v ch2o \left(sw02 + \frac{(1.0 - sw02) o2}{kmo2} \right)}{inita + age}$$

$$rate3 :=$$

$$\frac{v ch2o sw03 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 + \frac{(1.0 - sw04) no3}{kmno3} \right)}{inita + age}$$

$$rate4 := \frac{1}{inita + age} \left(v ch2o sw05 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 \right. \right. \right. \\
- \left. \left. sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 + \frac{(1.0 - sw04) no3}{kmno3} \right) \right) \left(sw06 \right. \\
+ \left. \frac{(1.0 - sw06) mno2}{kmmno2} \right) \left. \right)$$

$$rate5 := \frac{1}{inita + age} \left(v ch2o sw07 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 \right. \right. \right. \\
- \left. \left. sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 + \frac{(1.0 - sw04) no3}{kmno3} \right) - sw05 \left(1.0 \right. \right. \\
- \left. \left. sw02 - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 \right. \right. \right.$$

$$\begin{aligned}
& + \frac{(1.0 - sw04) no3}{kmno3} \Big) \Big) \left(sw06 + \frac{(1.0 - sw06) mno2}{kmmno2} \right) \Big) \left(sw08 \right. \\
& \left. + \frac{(1.0 - sw08) feoh3}{kmfeoh3} \right) \Big) \\
rate6 := & \frac{1}{inita + age} \left(v\ ch2o\ sw09 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 \right. \right. \right. \\
& - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \Big) \left(sw04 + \frac{(1.0 - sw04) no3}{kmno3} \right) - sw05 \left(1.0 \right. \\
& - sw02 - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 \right. \\
& + \frac{(1.0 - sw04) no3}{kmno3} \Big) \Big) \left(sw06 + \frac{(1.0 - sw06) mno2}{kmmno2} \right) - sw07 \left(1.0 - sw02 \right. \\
& - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 \right. \\
& + \frac{(1.0 - sw04) no3}{kmno3} \Big) - sw05 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 \right. \right. \\
& - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \Big) \left(sw04 + \frac{(1.0 - sw04) no3}{kmno3} \right) \Big) \left(sw06 \right. \\
& + \frac{(1.0 - sw06) mno2}{kmmno2} \Big) \Big) \left(sw08 + \frac{(1.0 - sw08) feoh3}{kmfeoh3} \right) \Big) \left(sw10 \right. \\
& \left. + \frac{(1.0 - sw10) so4}{kms04} \right) \Big) \\
rate7 := & \frac{1}{inita + age} \left(v\ ch2o\ sw11 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 \right. \right. \right. \\
& - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \Big) \left(sw04 + \frac{(1.0 - sw04) no3}{kmno3} \right) - sw05 \left(1.0 \right. \\
& - sw02 - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 \right. \\
& + \frac{(1.0 - sw04) no3}{kmno3} \Big) \Big) \left(sw06 + \frac{(1.0 - sw06) mno2}{kmmno2} \right) - sw07 \left(1.0 - sw02 \right. \\
& - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 \right. \\
& + \frac{(1.0 - sw04) no3}{kmno3} \Big) - sw05 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 \right. \right. \\
& - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \Big) \left(sw04 + \frac{(1.0 - sw04) no3}{kmno3} \right) \Big) \left(sw06 \right. \\
& + \frac{(1.0 - sw06) mno2}{kmmno2} \Big) \Big) \left(sw08 + \frac{(1.0 - sw08) feoh3}{kmfeoh3} \right) - sw09 \left(1.0 - sw02 \right. \\
& - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 \right. \\
& + \frac{(1.0 - sw04) no3}{kmno3} \Big) - sw05 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 \right. \right. \\
& - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \Big) \left(sw04 + \frac{(1.0 - sw04) no3}{kmno3} \right) \Big) \left(sw06 \right.
\end{aligned}$$

$$\begin{aligned}
& + \frac{(1.0 - sw06) mno2}{kmmno2} \Big) - sw07 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 \right. \right. \\
& \left. \left. - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 + \frac{(1.0 - sw04) no3}{kmno3} \right) - sw05 \left(1.0 \right. \right. \\
& \left. \left. - sw02 - \frac{(1.0 - sw02) o2}{kmo2} - sw03 \left(1.0 - sw02 - \frac{(1.0 - sw02) o2}{kmo2} \right) \left(sw04 \right. \right. \right. \\
& \left. \left. + \frac{(1.0 - sw04) no3}{kmno3} \right) \right) \left(sw06 + \frac{(1.0 - sw06) mno2}{kmmno2} \right) \Big) \left(sw08 \right. \\
& \left. + \frac{(1.0 - sw08) feoh3}{kmfeoh3} \right) \Big) \left(sw10 + \frac{(1.0 - sw10) so4}{kms04} \right) \Big) \Big)
\end{aligned}$$

$$rate8 := knit nh4 o2$$

$$rate9 := kmnox mn o2$$

$$rate10 := kfemno2 fe mno2$$

$$rate37 := kfemno2pr fe mno2pr$$

$$rate11 := kfeo2 fe o2$$

$$rate12 := kh2so2 (h2s + hs) o2$$

$$rate13 := kh2smno2 mno2 (h2s + hs)$$

$$rate38 := kh2smno2pr mno2pr (h2s + hs)$$

$$rate14 := kh2sfeoh3 feoh3 (h2s + hs)$$

$$rate40 := kh2sfeoh3mr feoh3mr (h2s + hs)$$

$$rate41 := kh2sfeoh3pr feoh3pr (h2s + hs)$$

$$rate15 := kaom ch4 so4$$

$$rate16 := kaomo2 ch4 o2$$

$$rate17 := kfeso2 fes o2$$

$$rate18 := kmnco3precip sw12 \left(\frac{mn co3}{KsMnCO3} - 1.0 \right)$$

$$rate19 := kfesprecip sw13 \left(\frac{fe hs}{hplus KsFeS} - 1.0 \right)$$

$$rate20 := kfesdiss (1.0 - sw13) fes \left(1.0 - \frac{fe hs}{hplus KsFeS} \right)$$

$$rate21 := kfeco3precip sw14 \left(\frac{fe co3}{KsFeCO3} - 1.0 \right)$$

$$rate22 := kpyr fes (h2s + hs)$$

$$rate23 := kfess0 fes s0$$

$$rate24 := sw15 kcal diss caco3 \left(1.0 - \frac{ca co3}{kspcal} \right)$$

$$rate25 := sw16 kapa (po4 - po4_{eq})$$

$$rate43 := sw20 kcal ppt \left(\frac{ca co3}{kspcal} - 5.0 \right)$$

$$rate44 := sw21 karadis arago \left(1.0 - \frac{ca co3}{kspara} \right)$$

$$rate45 := sw22 karappt \left(\frac{ca co3}{kspara} - 5.0 \right)$$

$$rate26 := (1.0 - sw17) kdis sw18 ch4g (ch4eq - ch4)$$

$$rate27 := sw17 kgas (ch4 - ch4eq)$$

$$rate28 := kdi sw19 s0$$

$$\begin{aligned}
\text{rate39} &:= \text{kmnage mno2} \\
\text{rate42} &:= \text{kfeage feoh3} \\
\text{kBSidis_z} &:= \text{kBSidis bd} + (\text{kBSidis} - \text{kBSidis bd}) e^{-\text{ad x_pos}} \\
\text{rate48} &:= \text{sw25} \left(\text{kBSidis bd} + (\text{kBSidis} - \text{kBSidis bd}) e^{-\text{ad x_pos}} \right) \text{BSi} \left(1.0 - \frac{\text{DSi}}{\text{BSisat}} \right) \\
\text{rate29} &:= \text{kf1 hplus hco3} - \text{kb1 h2co3} \\
\text{rate30} &:= \text{kf2 hplus co3} - \text{kb2 hco3} \\
\text{rate31} &:= \text{kf3 hplus hs} - \text{kb3 h2s} \\
\text{rate32} &:= \text{kf4 hplus boh4} - \text{kb4 boh3} \\
\text{rate33} &:= \text{kf5 po4 por(j)} - \text{kb5 spo4} (1. - \text{por(j)}) \\
\text{rate34} &:= \text{kf6 nh4 por(j)} - \text{kb6 snh4} (1. - \text{por(j)}) \\
\text{rate35} &:= \text{kf7 (feoh3 + feoh3mr + feoh3pr) po4} - \text{kb7 sfp} \\
\text{rate36} &:= \text{kf8 fe por(j)} - \text{kb8 sfe} (1. - \text{por(j)}) \\
\text{rate49} &:= \text{kf9 (feoh3 + feoh3mr + feoh3pr) DSi} - \text{kb9 sfSi} \\
\text{rate50} &:= \text{kf10 aloh hplus} - \text{kb10 al} \\
\text{rate51} &:= \text{kf11 aloh2 hplus} - \text{kb11 aloh} \\
\text{rate52} &:= \text{kf12 aloh3 hplus} - \text{kb12 aloh2} \\
\text{rate53} &:= \text{kf13 aloh4 hplus} - \text{kb13 aloh3} \\
\text{rate54} &:= \text{sw26 kglass glass} \left(\frac{\text{hplus}^3}{\text{al}} \right)^{1/3} \left(1.0 - \frac{\text{DSi al}^{0.36}}{\text{hplus}^{1.08} \text{kspglass}} \right) \\
\text{rate55} &:= \text{sw27 kplagio plagio} \left(\frac{\text{hplus}^3}{\text{al}} \right)^{0.35} \left(1.0 - \frac{\text{DSi}^{2.3} \text{al}^{1.7} \text{na}^{0.7} \text{ca}^{0.3}}{\text{kspplagio}} \right) \\
\text{rate56} &:= \text{sw28 kolive olive} \left(1.0 - \frac{\text{DSi mg}^{1.6} \text{fe}^{0.4}}{\text{hplus}^4 \text{kspolive}} \right) \\
\text{rate57} &:= \text{sw29 kpyrox pyrox} \left(1.0 - \frac{\text{DSi}^2 \text{fe}^{0.46} \text{mg}^{0.84} \text{ca}^{0.7}}{\text{hplus}^4 \text{ksppyrox}} \right) \\
\text{rate58} &:= \text{sw30 killite} \left(\frac{\text{DSi}^{3.5} \text{al}^{2.3} \text{kplus}^{0.6} \text{mg}^{0.25}}{\text{hplus}^8 \text{kspillite}} - 1.0 \right) \\
\text{rate59} &:= \text{sw31 ksmect} \left(\frac{\text{ca}^{0.165} \text{mg}^{0.33} \text{al}^{1.7} \text{DSi}^4}{\text{hplus}^6 \text{kpsmect}} - 1.0 \right) \\
\text{rate60} &:= \text{sw32 kkaoli} \left(\frac{\text{al}^2 \text{DSi}^2}{\text{hplus}^6 \text{kspkaoli}} - 1.0 \right)
\end{aligned} \tag{5.1.1}$$

Biogeochemistry – Stoichiometry

Stoichiometry of the biogeochemical reactions

d.sp.dt : rates of change of sp due to the sum of biogeochemical reactions

note that rateX must be referred to as rX

example:

dO2dt := -2*r1;

dhco3dt = -r2;

> SD := (1.0 - por(j)) / por(j);

```

SD1:= 1/por(j);
x  := 106;
y  := 12;
z  := 1;

```

$$SD := \frac{1.0 - por(j)}{por(j)}$$

$$SD1 := \frac{1}{por(j)}$$

$$x := 106$$

$$y := 12$$

$$z := 1$$

(6.1)

```

> dch2odt := 0.0; # PP
dagedt   := 0.0; # PP
do2dt    := -(x+2*y)/x*SD*r2-2.0*r8-2.0*r16-2.0*r12
-0.25*r11-2.0*SD*r17-0.5*r9;
dno3dt   := -(4.0*x+3.0*y)/(5.0*x)*SD*r3+r8;
dmno2dt  := -2.0*r4+r9/SD-r10-r13-r39; # FSF (r39 =
ageing);
dmno2prdt := -r37-r38+r39; # FSF PR MnO2 rate
dfeoh3dt := -4*r5-2.0*r14+1.0*r11/SD+2.0*r10+2.0*r37-
r42; #FSF (r42 = ageing);
dfeoh3mrdt := -2.0*r40+r42; #FSF MR FeOH3 rate;
dfeoh3prdt := -2.0*r41; #FSF PR FeOH3 rate;
dso4dt    := -0.5*SD*r6-r15+0.5*r12+SD*r17;#+r28;
dch4dt    := 0.5*SD*r7-r15-r16+r26-r27;
dch4gdt   := -r26+r27;

dnh4dt    := y/x*SD*( (r2+r3+r4+r5+r6+r7) -r3)-r8-r34*
SD1; # PP
dpo4dt    := z/x*SD*( r2+r3+r4+r5+r6+r7 )-r25-r33*SD1-
r35*SD1; # PP

#dnh4dt    := y/x*SD*(r1-r3)-r8-r34*SD1;
#dpo4dt    := z/x*SD*(r1)-r25-r33*SD1-r35*SD1;

dmndt     := 2.0*SD*r4-r9+(r10+r37)*SD+(r13+r38)*SD-1.0*
r18; #FSF
dfedt     := 4.0*SD*r5+2.0*SD*(r14+r40+r41)-1.0*r11+SD*
r17-1.0*r19-1.0*r21+SD*r20-r36*SD1-2.0*(r10+r37)*
SD+0.46*SD*r67 +0.4*SD*r66; #FSF

dh2sdt    := 0.5*SD*r6+r15-r12+r31-1.0*SD*(r14+r40+r41)
-1.0*r19-1.0*SD*r22+3.0*r28+SD*r20-(r13+r38)*SD; #FSF
dhsdt     := -r31;

```

```
dh2co3dt:=((x+y+2.0*z)/x)*SD*r2    +((x-3.0*y+10.0*z)/
(5.0*x))*SD*r3    -(3.0*x+y-2.0*z)*SD*r4/x    -((y-2.0*
z)/x)*SD*r6    +((x-2.0*y+4.0*z)/(2.0*x))*SD*r7    -
(7.0*x+y-2.0*z)/x*SD*r5    +2.0*r8    -r15    +r16
+2.0*r12    -4.0*SD*(r14+r40+r41)    +r29    +2.0*r11
+2.0*r19    +2.0*r28    -2.0*SD*r20    +2.0*r9    +2.0*
(r10+r37)*SD    -2.0*(r13+r38)*SD    -1.08*SD*r54 +8.0*
r58 -4.0*SD*r57 -4.0*SD*r56 +6.0*r59 +6.0*r60 ; #FSF +
r18+1.0*r21
```

```
dhco3dt:=-((y+2.0*z)/x)*SD*r2    +((4.0*x+3.0*y-10.0*z)
/(5.0*x))*SD*r3    +(4.0*x+y-2.0*z)*SD*r4/x    +((x+
y-2.0*z)/x)*SD*r6    +((y-2.0*z)/x)*SD*r7    +(8.0*x+
y-2.0*z)/x*SD*r5    -2.0*r8    +2.0*r15    -2.0*r12
+4.0*SD*(r14+r40+r41)    -r29    +r30    -2.0*r11    -2.0*
r19    -2.0*r28    +2.0*SD*r20    -2.0*r9    -2.0*(r10+
r37)*SD    +2.0*(r13+r38)*SD +1.08*SD*r54 -8.0*r58
+4.0*SD*r57 +4.0*SD*r56 -6.0*r59 -6.0*r60 ; #-2.0*
r18*SD-2.0*SD*r21
```

```
dco3dt:=-r30+SD*r24-r43+SD*r44-r45+SD*r46-r47-r18-r21;
#FSF r43: CaCO3 prec
```

```
dboh4dt:=-r32;
dboh3dt:=r32;
```

```
dhplusdt:=-r29-r30-r31-r32-r50-r51-r52-r53; #-1.08*SD*
r54+8.0*r58-4.0*SD*r57-4.0*SD*r56+6.0*r59+6.0*r60;
```

```
dcaco3dt:=-r24+1.0*r43/SD; #FSF r43: CaCO3 prec
dcadt:=SD*r24-r43+SD*r44-r45+SD*r46-r47+0.7*SD*
r57+0.3*SD*r55-0.165*r59; #FSF r43: CaCO3 prec ##
dsnh4dt:=r34/(1.0-por(j));
dsp04dt:=r33/(1.0-por(j));
dfesdt:=(1.0*r19/SD-1.0*r17-1.0*r22-1.0*r23-r20);
dfeco3dt:=(1.0*r21/SD);
dmnco3dt:=(1.0*r18/SD);
dfes2dt:=1.0*r22+1.0*r23;
ds0dt:=1.0*SD*(r14+r40+r41)-4.0*r28-1.0*SD*r23+1.0*SD*
(r13+r38); #FSF
dsfedt:=r36/(1.0-por(j));
dsfpdt:=r35/(1.0-por(j));
```

```
daragodt:=-r44+r45/SD;
dmgcaldt:=-r46+r47/SD ; ## Ikaite
dmgdt:=0.84*SD*r57 +1.6*SD*r56-0.25*r58-0.33*r59;
```

dBSidt := -r48;

dDSidt := r48*SD-r49*SD1+SD*r56+r54*SD+2.3*SD*r55-3.5*r58+2.0*SD*r57-4.0*r59-2.0*r60;

dsfSidt:= r49/(1.0 - por(j));

dalddt := 0.36*SD*r54-2.3*r58+0.0025*SD*r48-1.7*r59-2.0*r60+r50+1.7*SD*r55;

dalohdt := -r50 +r51;

daloh2dt:= -r51 +r52;

daloh3dt:= -r52 +r53;

daloh4dt:= -r53;

dglassdt:= -r54;

dplagiodydt:= -r55;

dolivedt:= -r56;

dpyroxdt:= -r57;

dillitedt:= r58/SD;

dsmechdt:= r59/SD;

dkaoliddt:= r60/SD;

dnadt := 0.7*SD*r55;

dkplusdt := -0.6*r58;

dch2odt := 0.

dagedt := 0.

$$\begin{aligned} do2dt := & -\frac{65}{53} \frac{(1.0 - por(j)) r2}{por(j)} - 2.0 r8 - 2.0 r16 - 2.0 r12 - 0.25 r11 \\ & - \frac{2.0 (1.0 - por(j)) r17}{por(j)} - 0.5 r9 \end{aligned}$$

$$dno3dt := -\frac{0.8679245284 (1.0 - por(j)) r3}{por(j)} + r8$$

$$dmno2dt := -2.0 r4 + \frac{r9 por(j)}{1.0 - por(j)} - r10 - r13 - r39$$

$$dmno2prdt := -r37 - r38 + r39$$

$$dfeoh3dt := -4 r5 - 2.0 r14 + \frac{1.0 r11 por(j)}{1.0 - por(j)} + 2.0 r10 + 2.0 r37 - r42$$

$$dfeoh3mrdt := -2.0 r40 + r42$$

$$dfeoh3prdt := -2.0 r41$$

$$\begin{aligned}
dso4dt &:= - \frac{0.5 (1.0 - por(j)) r6}{por(j)} - r15 + 0.5 r12 + \frac{(1.0 - por(j)) r17}{por(j)} \\
dch4dt &:= \frac{0.5 (1.0 - por(j)) r7}{por(j)} - r15 - r16 + r26 - r27 \\
dch4gdt &:= -r26 + r27 \\
dnh4dt &:= \frac{6}{53} \frac{(1.0 - por(j)) (r2 + r4 + r5 + r6 + r7)}{por(j)} - r8 - \frac{r34}{por(j)} \\
dpo4dt &:= \frac{1}{106} \frac{(1.0 - por(j)) (r2 + r3 + r4 + r5 + r6 + r7)}{por(j)} - r25 - \frac{r33}{por(j)} \\
&\quad - \frac{r35}{por(j)} \\
dmndt &:= \frac{2.0 (1.0 - por(j)) r4}{por(j)} - r9 + \frac{(r10 + r37) (1.0 - por(j))}{por(j)} \\
&\quad + \frac{(r13 + r38) (1.0 - por(j))}{por(j)} - 1.0 r18 \\
dfedt &:= \frac{4.0 (1.0 - por(j)) r5}{por(j)} + \frac{2.0 (1.0 - por(j)) (r14 + r40 + r41)}{por(j)} - 1.0 r11 \\
&\quad + \frac{(1.0 - por(j)) r17}{por(j)} - 1.0 r19 - 1.0 r21 + \frac{(1.0 - por(j)) r20}{por(j)} - \frac{r36}{por(j)} \\
&\quad - \frac{2.0 (r10 + r37) (1.0 - por(j))}{por(j)} + \frac{0.46 (1.0 - por(j)) r67}{por(j)} \\
&\quad + \frac{0.4 (1.0 - por(j)) r66}{por(j)} \\
dh2sdt &:= \frac{0.5 (1.0 - por(j)) r6}{por(j)} + r15 - r12 + r31 \\
&\quad - \frac{1.0 (1.0 - por(j)) (r14 + r40 + r41)}{por(j)} - 1.0 r19 - \frac{1.0 (1.0 - por(j)) r22}{por(j)} \\
&\quad + 3.0 r28 + \frac{(1.0 - por(j)) r20}{por(j)} - \frac{(r13 + r38) (1.0 - por(j))}{por(j)} \\
dhsdt &:= -r31 \\
dh2co3dt &:= - \frac{0.09433962264 (1.0 - por(j)) r6}{por(j)} + \frac{2.0 (r10 + r37) (1.0 - por(j))}{por(j)} \\
&\quad + \frac{1.132075472 (1.0 - por(j)) r2}{por(j)} + 2.0 r8 + r16 + 2.0 r12 + 2.0 r11 + 2.0 r9 \\
&\quad - r15 + 2.0 r19 + 2.0 r28 + r29 + 8.0 r58 + 6.0 r59 + 6.0 r60 \\
&\quad - \frac{3.094339623 (1.0 - por(j)) r4}{por(j)} - \frac{7.094339623 (1.0 - por(j)) r5}{por(j)} \\
&\quad - \frac{4.0 (1.0 - por(j)) (r14 + r40 + r41)}{por(j)} + \frac{0.1509433962 (1.0 - por(j)) r3}{por(j)} \\
&\quad + \frac{0.4056603774 (1.0 - por(j)) r7}{por(j)} - \frac{2.0 (r13 + r38) (1.0 - por(j))}{por(j)} \\
&\quad - \frac{2.0 (1.0 - por(j)) r20}{por(j)} - \frac{1.08 (1.0 - por(j)) r54}{por(j)} - \frac{4.0 (1.0 - por(j)) r57}{por(j)} \\
&\quad - \frac{4.0 (1.0 - por(j)) r56}{por(j)}
\end{aligned}$$

$$\begin{aligned}
dhco3dt &:= \frac{1.094339623 (1.0 - por(j)) r6}{por(j)} - \frac{2.0 (r10 + r37) (1.0 - por(j))}{por(j)} \\
&- \frac{0.1320754717 (1.0 - por(j)) r2}{por(j)} - 2.0 r8 - 2.0 r12 - 2.0 r11 - 2.0 r9 \\
&+ 2.0 r15 - 2.0 r19 - 2.0 r28 - r29 - 8.0 r58 - 6.0 r59 - 6.0 r60 + r30 \\
&+ \frac{4.094339623 (1.0 - por(j)) r4}{por(j)} + \frac{8.094339623 (1.0 - por(j)) r5}{por(j)} \\
&+ \frac{4.0 (1.0 - por(j)) (r14 + r40 + r41)}{por(j)} + \frac{0.8490566038 (1.0 - por(j)) r3}{por(j)} \\
&+ \frac{0.09433962264 (1.0 - por(j)) r7}{por(j)} + \frac{2.0 (r13 + r38) (1.0 - por(j))}{por(j)} \\
&+ \frac{2.0 (1.0 - por(j)) r20}{por(j)} + \frac{1.08 (1.0 - por(j)) r54}{por(j)} + \frac{4.0 (1.0 - por(j)) r57}{por(j)} \\
&+ \frac{4.0 (1.0 - por(j)) r56}{por(j)} \\
dco3dt &:= -r30 + \frac{(1.0 - por(j)) r24}{por(j)} - r43 + \frac{(1.0 - por(j)) r44}{por(j)} - r45 \\
&+ \frac{(1.0 - por(j)) r46}{por(j)} - r47 - r18 - r21 \\
dboh4dt &:= -r32 \\
dboh3dt &:= r32 \\
dhplusdt &:= -r29 - r30 - r31 - r32 - r50 - r51 - r52 - r53 \\
dcaco3dt &:= -r24 + \frac{1.0 r43 por(j)}{1.0 - por(j)} \\
dcadt &:= \frac{(1.0 - por(j)) r24}{por(j)} - r43 + \frac{(1.0 - por(j)) r44}{por(j)} - r45 \\
&+ \frac{(1.0 - por(j)) r46}{por(j)} - r47 + \frac{0.7 (1.0 - por(j)) r57}{por(j)} \\
&+ \frac{0.3 (1.0 - por(j)) r55}{por(j)} - 0.165 r59 \\
dsnh4dt &:= \frac{r34}{1.0 - por(j)} \\
dsp4dt &:= \frac{r33}{1.0 - por(j)} \\
dfesdt &:= \frac{1.0 r19 por(j)}{1.0 - por(j)} - 1.0 r17 - 1.0 r22 - 1.0 r23 - r20 \\
dfeco3dt &:= \frac{1.0 r21 por(j)}{1.0 - por(j)} \\
dmnco3dt &:= \frac{1.0 r18 por(j)}{1.0 - por(j)} \\
dfes2dt &:= 1.0 r22 + 1.0 r23 \\
ds0dt &:= \frac{1.0 (1.0 - por(j)) (r14 + r40 + r41)}{por(j)} - 4.0 r28 - \frac{1.0 (1.0 - por(j)) r23}{por(j)} \\
&+ \frac{1.0 (r13 + r38) (1.0 - por(j))}{por(j)}
\end{aligned}$$

$$\begin{aligned}
dsfedt &:= \frac{r36}{1.0 - por(j)} \\
dsfpdt &:= \frac{r35}{1.0 - por(j)} \\
daragodt &:= -r44 + \frac{r45 por(j)}{1.0 - por(j)} \\
dmgcaldt &:= -r46 + \frac{r47 por(j)}{1.0 - por(j)} \\
dmgdt &:= \frac{0.84 (1.0 - por(j)) r57}{por(j)} + \frac{1.6 (1.0 - por(j)) r56}{por(j)} - 0.25 r58 - 0.33 r59 \\
dBSidt &:= -r48 \\
dDSidt &:= \frac{r48 (1.0 - por(j))}{por(j)} - \frac{r49}{por(j)} + \frac{(1.0 - por(j)) r56}{por(j)} \\
&+ \frac{(1.0 - por(j)) r54}{por(j)} + \frac{2.3 (1.0 - por(j)) r55}{por(j)} - 3.5 r58 \\
&+ \frac{2.0 (1.0 - por(j)) r57}{por(j)} - 4.0 r59 - 2.0 r60 \\
dsfSidt &:= \frac{r49}{1.0 - por(j)} \\
daladt &:= \frac{0.36 (1.0 - por(j)) r54}{por(j)} - 2.3 r58 + \frac{0.0025 r48 (1.0 - por(j))}{por(j)} - 1.7 r59 \\
&- 2.0 r60 + r50 + \frac{1.7 (1.0 - por(j)) r55}{por(j)} \\
dalohdt &:= -r50 + r51 \\
daloh2dt &:= -r51 + r52 \\
daloh3dt &:= -r52 + r53 \\
daloh4dt &:= -r53 \\
dglassdt &:= -r54 \\
dplagiodt &:= -r55 \\
dolivedt &:= -r56 \\
dpyroxdt &:= -r57 \\
dillitedt &:= \frac{r58 por(j)}{1.0 - por(j)} \\
dsmectdt &:= \frac{r59 por(j)}{1.0 - por(j)} \\
dkaolidt &:= \frac{r60 por(j)}{1.0 - por(j)} \\
dnadt &:= \frac{0.7 (1.0 - por(j)) r55}{por(j)} \\
dkplusdt &:= -0.6 r58
\end{aligned}$$

(6.2)

Biogeochemistry – Equilibria

Specification of equilibrium constraints

eqrXnsId : set of kinetic reactions which are overruled by a thermodynamic constraint

equilibriumseqns[i] : Equilibrium constraint for reaction i

example:

```
eqrxnID := [r2,rX];
equilibriumeqns[1] := hplus*co3 - Keq*hco3;
equilibriumeqns[2] := ...;
> eqrxnId := [r29, r30, r31, r32, r33, r34, r35, r36,
r49, r50, r51, r52, r53] ;
> equilibriumeqns[1] := hplus*hco3-keq1*h2co3 ;
equilibriumeqns[2] := hplus*co3-keq2*hco3 ;
equilibriumeqns[3] := hplus*hs-keq3*h2s ;
equilibriumeqns[4] := hplus*boh4-keq4*boh3;
equilibriumeqns[5] := kspo4*po4*por(j) - spo4*(1. -
por(j)) ;
equilibriumeqns[6] := ksnh4*nh4*por(j) - snh4*(1. -
por(j)) ;
equilibriumeqns[7] := ksfp*(feoh3+feoh3mr+feoh3pr)*po4
- sfp ; #(assume forward reaction is dictated by solid
phase)#FSF Iron fractions;
equilibriumeqns[8] := ksfe*fe*por(j) - sfe*(1. - por
(j)) ;
equilibriumeqns[9] := ksfsi*(feoh3+feoh3mr+feoh3pr)*
(DSi) - (sfSi) ; #(assume forward reaction is dictated
by solid phase);
equilibriumeqns[10]:= aloh*hplus - eq1_al*al;
equilibriumeqns[11]:= aloh2*hplus - eq2_al*aloh;
equilibriumeqns[12]:= aloh3*hplus - eq3_al*aloh2;
equilibriumeqns[13]:= aloh4*hplus - eq4_al*aloh3;
```

```
eqrxnId := [r29, r30, r31, r32, r33, r34, r35, r36, r49, r50, r51, r52, r53]
equilibriumeqns1 := hplus hco3 - keq1 h2co3
equilibriumeqns2 := hplus co3 - keq2 hco3
equilibriumeqns3 := hplus hs - keq3 h2s
equilibriumeqns4 := hplus boh4 - keq4 boh3
equilibriumeqns5 := kspo4 po4 por(j) - spo4 (1. - por(j))
equilibriumeqns6 := ksnh4 nh4 por(j) - snh4 (1. - por(j))
equilibriumeqns7 := ksfp (feoh3 + feoh3mr + feoh3pr) po4 - sfp
equilibriumeqns8 := ksfe fe por(j) - sfe (1. - por(j))
equilibriumeqns9 := ksfsi (feoh3 + feoh3mr + feoh3pr) DSi - sfSi
equilibriumeqns10 := aloh hplus - eq1_al al
equilibriumeqns11 := aloh2 hplus - eq2_al aloh
equilibriumeqns12 := aloh3 hplus - eq3_al aloh2
equilibriumeqns13 := aloh4 hplus - eq4_al aloh3
```

(7.1)

Biogeochemistry - Parameters

Values of rates constants and parameters

In this section, all parameters defined in section 'Rate laws' should

be defined.

nparam: number of parameters to define

The list is given by **bio_name**; the values collected in **bio_val**.
note that for double precision, 10 should be written as 10.

example:

```
nparam:=4;
bio_name:=[kmo2hs,kf,kb,Keq];
vkf:=1.0*10^(5);
vKeq:=1.0*10^(-10.4);
vkb:=vkf*vKeq;
bio_val:=[1000.,vkf,vkb,vKeq];
```

```
> nparam := 139;
```

```
nparam := 139
```

(8.1)

```
> # calculate the dissociation constants, etc, based on
temp and salinity (baseline P = 0) from Millero (1995)
lnKC1_0 := 2.83655 - 2307.1266/ T - 1.5529413* ln(T) +
(- (0.20760841 + 4.0484/ T)* sqrt(S)) + (0.08468345* S
- 0.00654208* S* sqrt(S)) + (ln(1 - 0.001005* S));
lnKC2_0 := -9.226508 - 3351.6106/ T - 0.2005743* ln(T)+
(-0.106901773 - 23.9722/ T)* sqrt(S)+(0.1130822* S -
0.00846934* S^1.5 + ln(1 - 0.001005 * S));
lnKS_0 := 225.838 - 13275.3/T - 34.6435*ln(T) +
0.3449*S^0.5 - 0.0274*S ;
lnKB_0:=(-8966.90-2890.53*sqrt(S)-77.942*S+1.728*S^
(3.0/2.0)-0.0996*S*S)/T+(148.0248+137.1942*sqrt(S)
+1.62142*S)+((-24.4344-25.085*sqrt(S)-0.2474*S)*ln(T))
+ (0.053105*sqrt(S)*T);
```

```
# Now apply conversion for P (Millero 1995).
```

```
# units are M
```

```
deltav:= -25.5 + 0.1271*T_C + 0.0*T_C*T_C;
deltak:= (-3.08e-3 + 0.0877e-3*T_C + 0.0*T_C*T_C);
lnkpok0:= -(deltav/(R*T))*P + (0.5*deltak/(R*T))*P*P;
val_kkeq1 := exp(lnKC1_0)*exp(lnkpok0);
```

```
deltav:= -15.82 -0.0219*T_C;
deltak:= (1.13e-3 -0.1475e-3*T_C);
lnkpok0:= -(deltav/(R*T))*P + (0.5*deltak/(R*T))*P*P;
val_kkeq2:= exp(lnKC2_0)*exp(lnkpok0);
```

```
val_kkeq3 := exp(lnKS_0 + ((14.8 - 0.002*T_C + 0.0004
*T_C^2)*P + 0.5* (2.89 + 0.054*T_C) *P^2/1.01345)
/R/T);
```

```
deltav:= -29.48 + 0.1622*T_C -2.608e-3*T_C*T_C;
deltak:= (-2.84e-3+ 0.0794e-3*T_C);
```

```

lnkpok0:= -(deltav/(R*T))*P + (0.5*deltak/(R*T))*P*P;
val_kkeq4:= exp(lnKB_0)*exp(lnkpok0);

lnKC3_0:= -13847.26/T + 148.96502 - 23.6521* log(T)+
(118.67/T - 5.977 + 1.0495*log(T))*sqrt(S) - 0.01615*
S;
> deltav:= -25.6 +0.2324*T_C-3.6246e-3*T_C**2;
> deltak:= (5.13e-3 +0.07945e-3*T_C);
> lnkpok0:= -(deltav/(R*T))*P + (0.5*deltak/(R*T))*P*P;
> val_kkeq5:= exp(lnKC3_0)*exp(lnkpok0);
>

# calculate the solubility product for caco3
readlib(log10) :
tmp1 := -171.9065-0.077993*T+2839.319/T+71.595*log10
(T);
tmp2 := +(-0.77712+0.0028426*T+178.34/T)*sqrt(S);
tmp3 := -0.07711*S+0.0041249*S^1.5;
log10Kspc := tmp1 + tmp2 + tmp3;
ksca := 10.0^(log10Kspc);

deltav:= -48.7600 + 0.5304*T_C;
deltak:= (-0.0118 + 3.6920e-4*T_C);
lnkpok0:= -(deltav/(R*T))*P + (0.5*deltak/(R*T))*P*P;
val_kksca:= ksca * exp(lnkpok0);

# calculate the solubility product for aragonite
readlib(log10) :
tmp4 := -171.945-0.077993*T+2903.293/T+71.595*log10(T)
;
tmp5 := +(-0.068393+0.0017276*T+88.135/T)*sqrt(S);
tmp6 := -0.10018*S+0.0059415*S^1.5;
log10Kspa := tmp4 + tmp5 + tmp6;
ksar := 10.0^(log10Kspa);

deltav:= -35.0 + 0.5304*T_C;
deltak:= (-0.0118 + 3.6920e-4*T_C);
lnkpok0:= -(deltav/(R*T))*P + (0.5*deltak/(R*T))*P*P;
val_kksar:= ksar * exp(lnkpok0);

# calculate the solubility product for ikaite
readlib(log10) :
LogKspi := 0.15981 - 2011.1/T ; #Bischoff et al., 1993
ksik := 10**LogKspi ;
gammaCO3:= 0.0508; #CO3 activity coefficient (T/S)
from Pierot & Millero, 2017
gammaCa2:= 0.2053; #Ca2+ activity coefficient (T/S)
from Pierot & Millero, 2017

```

```

val_kksik:= ksik/(gammaCO3*gammaCa2);

# aluminium equilibrium constants (logK)
# LogK @ 5oC SUPERCRIPT database
# LogK' corrected to 10 oC / sal 35 (Pierrot and
Millero, 2017)
# unit correction: (mol/L) to (mol/cm3) applied
accordingly
LogK1_al := -5.297;
LogK2_al := -12.438;
LogK3_al := -19.012;
LogK4_al := -24.584;

kk1_al := (10**LogK1_al)*1.0e-3; # **mol/cm-3
conversion**
kk2_al := (10**LogK2_al)*1.0e-3; # **mol/cm-3
conversion**
kk3_al := (10**LogK3_al)*1.0e-3; # **mol/cm-3
conversion**
kk4_al := (10**LogK4_al)*1.0e-3; # **mol/cm-3
conversion**

# silicate minerals solubility constants
# Ksp values multiplied by a factor to convert to ***
mol cm-3***

## Basaltic glass: (0.36*Al + H4SiO4) ### from
Gislason and Oelkers, 2003 (SiO2 + 0.36Gibbsite = SiAl
(0.36)O2(OH(1.08))
## Ksp @ 10 oC SUPERCRIPT
## corrected to 10 oC / sal 35 (Pierrot and Millero,
2017)
LogKspglass:= 0.756 ;
Kspglass:= (10**LogKspglass)*0.144544; # ** conversion
(mol/L)^n to (mol/cm3)^n **

## Plagioclase - An70Ab30: (0.7*Ca + 0.3*Na + 1.7*
AlOH4 + 2.3*H4SiO4)
## Ksp @ 5oC after Stefansson 2001
## corrected to K'sp @ 5 oC / sal 35 (Pierrot and
Millero, 2017)
LogKspplagio := -18.248;
Kspplagio := (10**LogKspplagio)*1.0e-15 ; # **
conversion (mol/L)^n to (mol/cm3)^n **

## Olivine - Fo80Fa20 (1.6*Mg + 0.4*Fe + H4SiO4)
## Ksp @ 10oC after Stefansson 2001

```

```

## corrected to K'sp @ 10 oC / sal 35 (Pierrot and
Millero, 2017)
Log_Ksolive := 29.317 ;
Kspolive := (10**Log_Ksolive)*1.0e3 ; # ** conversion
(mol/L)^n to (mol/cm3)^n **

## Clinopyroxene - Di45Hed25En19Fs11:
## Ksp @ 5oC after Stefansson 2001
## corrected K'sp to 5 oC / sal 35 (Pierrot and
Millero, 2017)
LogKspyrox := 24.916 ;
Kspprox := (10**LogKspyrox)*1.0; # ** conversion
(mol/L)^n to (mol/cm3)^n **

##Illite ... ### 0.6K+ + 0.25Mg2+ + 2.3Al3+
+3.5H4SiO4!!!
## Ksp @ 10 oC from SUPERCRIPT database
## corrected K'sp to 10 oC / sal 35 (Pierrot and
Millero, 2017)
LogKsillite:= 13.044 ;
Kspillite:= (10**LogKsillite)*1.122*1.0e4; # **
conversion (mol/L)^n to (mol/cm3)^n **

##Smectite # Ca-Montmorillonite
##Ksp @ 5oC from SUPERCRIPT database
##corrected K'sp to 5oC / sal 35 (Pierrot and Millero,
2017)
LogKssmect:= 6.244 ;
Kpsmect:= (10**LogKssmect)*2.6002e-1; #** conversion
(mol/L) to (mol/cm3) **

##Kaolinite #Al2Si2O5(OH)4
##Ksp @ 5oC from SUPERCRIPT database
##Corrected K'sp to 5oC / sal 35 (Pierrot and Millero,
2017)
LogKskaoli:= 8.176 ;
Kspkaoli:= (10**LogKskaoli)* 1.0e6; #**conversion
(mol/L) to (mol/cm3)**

```

>

#

```

lnKCl_0 := -13.81164725
lnKC2_0 := -21.14789969
lnKS_0 := -15.54834188
lnKB_0 := -20.20576891

```



```

    deltav := -24.19087
    deltak := -0.00217669
    lnkpok0 := 0.01033668884
    val_kkeq1 := 0.000001014301296
    deltav := -16.04557
    deltak := -0.00038925
    lnkpok0 := 0.006858496573
    val_kkeq2 := 6.585101419 10-10
    val_kkeq3 := 1.792144094 10-7
    deltav := -28.08602272
    deltak := -0.00202218
    lnkpok0 := 0.01200216383
    val_kkeq4 := 1.698081511 10-9
    lnKC3_0 := -31.83859080
    deltav := -23.59081381
    deltak := 0.005948335
    lnkpok0 := 0.01009766720
    val_kkeq5 := 1.503353334 10-14
    tmp1 := -8.4115624
    tmp2 := 3.891545170
    tmp3 := -1.844736688
    log10Kspc := -6.364753918
    ksca := 4.317636555 10-7
    deltav := -43.29688
    deltak := -0.007997240
    lnkpok0 := 0.01849179953
    val_kksca := 4.398220198 10-7
    tmp4 := -8.2243647
    tmp5 := 4.331942300
    tmp6 := -2.276036419
    log10Kspa := -6.168458819
    ksar := 6.784864534 10-7
    deltav := -29.53688
    deltak := -0.007997240
    lnkpok0 := 0.01260952505
    val_kksar := 6.870960123 10-7
    LogKspi := -6.935268498
    ksik := 1.160730782 10-7
    gammaCO3 := 0.0508
    gammaCa2 := 0.2053
    val_kksik := 0.00001112958166
    LogK1_al := -5.297
    LogK2_al := -12.438
    LogK3_al := -19.012
    LogK4_al := -24.584
    kk1_al := 5.046612976 10-9

```

$kk2_al := 3.647539469 \cdot 10^{-16}$
 $kk3_al := 9.727472238 \cdot 10^{-23}$
 $kk4_al := 2.606153550 \cdot 10^{-28}$
 $LogKspglass := 0.756$
 $Kspglass := 0.8241382458$
 $LogKspplagio := -18.248$
 $Kspplagio := 5.649369748 \cdot 10^{-34}$
 $Log_Ksolive := 29.317$
 $Kspolive := 2.074913517 \cdot 10^{32}$
 $LogKspyrox := 24.916$
 $Ksppyrox := 8.241381150 \cdot 10^{24}$
 $LogKsillite := 13.044$
 $Kspillite := 1.241631886 \cdot 10^{17}$
 $LogKssmect := 6.244$
 $Kpsmect := 4.560440081 \cdot 10^5$
 $LogKskaoli := 8.176$
 $Kspkaoli := 1.499684836 \cdot 10^{14}$

(8.2)

```

> #units mol, l, yr
> val_xbiot      :=10.0;
val_inita       :=SWI_a;
val_nu          :=0.1;
val_kmo2        :=8.0e-9;
val_kmno3       :=5.0e-9;
val_kmmno2      :=5.0e-6;
val_kmfeoh3     :=1.25e-5;
val_kmso4       :=100.0e-9;
val_knit        :=1.0e7;
val_kmnox       :=5.0e7;
val_kfemno2     :=3.0e6;
val_kfemno2pr   :=3.0e2; #FSF
val_kfeo2       :=5.0e9;
val_kh2so2      :=1.0e7;
val_kh2smno2    :=1.0e2;
val_kh2smno2pr  :=1.0e-1; #FSF
val_kh2sfeoh3   :=1.0e2;
val_kh2sfeoh3mr:=1.0e-2; #FSF
val_kh2sfeoh3pr:=1.0e-4; #FSF
val_kaom        :=5.0e6;
val_kaomo2      :=1.0e13;
val_kfeso2      :=1.0e9;
val_kmnco3precip:=0.1e-7;
val_KsMnCO3     :=3.2e-15;
val_kfesprecip  :=5.0e-9;
val_kfesdiss    :=1.0e-3;
val_KsFeS       :=6.3e-6/0.77;
val_kfeco3precip:=1.0e-9;

```

```

val_KsFeCO3      :=4.0e-15;
val_kcaldiss     :=0.1e0;
val_kcalppt      :=0.1e-1; #FSF
val_karadis      :=0.1e0;
val_karappt      :=0.1e-1;
val_kmgcdiss     :=0.1e0; ## Ikaite
val_kmgcppt      :=0.1e-1; ## Ikaite
val_kpyr         :=60.0e6;
val_kfess0       :=2.5e4;
val_kapa         :=1.0e0;
val_po4_eq       :=10.0e-9;
val_kdis         :=0.0e0;
val_ch4eq        :=0.1e5;
val_kgas         :=0.0e0;
val_kdi          :=0.1e-2;
val_h2sstar      :=0.1e-4;
val_ksnh4        :=0.16e1;
val_kspo4        :=0.18e1;
val_ksfp         :=100.0;
val_ksfe         :=400.0;
val_kmnage       :=0.6; #FSF
val_kfeage       :=0.6; #FSF

val_keq1 := val_kkeq1*1e-3;
val_keq2 := val_kkeq2*1e-3;
val_keq3 := val_kkeq3*1e-3;
val_keq4 := val_kkeq4*1e-3;
val_keq5 := val_kkeq5*1e-6; #cst ionisation water Kw=
10-14mol/l; 10-20mol/cm3

val_kf1 := 1.0;
val_kb1 := val_keq1;
val_kf2 := 1.0 ;
val_kb2 := val_keq2;
val_kf3 := 1.0;
val_kb3 := val_keq3;
val_kf4 := 1.0;
val_kb4 := val_keq4;
val_kf5 := val_kspo4;
val_kb5 := 1.0;
val_kf6 := val_ksnh4;
val_kb6 := 1.0;
val_kf7 := val_ksfp;
val_kb7 := 1.0;
val_kf8 := val_ksfe;
val_kb8 := 1.0;
val_kspcal := val_kksca*1e-6;

```

```
val_kspara := val_kksar*1e-6;  
val_kspmgc := val_kspara*1.21; #Ikaite ##for MgCal,  
val_kspmgc := val_kspara*1.21;
```

```
val_kBSidis := 0.1e-2;  
val_ad := 0.2;  
val_bd := 0.05;  
val_BSisat := 8.0e-7;
```

```
val_ksfsi:= 0.0 ; #0.5e7;  
val_kf9 := val_ksfsi;  
val_kb9 := 1.0;
```

```
val_eq1_al:= kk1_al;  
val_eq2_al:= kk2_al;  
val_eq3_al:= kk3_al;  
val_eq4_al:= kk4_al;  
val_kf10 := 1.0;  
val_kb10 := val_eq1_al;  
val_kf11 := 1.0;  
val_kb11 := val_eq2_al;  
val_kf12 := 1.0;  
val_kb12 := val_eq3_al;  
val_kf13 := 1.0;  
val_kb13 := val_eq4_al;
```

```
val_kspglass := Kspglass;  
val_kglass:= 0.0; #0.2e1;
```

```
val_kspplagio:= Kspplagio;  
val_kplagio := 0.0 ; #0.1e0;
```

```
val_kspolive:= Kspolive;  
val_kolive := 0.0; #0.15e-3;
```

```
val_ksppyrox:= Ksppyrox;  
val_kpyrox := 0.0 ; #0.1e-6;
```

```
val_kspillite:= Kspillite;  
val_killite := 0.0; #0.2e-9;
```

```
val_kspsmect:= Kspsmect;  
val_ksmect:= 0.0 ; #0.1e-15;
```

```
val_kspkaoli:= Kspkaoli;  
val_kkaoli:= 0.0 ; #0.1e-15;
```

```
val_xbiot := 10.0  
val_inita := 10.
```

```

    val_nu := 0.1
    val_kmo2 := 8.0 10-9
    val_kmno3 := 5.0 10-9
    val_kmmno2 := 0.0000050
    val_kmfeoh3 := 0.0000125
    val_kms04 := 1.000 10-7
    val_knit := 1.0 107
    val_kmnox := 5.0 107
    val_kfemno2 := 3.0 106
    val_kfemno2pr := 300.
    val_kfeo2 := 5.0 109
    val_kh2so2 := 1.0 107
    val_kh2smno2 := 100.
    val_kh2smno2pr := 0.10
    val_kh2sfeoh3 := 100.
    val_kh2sfeoh3mr := 0.010
    val_kh2sfeoh3pr := 0.00010
    val_kaom := 5.0 106
    val_kaomo2 := 1.0 1013
    val_kfeso2 := 1.0 109
    val_kmnco3precip := 1. 10-8
    val_KsMnCO3 := 3.2 10-15
    val_kfesprecip := 5.0 10-9
    val_kfesdiss := 0.0010
    val_KsFeS := 0.000008181818182
    val_kfeco3precip := 1.0 10-9
    val_KsFeCO3 := 4.0 10-15
    val_kcaldiss := 0.1
    val_kcalppt := 0.01
    val_karadis := 0.1
    val_karappt := 0.01
    val_kmgcdiss := 0.1
    val_kmgcppt := 0.01
    val_kpyr := 6.00 107
    val_kfess0 := 25000.
    val_kapa := 1.0
    val_po4_eq := 1.00 10-8
    val_kdis := 0.
    val_ch4eq := 10000.
    val_kgas := 0.
    val_kdi := 0.001
    val_h2sstar := 0.00001
    val_ksnh4 := 1.6
    val_kspo4 := 1.8

```

```

    val_ksfp := 100.0
    val_ksfe := 400.0
    val_kmnage := 0.6
    val_kfeage := 0.6
    val_keq1 := 1.014301296 10-9
    val_keq2 := 6.585101419 10-13
    val_keq3 := 1.792144094 10-10
    val_keq4 := 1.698081511 10-12
    val_keq5 := 1.503353334 10-20
    val_kf1 := 1.0
    val_kb1 := 1.014301296 10-9
    val_kf2 := 1.0
    val_kb2 := 6.585101419 10-13
    val_kf3 := 1.0
    val_kb3 := 1.792144094 10-10
    val_kf4 := 1.0
    val_kb4 := 1.698081511 10-12
    val_kf5 := 1.8
    val_kb5 := 1.0
    val_kf6 := 1.6
    val_kb6 := 1.0
    val_kf7 := 100.0
    val_kb7 := 1.0
    val_kf8 := 400.0
    val_kb8 := 1.0
    val_kspcal := 4.398220198 10-13
    val_kspara := 6.870960123 10-13
    val_kspmge := 8.313861749 10-13
    val_kBSidis := 0.001
    val_ad := 0.2
    val_bd := 0.05
    val_BSisat := 8.0 10-7
    val_ksfsi := 0.
    val_kf9 := 0.
    val_kb9 := 1.0
    val_eq1_al := 5.046612976 10-9
    val_eq2_al := 3.647539469 10-16
    val_eq3_al := 9.727472238 10-23
    val_eq4_al := 2.606153550 10-28
    val_kf10 := 1.0
    val_kb10 := 5.046612976 10-9
    val_kf11 := 1.0
    val_kb11 := 3.647539469 10-16
    val_kf12 := 1.0

```

```

val_kb12 := 9.727472238 10-23
val_kf13 := 1.0
val_kb13 := 2.606153550 10-28
val_kspglass := 0.8241382458
val_kglass := 0.
val_kspplagio := 5.649369748 10-34
val_kplagio := 0.
val_kspolive := 2.074913517 1032
val_kolive := 0.
val_ksppyrox := 8.241381150 1024
val_kpyrox := 0.
val_kspillite := 1.241631886 1017
val_killite := 0.
val_kspsmect := 4.560440081 105
val_ksmect := 0.
val_kspkaoli := 1.499684836 1014
val_kkaoli := 0.

```

(8.3)

```

> bio_name := [sw01, sw02, sw03, sw04, sw05, sw06, sw07,
sw08, sw09, sw10, sw11, sw12, sw13, sw14, sw15, sw16,
sw17, sw18, sw19, sw20, sw21, sw22, sw23, sw24, sw25,
sw26, sw27, sw28, sw29, sw30, sw31, sw32, xbiot,
inita, nu, kmo2, kmno3, kmmno2, kmfeoh3, kmso4, knit,
kmnox, kfemno2, kfeo2, kh2so2, kh2smno2, kh2sfeoh3,
kaom, kaomo2, kfeso2, kmnco3precip, KsMnCO3,
kfesprecip, kfesdiss, KsFeS, kfeco3precip, KsFeCO3,
kcaldiss, kcalppt, karadis, karappt, kmgcds, kmgcppt,
kpyr, kfess0, kapa, po4_eq, kdis, h2sstar, ch4eq,
kgas, kdi, ksnh4, kspo4, ksfp, ksfe, kf1, kb1, kf2,
kb2, kf3, kb3, kf4, kb4, kf5, kb5, kf6, kb6, kf7, kb7,
kf8, kb8, kspcal, kspara, kspmgc, keq1, keq2, keq3,
keq4, kfemno2pr, kh2smno2pr, kmnage, kh2sfeoh3mr,
kh2sfeoh3pr, kfeage, keq5, kBSidis, ad, bd, BSisat,
ksfsi, kf9, kb9, eq1_al, eq2_al, eq3_al, eq4_al, kf10,
kb10, kf11, kb11, kf12, kb12, kf13, kb13, kspglass,
kglass, kspplagio, kplagio, kspolive, kolive,
ksppyrox, kpyrox, kspillite, killite, kspsmect,
ksmect, kspkaoli, kkaoli];

```

```

bio_name := [sw01, sw02, sw03, sw04, sw05, sw06, sw07, sw08, sw09, sw10, sw11, sw12,
sw13, sw14, sw15, sw16, sw17, sw18, sw19, sw20, sw21, sw22, sw23, sw24, sw25,
sw26, sw27, sw28, sw29, sw30, sw31, sw32, xbiot, inita, v, kmo2, kmno3, kmmno2,
kmfeoh3, kmso4, knit, kmnox, kfemno2, kfeo2, kh2so2, kh2smno2, kh2sfeoh3, kaom,
kaomo2, kfeso2, kmnco3precip, KsMnCO3, kfesprecip, kfesdiss, KsFeS, kfeco3precip,
KsFeCO3, kcaldiss, kcalppt, karadis, karappt, kmgcds, kmgcppt, kpyr, kfess0, kapa,
po4_eq, kdis, h2sstar, ch4eq, kgas, kdi, ksnh4, kspo4, ksfp, ksfe, kf1, kb1, kf2, kb2, kf3,
kb3, kf4, kb4, kf5, kb5, kf6, kb6, kf7, kb7, kf8, kb8, kspcal, kspara, kspmgc, keq1, keq2,

```

(8.4)

```
> bio_val :=[0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0,
0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0,
0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0,
0.0, 0.0, val_xbiot, val_inita, val_nu, val_kmo2,
val_kmno3, val_kmmno2, val_kmfeoh3, val_kmsO4,
val_knit, val_kmnox, val_kfemno2, val_kfeo2,
val_kh2so2, val_kh2smno2, val_kh2sfeoh3, val_kaom,
val_kaomo2, val_kfeso2, val_kmnco3precip, val_KSMnCO3,
val_kfesprecip, val_kfesdiss, val_KsFeS,
val_kfeco3precip, val_KsFeCO3, val_kcaldiss,
val_kcalppt, val_karadis, val_karappt, val_kmgcdiss,
val_kmgcppt, val_kpyr, val_kfess0, val_kapa,
val_po4_eq, val_kdis, val_h2sstar, val_ch4eq,
val_kgas, val_kdi, val_ksnh4, val_kspo4, val_ksfp,
val_ksfe, val_kf1, val_kb1, val_kf2, val_kb2, val_kf3,
val_kb3, val_kf4, val_kb4, val_kf5, val_kb5, val_kf6,
val_kb6, val_kf7, val_kb7, val_kf8, val_kb8,
val_kspcal, val_kspara, val_kspmgc, val_keq1,
val_keq2, val_keq3, val_keq4, val_kfemno2pr,
val_kh2smno2pr, val_kmnage, val_kh2sfeoh3mr,
val_kh2sfeoh3pr, val_kfeage, val_keq5, val_kBSidis,
val_ad, val_bd, val_BSisat, val_ksfsi, val_kf9,
val_kb9, val_eq1_al, val_eq2_al, val_eq3_al,
val_eq4_al, val_kf10, val_kb10, val_kf11, val_kb11,
val_kf12, val_kb12, val_kf13, val_kb13, val_kspglass,
val_kglass, val_kspplagio, val_kplagio, val_kspolive,
val_kolive, val_ksppyrox, val_kpyrox, val_kspillite,
val_killite, val_kspsmect, val_ksmect, val_kspkaoli,
val_kkaoli];
```

$$\text{bio_val} := [0., \\ 0., 0., 0., 0., 0., 0., 10.0, 10., 0.1, 8.0 \cdot 10^{-9}, 5.0 \cdot 10^{-9}, 0.0000050, 0.0000125, 1.000 \cdot 10^{-7}, \\ 1.0 \cdot 10^7, 5.0 \cdot 10^7, 3.0 \cdot 10^6, 5.0 \cdot 10^9, 1.0 \cdot 10^7, 100., 100., 5.0 \cdot 10^6, 1.0 \cdot 10^{13}, 1.0 \cdot 10^9, 1. \cdot 10^{-8}, \\ 3.2 \cdot 10^{-15}, 5.0 \cdot 10^{-9}, 0.0010, 0.000008181818182, 1.0 \cdot 10^{-9}, 4.0 \cdot 10^{-15}, 0.1, 0.01, 0.1, 0.01, \\ 0.1, 0.01, 6.00 \cdot 10^7, 25000., 1.0, 1.00 \cdot 10^{-8}, 0., 0.00001, 10000., 0., 0.001, 1.6, 1.8, 100.0, \\ 400.0, 1.0, 1.014301296 \cdot 10^{-9}, 1.0, 6.585101419 \cdot 10^{-13}, 1.0, 1.792144094 \cdot 10^{-10}, 1.0, \\ 1.698081511 \cdot 10^{-12}, 1.8, 1.0, 1.6, 1.0, 100.0, 1.0, 400.0, 1.0, 4.398220198 \cdot 10^{-13}, \\ 6.870960123 \cdot 10^{-13}, 8.313861749 \cdot 10^{-13}, 1.014301296 \cdot 10^{-9}, 6.585101419 \cdot 10^{-13}, \\ 1.792144094 \cdot 10^{-10}, 1.698081511 \cdot 10^{-12}, 300., 0.10, 0.6, 0.010, 0.00010, 0.6.] \quad (8.5)$$

1.503353334 10^{-20} , 0.001, 0.2, 0.05, $8.0 \cdot 10^{-7}$, 0., 0., 1.0, $5.046612976 \cdot 10^{-9}$,
 3.647539469 10^{-16} , $9.727472238 \cdot 10^{-23}$, $2.606153550 \cdot 10^{-28}$, 1.0, $5.046612976 \cdot 10^{-9}$, 1.0,
 3.647539469 10^{-16} , 1.0, $9.727472238 \cdot 10^{-23}$, 1.0, $2.606153550 \cdot 10^{-28}$, 0.8241382458, 0.,
 $5.649369748 \cdot 10^{-34}$, 0., $2.074913517 \cdot 10^{32}$, 0., $8.241381150 \cdot 10^{24}$, 0., $1.241631886 \cdot 10^{17}$,
 0., $4.560440081 \cdot 10^5$, 0., $1.499684836 \cdot 10^{14}$, 0.]

Switches

Switches can be used in the rate equations. Specify in **nswitches**, how many switches are in use, name them and define the switch expressions. The switch names must also appear in **bio_name** and must be assigned a dummy value there. The switch equals 1 if the switch expression is >0 , 0 otherwise. To reference the coordinates in the domain, use x_pos , y_pos and z_pos .

```
> nswitches := 32;
                                     nswitches := 32 (8.1.1)
```

```
> switchlist := [sw01, sw02, sw03, sw04, sw05, sw06,
                 sw07, sw08, sw09, sw10, sw11, sw12, sw13, sw14,
                 sw15, sw16, sw17, sw18, sw19, sw20, sw21, sw22,
                 sw23, sw24, sw25, sw26, sw27, sw28, sw29, sw30,
                 sw31, sw32] ;
switchlist := [sw01, sw02, sw03, sw04, sw05, sw06, sw07, sw08, sw09, sw10, sw11,
               sw12, sw13, sw14, sw15, sw16, sw17, sw18, sw19, sw20, sw21, sw22, sw23, sw24,
               sw25, sw26, sw27, sw28, sw29, sw30, sw31, sw32] (8.1.2)
```

```
> switchcrit := [(xbiot-x_pos), (o2-kmo2), -(sw02
               -1.0), sw03*(no3-kmno3), -sw03*(sw04-1.0), sw05*
               (mno2-kmmno2), -sw05*(sw06-1.0), sw07*(feoh3-
               kmfeoh3), -sw07*(sw08-1.0), sw09*(so4-kmso4), -sw09*
               (sw10-1.0), ((mn*co3/KsMnCO3)-1.0), ((fe*hs)/(KsFeS*
               hplus)-1.0), ((fe*co3/KsFeCO3)-1.0), (1.0-(ca*
               co3/kspcal)), (po4-po4_eq), (ch4-ch4eq), (ch4-0.0),
               (h2ssstar-(h2s+hs)), ((ca*co3/kspcal)-5.0), (1.0-(ca*
               co3/kspara)), ((ca*co3/kspara)-5.0), (1.0-(ca*
               co3/kspmgc)), ((ca*co3/kspmgc)-1.0), -(DSi-BSisat),
               (1.0 - DSi*al**0.36/hplus**1.08/kspglass), (1.0 -
               DSi**2.3*al**1.7*na**0.7*ca**0.3/kspplagio), (1.0 -
               DSi*mg**1.6*fe**0.4/hplus**4/kspolive), (1.0 - DSi**
               2 * fe**0.46 * mg**0.84 * ca**0.7 / hplus**4 /
               ksppyrox), (DSi**3.5 * al**2.3 * kplus**0.6 * mg**
               0.25 / hplus**8 / kspillite - 1.0), (ca**0.165 * mg*
               *0.33 * al**1.7 * DSi**4 / hplus**6 / kspsmect -
               1.0), (al**2 * DSi**2 / hplus**6 / kspkaoli - 1.0)];
switchcrit := [xbiot - x_pos, o2 - kmo2, 1.0 - sw02, sw03 (no3 - kmno3),
               -sw03 (sw04 - 1.0), sw05 (mno2 - kmmno2), -sw05 (sw06 - 1.0), sw07 (feoh3
```

-sw03 (sw04 - 1.0), sw05 (mno2 - kmmno2), -sw05 (sw06 - 1.0), sw07 (feoh3

$$\begin{aligned}
& - kmfeoh3), -sw07 (sw08 - 1.0), sw09 (so4 - kmso4), -sw09 (sw10 - 1.0), \\
& \frac{mn\ co3}{KsMnCO3} - 1.0, \frac{fe\ hs}{hplus\ KsFeS} - 1.0, \frac{fe\ co3}{KsFeCO3} - 1.0, 1.0 - \frac{ca\ co3}{kspcal}, po4 \\
& - po4_eq, ch4 - ch4eq, ch4, h2sstar - h2s - hs, \frac{ca\ co3}{kspcal} - 5.0, 1.0 - \frac{ca\ co3}{kspara}, \\
& \frac{ca\ co3}{kspara} - 5.0, 1.0 - \frac{ca\ co3}{kspmgc}, \frac{ca\ co3}{kspmgc} - 1.0, -DSi + BSisat, 1.0 \\
& - \frac{DSi\ al^{0.36}}{hplus^{1.08}\ kspglass}, 1.0 - \frac{DSi^{2.3}\ al^{1.7}\ na^{0.7}\ ca^{0.3}}{kspplagio}, 1.0 - \frac{DSi\ mg^{1.6}\ fe^{0.4}}{hplus^4\ kspolive}, 1.0 \\
& - \frac{DSi^2\ fe^{0.46}\ mg^{0.84}\ ca^{0.7}}{hplus^4\ ksppyrox}, \frac{DSi^{3.5}\ al^{2.3}\ kplus^{0.6}\ mg^{0.25}}{hplus^8\ kspillite} - 1.0, \\
& \left[\frac{ca^{0.165}\ mg^{0.33}\ al^{1.7}\ DSi^4}{hplus^6\ kpsmect} - 1.0, \frac{al^2\ DSi^2}{hplus^6\ kspkaoli} - 1.0 \right]
\end{aligned}$$

Transport - Molecular Diffusion

Specification of the molecular diffusion coefficients

diffdata: molecular diffusion coefficient at 0 degree celsius
(cm²/yr)

alphadata: temperature dependence of the diffusion coefficient (1/K)

the in situ molecular diffusion coefficient, corrected for tortuosity is calculated as:

$D(T, sal) = [(0.95 - 0.001 * sal) * D(T=0, sal=0) * (1 + muc * T[C])] / (1 - \ln(por^2))$

example:

```
diffdata := [100., 304., 0., 0., 100.];
alphadata:= [0.006, 0.04, 0., 0., 0.05];
[ch2o, o2, no3, mno2, feoh3, so4, ch4, nh4, po4, mn, fe, h2s, hs,
ch4g, h2co3, hco3, co3, boh4, boh3, hplus, caco3, ca, snh4, spo4,
fes, feco3, s0, fes2, sfe, sfp, mnco3]
```

```
> diffdata := [0.0, 380.449545, 394.5878727, 0.0, 0.0,
173.9205889, 263.9351889, 395.8731752, 112.35777,
123.3890416, 136.2420668, 331.6080494, 392.0172677,
5000.0, 320.0403267, 217.2161254, 176.0864448,
96.29573485, 110.0522684, 600.0, 0.0, 150.3803945,
0.0, 0.0, 0.0, 0.0, 173.9205889, 0.0, 0.0, 0.0, 0.0,
0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 112.26816, 0.0, 125.0,
0.0, 87.9966, 87.9966, 140.6684, 140.6684, 140.6684,
0.0, 0.0, 197.73072, 310.94496, 0.0, 0.0, 0.0, 0.0,
0.0];
```

```
alphadata := [0.0, 0.06, 0.038, 0.0, 0.0, 0.045,
0.0520, 0.041, 0.054, 0.05, 0.044, 0.06, 0.031, 0.0,
```

```
0.06, 0.048, 0.047, 0.048, 0.048, 0.06, 0.0, 0.045,
0.0, 0.0, 0.0, 0.0, 0.045, 0.0, 0.0, 0.0, 0.0, 0.0,
0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0,
0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0,
0.0] ;
```

```
diffdata := [0., 380.449545, 394.5878727, 0., 0., 173.9205889, 263.9351889, 395.8731752,
112.35777, 123.3890416, 136.2420668, 331.6080494, 392.0172677, 5000.0,
320.0403267, 217.2161254, 176.0864448, 96.29573485, 110.0522684, 600.0, 0.,
150.3803945, 0., 0., 0., 173.9205889, 0., 0., 0., 0., 0., 0., 0., 112.26816, 0.,
125.0, 0., 87.9966, 87.9966, 140.6684, 140.6684, 140.6684, 0., 0., 197.73072,
310.94496, 0., 0., 0., 0.]
```

```
alphadata := [0., 0.06, 0.038, 0., 0., 0.045, 0.0520, 0.041, 0.054, 0.05, 0.044, 0.06, 0.031, 0., (9.1)
0.06, 0.048, 0.047, 0.048, 0.048, 0.06, 0., 0.045, 0., 0., 0., 0.045, 0., 0., 0., 0., 0.,
0., 0., 0., 0., 0., 0., 0., 0., 0., 0., 0., 0., 0., 0., 0., 0., 0., 0.]
```

Transport - Boundary Conditions

Specification of upper and lower boundary conditions for each species.

There are 3 options

0. known concentration (Dirichlet, M/L^3)

1. known concentration gradient (Neumann, $M/L^3 \cdot L$)

2. known total (diffusive and advective) flux (Robin, $M/L^2/T$)

technical note: option 1 and 2 involve ghost points outside the domain. If the mixing parameters vary with depth, one needs to assign a mixing intensities at the ghost points. By default this is done by linear extrapolation. To overwrite this, the user has to edit `gridsetup.f` and `advdiffcoeff.f` (both at the bottom; explanations are given there)

type up: array defining the type of condition for each species at the upper boundary (0, 1 or 2)

bnddata up: array containing the values specified at the upper boundary.

type down: array defining the type of condition for each species at the lower boundary

bnddata down: array containing the values specified at the upper boundary.

example:

```
type_up := [0,0,2];
```

```
bnddata_up := [1.4,0.001,0.001];
```

```
type_down := [1,1,1];
```

```
bnddata_down := [0.,0.,0.];
```

```
[ch2o, o2, no3, mno2, feoh3, so4, ch4, nh4, po4, mn, fe, h2s, hs, ch4g, h2co3, hco3, co3, boh4, boh3,
hplus, caco3, ca, snh4, spo4, fes, feco3, s0, fes2, sfe, sfp, mnco3] ;
```

```
> #set upper boundary cond TCup from given TAup and
hplusup
#by def: keq4=(hplus*BOH4)/BOH3; keq3=(hplus*HS)/H2S ;
and keq5=hplus*OHminus
#TAup and hplusup to be set by the user:
TAup :=2.306e-6; #mol/cm3
TBup := 0.416e-6;
```

```

TSup := 0.0;
hplusup := 10**(-SWI_pH)*1e-3; ## value in brackets is
'the actual pH-value #FSF

term1 := -(val_keq4*TBup)/(hplusup+val_keq4)-
(val_keq3*TSup)/(hplusup+val_keq3)-val_keq5/hplusup+
hplusup;
term2 := 1+val_keq1/hplusup+(val_keq1*val_keq2)/
(hplusup*hplusup);
term3 := val_keq1/hplusup+(2*val_keq1*val_keq2)/
(hplusup*hplusup);
TCup := (TAup+term1)*term2/term3;

co3up := TCup*val_keq1*val_keq2/(val_keq1*val_keq2+
val_keq1*hplusup+hplusup*hplusup);
hco3up := TCup*val_keq1*hplusup/(val_keq1*val_keq2+
val_keq1*hplusup+hplusup*hplusup);
h2co3up := TCup - co3up - hco3up;

boh4up := TBup*hplusup/(val_keq4+hplusup);
boh3up := TBup-boh4up;

ohup := val_keq5/hplusup;
caup := 10.0e-6;          #report manually the value
from bnddata_up
Omegaup := (co3up*caup)/val_kspcal;

## Set upper boundary conditions for Aluminium species
based on T_Al and pH ##
## Al eq constants LogK_Al defined in parameters above

T_Al_up:= 50.0e-12; ## Total dissolved aluminium
concentration at the SWI

## Al_DNT = common denominator for Al speciation

Al_up := (T_Al_up)/(1.0 + val_eq1_al/hplusup +
(val_eq1_al*val_eq2_al)/hplusup**2 +
(val_eq1_al*val_eq2_al*val_eq3_al)/hplusup**3 +
(val_eq1_al*val_eq2_al*val_eq3_al*val_eq4_al)/hplusup*
*4);

AlOH_up:= val_eq1_al/hplusup * Al_up;

AlOH2_up:= (val_eq1_al*val_eq2_al)/hplusup**2 * Al_up;

```

```
AlOH3_up:= (val_eq1_al*val_eq2_al*val_eq3_al)/hplusup*
*3 * Al_up;
```

```
AlOH4_up:= (val_eq1_al*val_eq2_al*val_eq3_al*
val_eq4_al)/hplusup**4 * Al_up;
```

```
type_up := [0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0,
0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0,
0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0,
0, 0, 0, 0, 0] ;
```

```
bnddata_up := [SWI_TOC, 195.0e-9, 17.3e-9, 1.36e-4,
2.21e-4, 28.0e-6, 0.0, 0.0, 1.0e-9, 0.0, 0.0, 0.0,
0.0, 0.0, h2co3up, hco3up, co3up, boh3up, boh4up,
hplusup, 1.78e-3, caup, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0,
0.0, 0.0, 0.0, 0.0, 1.36e-4, 2.21e-4, 2.21e-4,
9.53e-4, 0.0, 0.523e-4, 1.5e-03, 17.3e-9, 0.0, Al_up,
AlOH_up, AlOH2_up, AlOH3_up, AlOH4_up, 2.4e-3, 0.0,
469.1e-6, 10.2e-6, 0.85e-3, 0.0, 0.0, 0.0, 0.0] ;
```

```
type_down := [1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1,
1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1,
1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1,
1, 1, 1, 1, 1] ;
```

```
bnddata_down := [0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0,
0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0,
0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0,
0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0,
0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0,
0.0, 0.0, 0.0, 0.0] ;
```

$TA_{up} := 0.000002306$

$TB_{up} := 4.16 \cdot 10^{-7}$

$TS_{up} := 0.$

$hplus_{up} := 7.943282347 \cdot 10^{-12}$

$term1 := -7.515250686 \cdot 10^{-8}$

$term2 := 139.2789082$

$term3 := 148.8648487$

$TC_{up} := 0.000002087195237$

$co3_{up} := 1.586379795 \cdot 10^{-7}$

$hco3_{up} := 0.000001913571534$

$h2co3_{up} := 1.4985724 \cdot 10^{-8}$

$boh4_{up} := 3.427321595 \cdot 10^{-7}$

$boh3_{up} := 7.32678405 \cdot 10^{-8}$

- 3: individual files for each species. listinput contains the species number (length: number of species), filenames are given in "file_in_names".inp, containing in column 1 the conc, in column 2 depth.

example (only relevant input data is given for each option, assuming 17 species):

NOTE that you have to provide something for all ncomp species in the arrays iniconc, listinput and file_in_names, EVEN IF YOU DON'T USE IT WITH THE OPTION YOU SELECTED

vic:=1;

vic:=2;iniconc:=[0,2,...];

vic:=3;listinput:=[1,2,...,17];file_in_names:=[o2,no3,...,sp17];

```
> vic := 2 ;
iniconc := [SWI_TOC, 195.0e-9, 17.3e-9, 0.0, 0.0,
28.0e-6, 0.0, 0.0, 1.0e-9, 0.0, 0.0, 0.0, 0.0, 0.0,
h2co3up, hco3up, co3up, boh3up, boh4up, hplusup,
1.78e-3, caup, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0,
0.0, 0.0, 0.0, 0.0, 0.0, 9.53e-4, 0.0, 0.523e-4,
1.5e-03, 17.3e-9, 0.0, Al_up, AlOH_up, AlOH2_up,
AlOH3_up, AlOH4_up, 0.72e-3, 0.0, 469.1e-6, 10.2e-6,
0.17e-3, 0.0, 0.0, 0.0, 0.0] ;
```

```
listinput := [1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12,
13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25,
26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38,
39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51,
52, 53, 54, 55] ;
```

```
file_in_names := [dummy_1, dummy_2, dummy_3, dummy_4,
dummy_5, dummy_6, dummy_7, dummy_8, dummy_9, dummy_10,
dummy_11, dummy_12, dummy_13, dummy_14, dummy_15,
dummy_16, dummy_17, dummy_18, dummy_19, dummy_20,
dummy_21, dummy_22, dummy_23, dummy_24, dummy_25,
dummy_26, dummy_27, dummy_28, dummy_29, dummy_30,
dummy_31, dummy_32, dummy_33, dummy_34, dummy_35,
dummy_36, dummy_37, dummy_38, dummy_39, dummy_40,
dummy_41, dummy_42, dummy_43, dummy_44, dummy_45,
dummy_46, dummy_47, dummy_48, dummy_49, dummy_50,
dummy_51, dummy_52, dummy_53, dummy_54, dummy_55] ;
```

vic := 2

```
iniconc := [0.002083333333, 1.950 10-7, 1.73 10-8, 0., 0., 0.0000280, 0., 0., 1.0 10-9, 0., 0.,
0., 0., 0., 1.4985724 10-8, 0.000001913571534, 1.586379795 10-7, 7.32678405 10-8,
3.427321595 10-7, 7.943282347 10-12, 0.00178, 0.0000100, 0., 0., 0., 0., 0., 0., 0.,
0., 0., 0., 0., 0.000953, 0., 0.0000523, 0.0015, 1.73 10-8, 0., 7.857186444 10-14,
4.991913586 10-11, 2.292276799 10-15, 2.807159301 10-26, 9.210157539 10-43,
0.00072, 0., 0.0004691, 0.0000102, 0.00017, 0., 0., 0., 0.]
```

```
listinput := [1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24,
25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47,
```

```

48, 49, 50, 51, 52, 53, 54, 55]
file_in_names := [dummy_1, dummy_2, dummy_3, dummy_4, dummy_5, dummy_6,
dummy_7, dummy_8, dummy_9, dummy_10, dummy_11, dummy_12, dummy_13,
dummy_14, dummy_15, dummy_16, dummy_17, dummy_18, dummy_19, dummy_20,
dummy_21, dummy_22, dummy_23, dummy_24, dummy_25, dummy_26, dummy_27,
dummy_28, dummy_29, dummy_30, dummy_31, dummy_32, dummy_33, dummy_34,
dummy_35, dummy_36, dummy_37, dummy_38, dummy_39, dummy_40, dummy_41,
dummy_42, dummy_43, dummy_44, dummy_45, dummy_46, dummy_47, dummy_48,
dummy_49, dummy_50, dummy_51, dummy_52, dummy_53, dummy_54, dummy_55]

```

(12.1)

Output

noutput: number of species to be printed
nroutput: number of rates to be printed
listoutput: species number to print
listroutput: rate number to print
file_names: Respective file name for each of the species to print
file_rnames: Respective file name for each of the rates to print
time_iniout: First time (in years) for which a printout is requested
time_intvout: time interval (in years) at which the printing is performed, starting from time_iniout

example:
noutput:=4;
listoutput:=[1,2,3,5];
file_names:=[o2, so4, MnOx, hco3];
time_iniout:=10.;
time_intvout:= 100.;

```

> noutput := 55;
nroutput := 60;
listoutput := [1,2,3,4,5,6,7,8,9,10,11,12,13,14,15,16,
17,18,19,20,21,22,23,24,25,26,27,28,29,30,31,32,33,34,
35,36,37,38,39,40,41,42,43,44,45,46,47,48,49,50,51,52,
53,54,55] ;

listroutput := [1,2,3,4,5,6,7,8,9,10,11,12,13,14,15,
16,17,18,19,20,21,22,23,24,25,26,27,28,29,30,31,32,33,
34,35,36,37,38,39,40,41,42,43,44,45,46,47,48,49,50,51,
52,53,54,55,56,57,58,59,60] ;

file_names := [ch2o, zzo2, zno3, mno2, feoh3, zso4,
zch4, znh4, zpo4, zzm, zzfe, zh2s, zzhs, ch4g, h2co,
hco3, zco3, boh4, boh3, zzph, caco, zzca, snh4, spo4,
zfes, feco, zzs0, fes2, zsfe, zfp, mnco, age, mno2pr,
feoh3mr, feoh3pr, arago, mgcal, mg, BSi, DSi, sfSi,
al, aloh, aloh2, aloh3, aloh4, glass, plagio, na,
kplus, olive, pyrox, illite, smect, kaoli] ;

file_rnames := [xrate1, xrate2, xrate3, xrate4,

```



```

xrate5, xrate6, xrate7, xrate8, xrate9, xrate10,
xrate11, xrate12, xrate13, xrate14, xrate15, xrate16,
xrate17, xrate18, xrate19, xrate20, xrate21, xrate22,
xrate23, xrate24, xrate25, xrate26, xrate27, xrate28,
xrate29, xrate30, xrate31, xrate32, xrate33, xrate34,
xrate35, xrate36, xrate37, xrate38, xrate39, xrate40,
xrate41, xrate42, xrate43, xrate44, xrate45, xrate46,
xrate47, xrate48, xrate49, xrate50, xrate51, xrate52,
xrate53, xrate54, xrate55, xrate56, xrate57, xrate58,
xrate59, xrate60] ;

```

```

time_iniout := 0.9*tot_time ;
time_intvout := tot_time/10.0 ;

```

```

noutput := 55

```

```

nroutput := 60

```

```

listoutput := [1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24,
25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47,
48, 49, 50, 51, 52, 53, 54, 55]

```

```

listroutput := [1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23,
24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46,
47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60]

```

```

file_names := [ch2o, zzo2, zno3, mno2, feoh3, zso4, zch4, znh4, zpo4, zzm, zzfe, zh2s, zzhs,
ch4g, h2co, hco3, zco3, boh4, boh3, zzph, caco, zzca, snh4, spo4, zfes, feco, zzs0, fes2,
zsf, zfp, mnco, age, mno2pr, feoh3mr, feoh3pr, arago, mgcal, mg, BSi, DSi, sfSi, al,
aloh, aloh2, aloh3, aloh4, glass, plagio, na, kplus, olive, pyrox, illite, smect, kaoli]

```

```

file_rnames := [xrate1, xrate2, xrate3, xrate4, xrate5, xrate6, xrate7, xrate8, xrate9, xrate10,
xrate11, xrate12, xrate13, xrate14, xrate15, xrate16, xrate17, xrate18, xrate19, xrate20,
xrate21, xrate22, xrate23, xrate24, xrate25, xrate26, xrate27, xrate28, xrate29, xrate30,
xrate31, xrate32, xrate33, xrate34, xrate35, xrate36, xrate37, xrate38, xrate39, xrate40,
xrate41, xrate42, xrate43, xrate44, xrate45, xrate46, xrate47, xrate48, xrate49, xrate50,
xrate51, xrate52, xrate53, xrate54, xrate55, xrate56, xrate57, xrate58, xrate59, xrate60]

```

```

time_iniout := 361.4400000

```

```

time_intvout := 40.16000000

```

(13.1)

Optimization

there are several optimization options available. here you can specify what needs to be optimized and where the data is stored. to identify what kind of algorithm you want to use please select the appropriate options in drivervalues.f

nopt_v: number of parameters to be optimized

ntopt_v: number of time points where measurements are available

nparam_opt: total number of parameters. can include the nparam but also the physical parameters.

maxxmeas_v: maximum number of depth points at any given time measured (used to make array sizes)

maxspmeas_v: maximum number of species measured at any given time (used to make array sizes)

opt_name: names of the parameters. they have to match the names given above, so best you make a copy paste!

idpar_v: identify the parameters to be optimized from the parameter

```

list opt_name
filemeas_name: name of the files containing the measured data
if nopt_v is set to 0 then the rest of the input doesn't matter
example:
nopt_v := 2; # number of parameters to be optimized
ntopt_v := 3; # number of timepoints with measurements
nparam_opt := nparam; # total number of parameters, set equal to all parameters except physical ones
maxxmeas_v := 20; # maximum 20 points in a profile at any given time
maxspmeas_v := 2; # maximum 2 species measured at one timepoint
opt_name := bioname; # (note that this does not include the physical parameters! if you want them to be
adapted you need to specify them explicitly)
idpar_v := [1,3]; # optimize parameters 1 and 3 in the above list
filemeas_name := [meas1.dat, meas2.dat, meas3.dat]; # filenames with measurements at timepoints

> nopt_v := 0 ;
  ntopt_v := 0 ;
  nparam_opt := 0 ;
  maxxmeas_v := 0;
  maxspmeas_v := 0;
  opt_name := [ ];
  idpar_v := [ ];
  filemeas_name := [ ];

                                nopt_v := 0
                                ntopt_v := 0
                                nparam_opt := 0
                                maxxmeas_v := 0
                                maxspmeas_v := 0
                                opt_name := [ ]
                                idpar_v := [ ]
                                filemeas_name := [ ]

```

(14.1)

Maple specific info

```

dir_f: directory where the FORTRAN routines and Maple spread.m files
are parsed
format Mac: "Macintosh HD:UU:...:code"
format PC: "C:\\maple\\...\\code"
WAS: dir_f := "C:\\Dokumente und Einstellungen\\centler\\Desktop\\Labor\\Simulations":
currentdir(dir_f):
save "spread.m" ;
> dir_f :=
  "/Users/sandra/Desktop/Models/BRNSPackage-2013-07-03/"
  :
  currentdir(dir_f) :
  parse(sprintf("save %q,\"spread.m\";", anames()),
  statement) ;

> "now execute processor - make sure the directories are
  set correctly";

                                "now execute processor - make sure the directories are set correctly"

```

(15.1)

