

Documentation: data input

- **Transient 1D reactive transport class:**
 - **Line 1:** Name of the problem
 - **Line 2:** Section separation
 - **Line 3:** Section name
 - **Line 4:** *Num_time* (integer)
 - Number of time steps
 - **Line 5:** *Delta_t* (real)
 - Time step
 - Note: If discretization is not uniform, you have to write the different time steps on the same line
 - **Line 6:** *int_method* (integer)
 - Time integration method:
 1. Explicit Euler
 2. Euler fully implicit
 3. Crank-Nicolson
 - **Line 7:** end of section

```
'TITLE OF THE PROBLEM: Yeso en equilibrio'
```

```
'-----'
```

```
'TIME'
```

```
10          ! number of time steps
```

```
1d1         ! time step ( we assume uniform time discretisation )
```

Δt

```
1          ! time integration method for chemical reactions
```

```
'*'
```

```
'-----'
```

```
'end'
```

- **Transient 1D transport class:**

- **Line 1:** Name of the problem
- **Line 2:** Section separation
- **Line 3:** Section name
- **Line 4:** *Flag_porosity* (logical), *porosity* (real scalar or vector)
 - *Flag_porosity=true*: porosity is homogenous
 - *Flag_porosity=false*: porosity is heterogeneous (in this case you must write the porosities in the same line)
- **Line 5:** end of section
- **Line 6:** Section separation
- **Line 7:** Section name
- **Line 8:** *num_mix_ratios* (integer), *mixing_ratios* (real vector)
 - *num_mix_ratios*: number of mixing ratios in each target
 - *mixing_ratios*: mixing ratios in each target (contains sink/sources and boundary terms)
- **Line 9:** end of mixing ratios pseudomatrix
- **Line 10:** end of section
- **Line 11:** section separation
- **Line 12:** name of section
- **Line 13:** *ind_tar_wat* (integer), *mix_wat_indices* (integer vector)
 - *ind_tar_wat*: index of target water
 - *mix_wat_indices*: indices of target waters associated to each mixing ratio
- **Line 14:** end of section
- **Line 15:** end of file

'TITLE OF THE PROBLEM: Yeso en equilibrio'

'TRANSPORT PROPERTIES'

.true. 5d-1 ! in this line: flag for homogeneous property, porosity

$$\phi$$

'MIXING RATIOS'

! in each line: number of mixing ratios, mixing ratios at each target (including sink/sources and boundary terms)

1 1.00000E+00
3 9.20000E-01 4.20000E-02 3.80000E-02
3 9.20000E-01 4.20000E-02 3.80000E-02
3 9.20000E-01 4.20000E-02 3.80000E-02
3 9.20000E-01 4.20000E-02 3.80000E-02
3 9.20000E-01 4.20000E-02 3.80000E-02
3 9.20000E-01 4.20000E-02 3.80000E-02
3 9.20000E-01 4.20000E-02 3.80000E-02
3 9.20000E-01 4.20000E-02 3.80000E-02
3 9.20000E-01 4.20000E-02 3.80000E-02
1 1.00000E+00

$$\lambda_{ij}$$

0 ! indicates the end of the mixing ratios pseudomatrix

'MIXING WATERS'

! target water index, water indices in 'MIXING RATIOS'

1 1
2 2 1 3
3 3 2 4
4 4 3 5
5 5 4 6
6 6 5 7
7 7 6 8
8 8 7 9
9 9 8 10
10 10 9 11
11 11

$$\mathbf{c}_i^{k+1} = \sum_{j \in I_i} \lambda_{ij} \mathbf{c}_j^k$$

$$I_1, I_2, I_3, \dots$$

'end'

- **Chemical system class:**
 - **Line 1:** Name of the problem
 - **Line 2:** Section separation
 - **Line 3:** Section name
 - If Section name = 'PRIMARY AQUEOUS SPECIES' or 'AQUEOUS COMPLEXES':
 0. **Line 3a:** *sp_name* (string)
 - Name of species in database "master25_modif.dat"
 1. **Line 3b:** end of section
 - If Section name = 'MINERALS' or 'GASES':
 0. **Line 3a:** *sp_name* (string), *flag_eq* (logical), *flag_cst_act* (logical)
 - *sp_name*: Name of species in database "master25_modif.dat"
 - *flag_eq*: TRUE if equilibrium reaction, FALSE otherwise
 - *flag_cst_act*: TRUE if species has constant activity, FALSE otherwise
 1. **Line 3b:** end of section
 - If Section name = 'REDOX REACTIONS':
 0. **Line 3a:** *react_name* (string), *flag_eq* (logical)
 - *react_name*: Name of redox reaction in database "reacciones_monod.dat"
 - *flag_eq*: TRUE if equilibrium reaction, FALSE otherwise
 1. **Line 3b:** end of section
 - If Section name = 'SURFACE COMPLEXES':
 0. **Line 3a:** *sp_name* (string)
 - *sp_name*: Name of species in database "master25_modif.dat"
 1. **Line 3b:** end of section
 - **Line 4:** end of section
 - **Line 5:** end of file

```
'TITLE OF THE PROBLEM: Gypsum in equilibrium'

'-----'

'PRIMARY AQUEOUS SPECIES'

'h2o'                ! name of primary aqueous species

'ca+2'

'so4-2'

'*'

'AQUEOUS COMPLEXES'

'*'                ! name of aqueous complex

'MINERALS'

'gypsum' .true. .true.    ! name of mineral, flag equilibrium, flag constant activity

'*' .true. .true.

'GASES'

'*' .true. .true.        ! name of gas, flag equilibrium, flag constant activity

'SURFACE COMPLEXES'

'*'                ! name of surface complex

'-----'

'end'
```

- **Water type class:**

- **Line 1:** Name of the problem
- **Line 2:** Section separation
- **Line 3:** Section name: 'INITIAL AND BOUNDARY WATER TYPES'
- **Line 4:** *act_coeffs_model* (integer)
 - 0. Ideal
 - 1. Debye-Hückel
 - 2. Debye-Hückel extended
 - 3. Davies
 - 4. Truesdell-Jones
- **Line 5:** *num_wat_types* (integer)
 - 0. Number of water types (include initial, recharge, boundary etc)
- **Line 6:** *ind_wat_type* (integer), *temp* (real)
 - *ind_wat_type*: index of water type
 - *temp*: temperature of solution
- **Line 6a:** *name* (string)
 - **Name of water type**
- **Line 6b:** water type data begins
 - *Icon*: initial condition type
 - 1. prescribed primary concentration
 - 2. prescribed total concentration
 - 3. prescribed activity
 - 4. primary species is in equilibrium with a phase
 - *Guess*: Initial guess for primary species concentration
 - *Ctot*:
 - 0. *Icon*=1: primary species concentration
 - 1. *Icon*=2: total concentration of primary species
 - 2. *Icon*=3: activity of primary species
 - 3. *Icon*=4: activity of phase that is in equilibrium with this primary species
 - *Constrain*: if *icon*=4, it is the phase in equilibrium with the primary species
- **Line 6c:** *prim_sp_name* (string), *icon* (integer), *guess* (real), *ctot* (real), *constrain* (string)
- **Line 6d:** end of water type data
- **Line 7:** end of section

'TITLE OF THE PROBLEM: Yeso en equilibrio'

'-----'

'INITIAL AND BOUNDARY WATER TYPES'

0 ! activity coefficients model

2 ! number of water types

1 25.0 ! index water type, temp (C)

'entrada' ! name of water type

' icon guess ctot constrain'

'h2o' 1 1d0 1d0 ''

'ca+2' 1 1d-3 1d-3 ''

'so4-2' 4 1d-5 1d0 'gypsum'

'*' 0 0.0 0.0 ''

2 25.0 ! index water type, temp (C)

'inicial' ! name of water type

' icon guess ctot constrain'

'h2o' 1 1d0 1d0 ''

'ca+2' 1 1d-5 1d-5 ''

'so4-2' 4 1d-5 1d0 'gypsum'

'*' 0 0.0 0.0 ''

'-----'

- **Solid type class:**

- **Line 1:** Section name: 'INITIAL MINERAL ZONES'
- **Line 2:** *num_min_zones* (integer)
 - Number of mineral zones
- **Line 3a:** *ind_min_zone* (integer), *temp* (real)
 - *ind_min_zone*: index of mineral zone
 - *temp*: temperature of mineral zone
- **Line 3b:** mineral zone data begins
- **Line 3c:** *min_name* (string), *vol_frac* (real), *react_surf* (real)
 - *min_name*: name of mineral in mineral zone (must be in the chemical system)
 - *vol_frac*: initial volumetric fraction of mineral
 - *react_surf*: specific surface area
- **Line 3d:** end of block
- **Line 4:** end of section

INITIAL MINERAL ZONES'			
1			! number of mineral zones
1	25.0		! index of mineral zone, temp (C)
'mineral	vol.frac.	area(m2/m3rock)'	
'gypsum'	0.50	0.1d+3	
'*	0.00	0.00	
* _____			

- **Gas type class:**

- **Line 1:** Section name: 'INITIAL GAS ZONES'
- **Line 2:** *num_gas_zones* (integer)
 - Number of gas zones
- **Line 3a:** *ind_gas_zone* (integer), *temp* (real), *vol* (real)
 - *ind_gas_zone*: index of gas zone
 - *temp*: temperature of gas zone
 - *vol*: total volume of gas
- **Line 3b:** gas zone data begins
- **Line 3c:** *gas_name* (string), *part_press* (real)
 - *gas_name*: name of gas in gas zone (must be in the chemical system)
 - *part_press*: initial partial pressure (equivalent to activity) of gas
- **Line 3d:** end of block
- **Line 4:** end of section
- **Line 5:** end of file

```

'INITIAL GAS ZONES'

1                                     ! number of initial gas zones

1          25          1d0          ! index gas type, temp (°C), volume

'gas          partial pressure'          ! pressure in atm

'co2(g)'          3.5d-4

'o2(g)'          2d-1

'n2(g)'          7.99965d-1

'***          0.00

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'end'

```

- **Chemistry class:**

- **Line 3:** Section name: 'TARGET WATERS'
- **Line 4:** *num_tar_wat* (integer)
 - Number of target waters
- **Line 5:** Subsection name: 'external waters'
- **Line 5a:** *num_ext_wat* (integer)
 - Number of external target waters
- **Line 5b:** *ind_ext_wat* (integer), *iwtype* (integer)
- **Line 6a:** *ind_tar_wat* (integer), *iwtype* (integer), *ind_sol_zone* (integer), *ind_gas_zone* (integer)
 - *ind_tar_wat*: Index target water
 - *iwtype*: Index water type
 - *ind_sol_zone*: Index solid zone
 - *ind_gas_zone*: Index gas zone
- **Line 3d:** end of section

'TITLE OF THE PROBLEM: Yeso en equilibrio'

'TARGET WATERS'

12 ! number of target waters

'external waters'

2 ! number of external target waters (recharge, boundary, etc)

```
1      3      ! target water index, water type index
```

2 2

'initial target waters'

3 2 1 0 ! target water index, water type index, solid zone index, gas zone index

4 2 1 0

5 2 1 0

6 2 1 0

7 2 1 0

8 2 1 0

9 2 1 0

10 2 1 0

11 2 1 0

12 2 1 0

'end'