



# **The PFLOTRAN Reaction Sandbox**

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**Abstract.** As modern reactive transport simulators evolve to accommodate the demands of a user community, researchers need a platform for prototyping new biogeochemical processes, many of which are niche and specific to laboratory or field experiments. The PFLOTRAN Reaction Sandbox leverages modern, object oriented Fortran in an attempt to provide such an environment within an existing reactive transport simulator. This work describes the PFLOTRAN Reaction Sandbox con-

5 cept and implementation through several illustrative examples. Reaction Sandbox Biodegradation Hill customizes the existing microbially-mediated biodegradation reaction formulation within PFLOTRAN to better match empirical data. Reaction Sandbox Simple provides an isolated environment for testing numerous preconfigured kinetic rate expressions and developing user intuition. Reaction Sandbox Example serves as a template for creating new sandboxes within PFLOTRAN.

# 1 Introduction

10 Modern reactive transport simulators incorporate sophisticated networks of reactions to simulate complex biogeochemical processes within the Earth's subsurface environment in support of scientific research in climate change, fate and transport of contaminants, and water resources management (Steefel et al., 2005). As these simulators mature and evolve over time, they can accumulate a large number of chemical reactions with a wide range of implementations. Many reactions are somewhat common among reactive transport codes [e.g. aqueous complexation, mineral precipitation-dissolution, radioactive decay and sorption (Steefel et al., 2015)], while others are niche, prototypical or problem dependent.

There is a multitude of possible reactions to include within a simulator, and the approach to implementing these reactions often varies among simulators. For instance, sorption processes includes absorption, adsorption and ion exchange. Surface complexation is an adsorptive process that can be simulated using a constant capacitance, (diffuse) double layer, triple layer, or non-electrostatic model (Bethke, 2007). Depending on the reaction timescales, surface complexation may be simulated

20 assuming local equilibrium or driven by a kinetic rate expression, and kinetic approaches may include single and multi-rate models, the latter utilizing a distribution of rate constants associated with size fractions of sediment material (Liu et al., 2008).

It is also common for researchers to develop one-off implementations of reactions when existing capabilities cannot replicate biogeochemical phenomenon observed in field or laboratory experiments. For example, Tutolo et al. (2018) demonstrated that the brucite silicification is a serpentinization rate-limiting reaction that is exponentially dependent upon aqueous silica activity.

25 Their brucite silicification reaction required the implementation of a custom rate law that was exponentially dependent upon the activity of aqueous silica. These one-off implementations quick increase the diversity of reactions supported by a simulator.





As the number of reactions implemented within a simulator grows, and the reactions become increasingly diverse, adding new reactions can challenge ongoing simulator development and maintenance, especially within an open source community where code development is often crowdsourced.

- 30 Due to time and funding limitations, it is difficult for code developers to satisfy the needs of the user community by implementing all variants of a reactive process. Nor does it make sense when reactions are problem-specific and may never be used in the future. Meanwhile, the end users who are requesting customization are often non-computational domain scientists with less interest in or limited understanding of numerical methods, programming paradigms, code abstractions, and data layout. Their focus is more on improving predictive accuracy through the refinement of reaction conceptual models, not code development.
- 35 This raises the question as to what steps can be taken in the design and implementation of a reactive transport simulator to facilitate code development and long-term maintenance while flattening the (often) steep learning curve for new developers.

The PFLOTRAN Reaction Sandbox is an attempt to provide such an environment within an existing reactive transport simulator. The purpose of the Reaction Sandbox is to provide a means for testing alternative implementations for kinetically formulated rate expressions or networks of these reactions in conjunction with the existing reactive transport capability within

40 PFLOTRAN. Within computer science, the term *sandbox* often refers to an environment for implementing and vetting new, untested algorithms in isolation. The purpose of a sandbox is to limit the impact on the remainder of the code.

The following sections document the implementation of the PFLOTRAN Reaction Sandbox and demonstrate its application on several problem scenarios. Section 2 provides an overview of PFLOTRAN by presenting the governing equations for reactive transport and the numerical methods employed to solve the resulting discrete nonlinear systems of equations. The

45 section also presents PFLOTRAN's conventional reactive transport capability and discusses the code's modular object orienteddesign that facilitates the implementation of the Reaction Sandbox. Section 3 describes the foundational (Fortran) Reaction Sandbox class upon which all Reaction Sandbox classes are coded and the high-level programming interface through which the remainder of the code accesses sandbox reactions. Section 4 documents several example sandboxes from which a researcher may derive their own implementation. Finally, Section 5 summarizes the approach.

# 50 2 Background

PFLOTRAN is an massively parallel, reactive multiphase flow and transport simulator for modeling subsurface earth system processes (Hammond et al., 2014). The code has been developed under open source GNU LGPL licensing since 2009 and contains contributions from an international group of developers. PFLOTRAN is written in Fortran 2003/2008 which facilitates object oriented design through the use of nested derived types, classes (extensible derived types with member procedures) and

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procedures pointers. The code is designed as a nested hierarchy of objects. The top-level simulation object contains pointers to all process models, solvers and supporting data (state variables, parameters, etc.) needed to run a simulation.

PFLOTRAN simulates biogeochemical transport through its reactive transport process model. Supported biogeochemical reaction capabilities include aqueous speciation with ion activity models, general N<sup>th</sup>-order forward (and reversible) kinetic reactions, microbially-mediated reactions, mineral precipitation-dissolution, radioactive decay and ingrowth and sorption (An-





60 dre et al., 2021). These reactions are referred to as *conventional* reactive transport capability as the implementations are based on commonly accepted and well-documented in the literature.

#### 2.1 Governing Equations

PFLOTRAN's governing mass conservation equation for reactive transport of aqueous species j is

$$\frac{\partial}{\partial t} (\phi s \Psi_j) + \nabla \cdot (\boldsymbol{q} - \phi s \boldsymbol{D} \nabla) \Psi_j = Q_j - \sum_r \nu_{jr} I_r, \tag{1}$$

65 with porosity  $\phi$ , liquid saturation *s*, total aqueous component concentration  $\Psi_j$ , Darcy fluid flux *q*, diagonal hydrodynamic dispersion tensor *D* and source term  $Q_j$ .  $\nu_{jr}$  represents the stoichiometry of species *j* in kinetic reaction  $I_r$ . Following the continuum formulation for reactive transport (Lichtner, 1985) and assuming local equilibrium (Rubin, 1983), the total aqueous component concentration of species *j* is the sum of the free ion concentration  $c_j$  and its stoichiometric contribution  $\nu_{ji}$  to each secondary aqueous complex  $X_i$ ,

$$70 \quad \Psi_j = c_j + \sum_i \nu_{ji} X_i. \tag{2}$$

Aqueous complex  $X_i$  is calculated through mass action as

$$X_i = \frac{K_i}{\gamma_i} \prod_{j'} \left( \gamma_{j'} c_{j'} \right)^{\nu_{j'i}} \tag{3}$$

with equilibrium constant  $K_i$ , activity coefficients  $\gamma_i$  and  $\gamma_{j'}$ , stoichiometry  $\nu_{j'i}$ , and primary aqueous species free ion concentration  $c_{j'}$ . The governing mass conservation equation for the  $j^{th}$  primary immobile species  $\Phi$  is

75 
$$\frac{\partial}{\partial t}(\Phi_j) = -\sum_r \nu_{jr} I_r.$$
 (4)

Ignoring the advection, hydrodynamic dispersion and source terms and assuming constant porosity and saturation, the discrete (finite volume) forms of Eqs. (1) and (4) are

$$\frac{\phi s V}{\Delta t} \left( \Psi_j^{k+1} - \Psi_j^k \right) \dots = \dots - V \sum_r \nu_{jr} I_r$$
(5)

and

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$$0 \quad \frac{V}{\Delta t} \left( \Phi_j^{k+1} - \Phi_j^k \right) = -V \sum_r \nu_{jr} I_r, \tag{6}$$

respectively. Units for these equations are [mole/second]. Rate expression  $I_r$  represents the primary species mass consumed or produced by each kinetic reaction and has units of [mole/m<sup>3</sup><sub>bulk</sub>-second].

# 2.2 Numerical Solution Technique

PFLOTRAN employs the finite volume method for spatial discretization, backward Euler time integration and Newton's method for solving the resulting nonlinear system of equations. Newton's method converges to a solution for the primary





species concentrations x by iteratively evaluating the residual function

$$f(x^p) = 0 \tag{7}$$

and Jacobian J, and solving the linear system

$$J\delta x = -f(x^p) \tag{8}$$

90 for the concentration update  $\delta x$ 

$$x^{p+1} = x^p + \delta x. \tag{9}$$

p is iteration number.

The residual function f is evaluated by rearranging Eqs. (5) and (6) and setting them equal to zero. For example, the residual for aqueous degree of freedom  $c_n$  is

95 
$$f(c_n^p) = \frac{\phi s V}{\Delta t} \left( \Psi_n^{k+1} - \Psi_n^k \right) \dots + V \sum_r \nu_{nr} I_r = 0,$$
 (10)

while for immobile degree of freedom  $\Phi_n$ , it is

$$f(\Phi_n^p) = \frac{V}{\Delta t} \left( \Phi_n^{k+1} - \Phi_n^k \right) + V \sum_r \nu_{nr} I_r = 0.$$
(11)

Since the focus of this research is chemical reaction, the advection, dispersion and source terms (represented by ellipses ...) are ignored. The Jacobian contains derivatives of the residual with respect to the primary unknowns, i.e.

100 
$$J_{n,m} = \frac{\partial f(x_n^p)}{\partial x_m^p}.$$
 (12)

Units for internal PFLOTRAN variables are documented in Table 1. These units must be considered in the development of a Reaction Sandbox.

# 2.3 Object-Oriented Fortran

long-term maintenance of a code.

Modern programming paradigms facilitate the modularity, extensibility and overall longevity of a software product. One common feature among most modern programming paradigms is support for object-oriented design, where objects contain the data structures and procedures necessary to provide functionality, and interfaces are set up for interaction between objects. One major benefit of object oriented design is that when programmed correctly, modifications to an object's data structures and procedures have little to no impact on other portions of the code. This modularity greatly facilitates the initial development and

Modern Fortran facilitates object-oriented design through the use of derived types and classes. A Fortran derived type is a container that encapsulates other Fortran data types (e.g. logicals, integers, reals, other derived types, or pointers to data types). A Fortran class is an extensible version of the derived type that supports inheritance of (type-bound) member variables and





Table 1. Units for PFLOTRAN internal variables.

Variables	Symbol	Units
Porosity	$\phi$	m <sup>3</sup> pore / m <sup>3</sup> bulk volume
Liquid saturation	s	m <sup>3</sup> liquid / m <sup>3</sup> pore
Volume	V	m <sup>3</sup> bulk volume
Time	t	seconds
Aqueous free ion concentration	c	mole / kg water (or molality)
Total aqueous component concentration	$\Psi$	mole / L water (or molarity)
Immobile concentration	Φ	mole / m <sup>3</sup> bulk volume
Stoichiometry	ν	-
Kinetic rate	$I_r$	mole / m <sup>3</sup> bulk volume - second
Residual function	$f(x^p)$	mole / second
Derivative of residual wrt free ion concentration	$\frac{\partial f(x_n^p)}{\partial c_m^p}$	kg water / second
Derivative of residual wrt immobile concentration	$\frac{\partial f(x_n^p)}{\partial \Phi_m^p}$	m <sup>3</sup> bulk volume / second

procedures. A child class, created by extending the parent derived type, inherits all parent class variables and procedures. New member variables may be added to child classes and member procedures may be overridden. By default, a member procedure

receives the class object as the first entry in function or subroutine argument lists. (A class object is simply an instantiation of 115 the class.) The procedure uses the object's member variables and the remaining arguments to perform calculations. Chapman (2018) describes Fortran classes in greater detail.

PFLOTRAN is designed as a nested hierarchy of dynamically allocated objects from the highest-level simulation object down to low level, cell-centric auxiliary objects that store all state variables for each grid cell. Given a pointer to the top-level simulation object, the developer has access to all underlying data structures or objects. This hierarchy engenders modularity 120 and structure within the code. PFLOTRAN employs modern Fortran 2003/2008 where all objects are instantiations of Fortran derived types or classes. It also leverages pointers to procedures and common procedure interfaces to allow users to choose from a suite of constitutive relations, equations of state, flux algorithms, etc., in support of run time options. The modularity afforded through object-oriented Fortran has greatly facilitated the incorporation of new algorithms within PFLOTRAN, as

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# will be shown through the implementation of the Reaction Sandbox.

# **3** Reaction Sandbox Approach

The PFLOTRAN Reaction Sandbox provides a simplified interface for implementing new kinetically formulated reactions (rate expressions) within PFLOTRAN that are tailored to specific user needs. The term  $I_r$  on the right side of Eqs. (1) and (4) represents these kinetic reactions in the governing equations. The Reaction Sandbox isolates one-off or application-specific

reactions from the remainder of the PFLOTRAN code base and facilitates code development and long-term maintenance. For

the domain scientist desiring to implement a new reaction, the Reaction Sandbox reduces complexity by exposing only the

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limited set of variables that are necessary to calculate rate expressions. Thus, the researcher is better shielded from the intricate details of code development. For kinetic reactions that have the potential for wider acceptance, the Reaction Sandbox provides a venue for vetting reactions in isolation prior to adoption within the main code base. This section describes the Reaction
135 Sandbox concept and defines the underlying data structures and user interface.

#### 3.1 Concept

Figure 1 illustrates the hierarchical structure of (hypothetical) Reaction Sandboxes within PFLOTRAN where reaction classes are derived as descendants of the Reaction Sandbox Base Class. Each Reaction Sandbox class is implemented as a separate



Figure 1. Schematic of a hypothetical Reaction Sandbox class hierarchy.

module within the PFLOTRAN source code. The end-user specifies the reaction sandboxes to be employed within the PFLO-TRAN input file, and a linked list of sandbox reactions is constructed during simulation initialization. Figure 2 illustrates a



**Figure 2.** Schematic of a linked list of Reaction Sandboxes composed of two reaction objects instantiated from the class hierarchy shown in Fig. 1.

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representative linked list of sandboxes composed of two reaction objects from Fig. 1. The outer Reaction Sandbox module loops over the linked list to perform all operations (e.g. setup, evaluation, destruction, etc.) as illustrated later in Code Block 2. Linked lists provide great flexibility for the code developer as all reactions may be inserted or appended in any order. Newly developed reaction classes are added to the source code as daughter classes in new modules, and instantiated objects are added

145 to the list during simulation initialization.





# 3.2 Implementation

The Reaction Sandbox is founded upon two files within the PFLOTRAN source code: reaction\_sandbox\_base.F90 and reaction\_sandbox.F90.

# 3.2.1 reaction\_sandbox\_base.F90

150 reaction\_sandbox\_base.F90 defines the base (or parent) Fortran class reaction\_sandbox\_base\_type that the developer extends to create new (child) Reaction Sandbox classes. Code Block 1 illustrates the member variables and procedures within the reaction\_sandbox\_base\_type class.

Code Block 1. reaction\_sandbox\_base\_type in reaction\_sandbox\_base.F90.

```
type, abstract, public :: reaction_sandbox_base_type
class(reaction_sandbox_base_type), pointer :: next
contains
procedure, public :: ReadInput => BaseReadInput
procedure, public :: Setup => BaseSetup
procedure, public :: Evaluate => BaseEvaluate
160 procedure, public :: UpdateKineticState => BaseUpdateKineticState
procedure, public :: AuxiliaryPlotVariables => BaseAuxiliaryPlotVariables
procedure, public :: Destroy => BaseDestroy
end type reaction_sandbox_base_type
```

- 165 This class contains a single member variable next, a pointer to the next class of the same reaction\_sandbox\_base\_type type that enables the creation of an abstract linked list of Reaction Sandbox objects. The class also contains empty member procedures with prescribed subroutine interfaces (or argument lists) that the developer overrides in child classes. With the exception of BaseEvaluate, these procedures are empty and return immediately if not overridden by the child class. BaseEvaluate must be extended, and error messaging is incorporated within BaseEvaluate to ensure correct implementation. The other
- 170 member procedures are optional and do not require implementation in the child classes. Appendix A documents the member procedure interfaces describing the subroutine argument lists.

# 3.2.2 reaction\_sandbox.F90

reaction\_sandbox.F90 serves as the main driver interface for the Reaction Sandbox, providing subroutines that manage the creation, reading, setup, execution and destruction of all Reaction Sandbox objects. For example, the subroutine RSandboxRead instantiates Reaction Sandbox objects based on the keywords parsed from the REACTION SANDBOX block in

the input file. RSandboxEvaluate calculates reaction rates by traversing the linked list of Reaction Sandboxes evaluating individual rates as shown in Code Block 2 (subroutine argument lists have been omitted for simplicity).





```
Code Block 2. Abbreviated version of subroutine RSandboxEvaluate in reaction_sandbox.F90.
    subroutine RSandboxEvaluate (...)
180
       . . .
       class (reaction_sandbox_base_type), pointer :: cur_reaction
       cur_reaction => rxn_sandbox_list
      do
185
         if (.not.associated(cur_reaction)) exit
         call cur_reaction%Evaluate (...)
         cur_reaction => cur_reaction%next
       enddo
```

190 end subroutine RSandboxEvaluate

With the exception of RSandboxRead, all member procedures are executed from within a linked list loop similar to that shown in Code Block 2. Thus, subroutines within reaction\_sandbox.F90 serve as interfaces to the linked lists of Reaction Sandbox objects, and all information is exchanged through these routines. The outer RSandboxXXX subroutines defined in reaction sandbox.F90 may be called from other PFLOTRAN modules (e.g. see the call to RSandboxEvaluate

```
195
```

within RReaction within reaction.F90), but member procedures within the Reaction Sandbox classes may not. The following section describes several example Reaction Sandboxes. These examples are implemented within the PFLO-

TRAN source code and may serve as templates for future sandboxes.

#### **Example Reaction Sandboxes** 4

- 200 This section illustrates the implementation of a few Reaction Sandboxes within PFLOTRAN. Reaction Sandbox Biodegradation Hill and Reaction Sandbox Flexible Biodegradation Hill were developed to demonstrate the implementation of a microbially-mediated biodegradation reaction. Reaction Sandbox Simple provides a set of preconfigured reactions that may be uncommented, compiled and run to better understand kinetic rate expressions. Reaction Sandbox Example implements a firstorder decay reaction. The comments within reaction\_sandbox\_example.F90 detail the steps necessary to implement a new Reaction Sandbox class.
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# 4.1 Biodegradation

# 4.1.1 Biodegradation Conceptual Model

Consider a microbially-mediated biodegradation reaction with biomass growth and decay over time. The reaction could be expressed as

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$$A_{aq} + 0.25B_{aq} \to 0.33C_{aq} + D_{aq},$$
 (R1)

with electron donor  $A_{aq}$  and acceptor  $B_{aq}$  and products  $C_{aq}$  and  $D_{aq}$  [M]. The reaction is mediated by the immobile biomass species  $X_{im}$  [mole biomass/m<sup>3</sup> bulk] and inhibited above a  $C_{aq}$  concentration of  $10^{-4}$ . The reaction rate  $I_r$  [mole/m<sup>3</sup> bulk/sec] can be calculated using Michaelis-Menten kinetics as

$$I_r = k_{\max} X_{im} \frac{A_{aq}}{K_{A_{aq}} + A_{aq}} \times \frac{B_{aq}}{K_{B_{aq}} + B_{aq}} \times \frac{I_{C_{aq}}}{I_{C_{aq}} + C_{aq}}$$
(13)

with maximum specific utilization rate constant  $k_{max}$  [mole/sec-mole biomass], half saturation constants  $K_{A_{aq}}$  and  $K_{B_{aq}}$  [M], and inhibitor concentration  $I_{C_{aq}}$  [M]. The rate of biomass growth and decay can be modeled as

$$\frac{dX}{dt} = yield_{X_{im}}I_r - k_{\text{decay}}X \tag{14}$$

with yield  $yield_{X_{im}}$  [mole biomass/mole] and decay rate constant  $k_{decay}$  [1/sec]. These rate expressions are implemented as microbial and biomass decay reactions within PFLOTRAN and enabled through the MICROBIAL\_REACTION and IMMO-BILE\_DECAY\_REACTION keywords in CHEMISTRY block of the input file.

Figure 3 shows PFLOTRAN simulation results for an example batch experiment run over seven days employing the reactions in Eqs. (13) and (14), reaction parameters in Table 2, stoichiometries in Reaction (R1), and initial conditions in Table 3. Results plotted in the figure may be replicated through the following commands:

cd \$PFLOTRAN\_DIR/regression\_tests/default/reaction\_sandbox

\$PFLOTRAN\_DIR/src/pflotran/pflotran -input\_prefix biodegradation

python biodegradation\_vs\_data.py biodegradation-obs-0.pft

The plot shows the time evolution of aqueous and immobile species concentrations as the week-long simulaton runs with a maximum time step size of one hour. The results demonstrate that electron donor  $A_{aq}$  is the limiting substrate, as the concentations for acceptor  $B_{aq}$  and the reaction products  $C_{aq}$  and  $D_{aq}$  plateau when  $A_{aq}$  is nearly depleted at approximately

four days. Biomass concentration increases through 2.25 days at which time the first-order decay rate exceeds the growth rate, and the population begins to fade. Superimposed on the figure are hypothetical experimental results for species  $A_{aq}$  (shown as circles) that deviate from the (blue) simulated curve in the log scale plot. In this hypothetical case, Eq. (13) is somewhat inaccurate in predicting the tailing behavior of species  $A_{aq}$  at late times.







Figure 3. Time evolution of aqueous and immobile biomass species in a week-long example batch biodegradation experiment.

**Table 2.** Microbially-mediated reaction parameters for the example batch biodegradation experiment. n only applies to the reaction incorporating the Hill function (i.e. Eq. (15)). M signifies molarity or mole per liter of water.

Parameter	Value	Units	
$k_{\max}$	9.e-2	mole/mole biomass-sec	
$K_{A_{aq}}$	2.e-4	М	
$K_{B_{aq}}$	1.25e-5	М	
$I_{C_{aq}}$	2.5e-4	М	
$yield_{X_{im}}$	1.e-4	mole biomass/mole	
$k_{ m decay}$	1.e-6	1/sec	
n	1.2	-	

 Table 3. Initial concentrations for the example batch biodegradation experiment.

Species	Concentration	Units
$A_{aq}$	1.e-3	М
$B_{aq}$	5.e-4	М
$C_{aq}$	1.e-10	М
$D_{aq}$	1.e-10	М
$X_{im}$	1.e-4	mole/m <sup>3</sup> bulk volume

The discrepancy in  $A_{aq}$  concentration can be resolved by employing a Hill function (Hill, 1910) within the monod expression 235 for  $A_{aq}$ . Here, the  $A_{aq}$  concentration and corresponding half saturation constant are raised to the power n.

$$I_{r} = k_{\max} X_{im} \frac{A_{aq}^{n}}{K_{A_{aq}}^{n} + A_{aq}^{n}} \times \frac{B_{aq}}{K_{B_{aq}} + B_{aq}} \times \frac{I_{C_{aq}}}{I_{C_{aq}} + C_{aq}}$$
(15)





However, the Hill function is not an option in PFLOTRAN; the code must be altered to accommodate this new feature. To further complicate matters, the researcher cannot modify the existing Monod expression within PFLOTRAN since in doing so, the Hill function would be applied to both the A<sub>aq</sub> and B<sub>aq</sub> Monod expressions. One possible solution is to implement Eq. (15)
as a new reaction, and the Reaction Sandbox is designed to facilitate this process.

# 4.1.2 Reaction Sandbox Biodegradation Hill

Reaction Sandbox Biodegradation Hill implements the enhanced biodegradation reaction that incorporates the Hill function in Eq. (15) and the biomass growth and decay reaction in Eq. (14). The reactions are encoded in reaction\_sandbox\_biohill.F90. The reaction\_sandbox\_biohill\_type class is presented in Code Block 3 where integer IDs for each species are stored as class variables and the Setup and Evaluate procedures are redirected to local BioHillXXX implementations.

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Code Block 3. reaction sandbox biohill\_type in reaction sandbox biohill.F90.

```
type, public, &
      extends(reaction_sandbox_base_type) :: reaction_sandbox_biohill_type
      ! Aqueous species
      PetscInt :: species Aag id
250
      PetscInt :: species_Baq_id
      PetscInt :: species_Caq_id
      PetscInt :: species Dag id
      ! Immobile species (e.g. biomass)
      PetscInt :: species_Xim_id
255
    contains
      procedure, public :: Setup => BioHillSetup
      procedure, public :: Evaluate => BioHillEvaluate
    end type reaction_sandbox_bio_type
260
```

Code Block 4 illustrates the assignment of species IDs in an abbreviated version of BioHillSetup based on keywords for aqueous species Aaq, Baq, Caq and Daq and immobile species Xim specified in the input file. BioHillEvaluate utilizes these IDs to access species concentrations and entries in the residual vector.

Code Block 4. Subroutine BioHillSetup in reaction\_sandbox\_biohill.F90.



. . .

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! Immobile species word = 'Xim' this%species\_Xim\_id = & GetImmobileSpeciesIDFromName(word, reaction%immobile, option)

# end subroutine BioHillSetup

Code Block 5 presents an abbreviated version of the implementation of BioHillEvaluate where the naming convention for local variables corresponds closely to reaction parameters and state variables defined in Eqs. (14) and (15). Note that in the calculation of rate I\_r, the concentration of Aaq and half saturation constant K\_Aaq are raised to the power of n.

Code Block 5. Subroutine BioHillEvaluate in reaction\_sandbox\_biohill.F90.

```
subroutine BioHillEvaluate(this, Residual, Jacobian, compute_derivative, &
                                  rt_auxvar, global_auxvar, material_auxvar, reaction, &
285
                                  option)
       . . .
      k max = 9.d-2
      k_decay = 1.d-6
      K_Aaq = 2.d-4
      K Baq = 1.25 d-5
290
      I\_Caq = 2.5 d-4
      yield = 1.d-4
      n = 1.2 d0
295
      stoichA = -1.d0
      stoichB = -0.25 d0
      stoichC = 0.33d0
      stoichD = 1.d0
300
      I_r = k_max * Xim * Aaq **n / (K_Aaq **n + Aaq **n) * &
                            Baq / (K_Baq + Baq) * \&
                            I_Caq / (I_Caq + Caq)
      I = I_r * volume
305
```



310



```
RateA = stoichA * I

...

RateX = yield * I - k_decay * Xim * volume

...

end subroutine BioHillEvaluate
```

Figure 4 illustrates a more accurate the simulation result for the example batch reaction where the Hill function exponent n is set to 1.2 in the Monod expression for  $A_{aq}$ . It is clear that the addition of the Hill function improves the model's ability to capture the tailing of species  $A_{aq}$  at low concentrations.



Figure 4. Improved match to hypothetical experimental results for species  $A_{aq}$  after incorporating the Hill function in Eq. (15). Compare log-scale plot to the solution without the Hill function in Fig. 3.

Results plotted in Fig. 4 may be replicated through the following commands:

```
cd $PFLOTRAN_DIR/regression_tests/default/reaction_sandbox
$PFLOTRAN_DIR/src/pflotran/pflotran -input_prefix biodegradation_hill
python biodegradation_vs_data.py biodegradation_hill-obs-0.pft
```

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The implementation of the enhanced biodegration reaction in Reaction Sandbox Biodegradation Hill is somewhat rigid. All reaction parameters (rate constants, half saturation constants, stoichiometries, etc.) are hardcoded within the source code as shown in Code Block 5 and may not be changed without code modifications. The implementation may be generalized, and this is demonstrated in Reaction Sandbox Flexible Biodegradation Hill.



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# 4.1.3 Reaction Sandbox Flexible Biodegradation Hill

Reaction Sandbox Flexible Biodegradation Hill employs the same biodegradation, growth and decays reactions as Reaction
Sandbox Biodegradation Hill with the added flexibility of specifying reaction parameters at run time through the input file, eliminating the need to re-compile PFLOTRAN every time a parameter changes. The new class is implemented in reac-tion\_sandbox\_flexbiohill.rype class extends reaction\_sandbox\_biohill\_type as presented in Code Block 6. The child class inherits the species integer IDs from the parent and adds member variables for storing all reaction parameters and a logical flag for specifying the units of half saturation constants K\_Aaq and K\_Baq and inhibitor concentration I\_Caq (molality versus molarity). The dynamic stoich array enables the use of do loops in the Evaluate routine. The class redirects the ReadInput, Setup, Evaluate and Destroy procedures to local implementations, though only the implementation of FlexBioHillEvaluate will be described below.

Code Block 6. reaction\_sandbox\_flexbiohill\_type in reaction\_sandbox\_flexbiohill.F90.

```
type, public, &
335
      extends(reaction_sandbox_biohill_type) :: reaction_sandbox_flexbiohill_type
      PetscReal :: k max
      PetscReal :: K_Aaq_n
      PetscReal :: K_Baq
      PetscReal :: I Caq
340
      PetscReal :: vield
      PetscReal :: k decay
      PetscReal :: n
      PetscBool :: molarity units
      PetscReal, pointer :: stoich(:)
345
      contains
      procedure, public :: ReadInput => FlexBioHillReadInput
      procedure, public :: Setup => FlexBioHillSetup
      procedure, public :: Evaluate => FlexBioHillEvaluate
      procedure, public :: Destroy => FlexBioHillDestroy
    end type reaction_sandbox_flexbiohill_type
350
```

The FlexBioHillEvaluate routine shown in Code Block 7 is more succint than BioHillEvaluate (Code Block 5). Reaction parameters are no longer hardwired, but read from the input file in FlexBioHillReadInput and stored as class member variables. The use of the class member stoich array within the do loops eliminates the need to hardwire Residual array indexing and reduces the number of lines of code. In addition, the full implementation within the source code demonstrates the ability to choose between molality versus molarity for aqueous concentrations and half saturation constants, a

14





useful option since laboratory data is often available in either format. There is also support for calculating analytical derivatives (for the Jacobian), which can be programmed much more concisely with the stoich array and do loops.

**Code Block 7.** FlexBioHillEvaluate in reaction\_sandbox\_flexbiohill.F90.

```
360
    subroutine FlexBioHillEvaluate(this, Residual, Jacobian, compute_derivative, &
                                    rt_auxvar, global_auxvar, material_auxvar, &
                                    reaction, option)
      . . .
      I_r = this%k_max * Xim * Aaq**this%n / (this%K_Aaq_n + Aaq**this%n) * &
365
                                Baq / (this%K_Baq + Baq) * &
                                this%I_Caq / (this%I_Caq + Caq)
      I = I_r * volume
      do icomp = 1, reaction%ncomp
370
        Residual(icomp) = Residual(icomp) - this%stoich(icomp) * I
      enddo
      Residual(Xim_offset) = Residual(Xim_offset) + this%k_decay * Xim * volume
    end subroutine FlexBioHillEvaluate
375
```

Flexible Biodegradation Hill produces results identical to Biodegradation Hill when given identical initial conditions and Biodegradation Hill's reaction parameters. These results may be compared by running the script compare\_biodegradation\_result after completing both simulations, i.e.

380

\$PFLOTRAN\_DIR/src/pflotran/pflotran -input\_prefix biodegradation\_hill
\$PFLOTRAN\_DIR/src/pflotran/pflotran -input\_prefix flexible\_biodegradation\_hill
python compare\_biodegradation\_results.py

cd \$PFLOTRAN DIR/regression tests/default/reaction sandbox

# 4.2 Reaction Sandbox Simple

385

Reaction Sandbox Simple provides a framework for evaluating reactive transport in 1D using common kinetic rate expressions with a preconfigured set of six aqueous ( $A_{aq}$ ,  $B_{aq}$ ,  $C_{aq}$ ,  $D_{aq}$ ,  $E_{aq}$ ,  $F_{aq}$ ) and two immobile species ( $X_{aq}$ ,  $Y_{aq}$ ). The conceptual model consists of a 100 meter, liquid-saturated column with a Dirichlet boundary condition at the inlet (x=0) and a zero-gradient boundary condition at the outlet (x=100). Grid spacing is set to 1 m resolution. The prescribed Darcy velocity is 1 m/y with a pore water velocity of 4 m/y (porosity = 0.25). Throughout the simulation, solutes enter at the inlet and react within the domain. Simulation results are stored in two formats: (1) snapshots of the entire domain at select times and (2) continuous





390 observation at a mid-column observation point at x=49.5 m. Table 4 summarizes simulation parameters, while Table 5 describes the initial and boundary concentrations.

**Table 4.** Reaction Sandbox Simple conceptual model.

Parameter	Value
Column length	100 m
Cross-sectional area	$1 \text{ m}^2$
Grid resolution	1 m
Prescribed Darcy velocity	1 m/y
Pore water velocity	4 m/y
Porosity	0.25
Tortuosity	1.0
Water density	$1000 \text{ kg/m}^3$
Aqueous diffusion coefficient	$10^{-9} \text{ m}^2$
Observation point location	49.5 m
Initial timestep size	1 h
Maximum timestep size	0.25 y
Final simulation time	25 у
Observation output frequency	Every timestep
Snapshot file output times	0, 6.25, 12.5, 18.75, 25 y

Table 5. Initial and boundary concentrations in the Reaction Sandbox Simple input file.

Species	Initial Concentration	Boundary Concentration	Units
$A_{aq}$	1.e-10	1.e-3	М
$B_{aq}$	1.e-10	1.e-3	М
$C_{aq}$	1.e-10	1.e-10	М
$D_{aq}$	1.e-10	1.e-10	Μ
$E_{aq}$	1.e-10	1.e-10	М
$F_{aq}$	1.e-10	1.e-10	М
$X_{im}$	1.e-4	N/A	mole/m <sup>3</sup> bulk volume
$Y_{im}$	1.e-10	N/A	mole/m <sup>3</sup> bulk volume

The implementation of the reaction\_sandbox\_simple\_type class is nearly identical to that of reaction\_sandbox\_biohill\_type in Code Block 3 with the addition of member variables <code>species\_Eaq\_id</code>, <code>species\_Faq\_id</code> and <code>species\_Yim\_id</code>. Member procedure <code>SimpleSetup</code> links these integer IDs to their respective names from the input file. The user may evaluate reactions in this sandbox with the following steps:





1. Uncomment a rate expression block within subroutine SimpleEvaluate.

cd \$PFLOTRAN\_DIR/src/pflotran

[emacs,gedit,nano,vi] reaction\_sandbox\_simple.F90

Remove "!uncomment:" prefixes from lines within chosen rate expression in subroutine SimpleEvaluate.

400

2. Compile the PFLOTRAN executable.

Save the file.

make pflotran

3. Navigate to \$PFLOTRAN\_DIR/regression\_tests/default/reaction\_sandbox

cd \$PFLOTRAN\_DIR/regression\_tests/default/reaction\_sandbox

405 4. Change the x-direction grid resolution in reaction\_sandbox\_simple.in to 100.

[emacs,gedit,nano,vi] reaction\_sandbox\_simple.in

NXYZ 10 1 1  $\rightarrow$  NXYZ 100 1 1

Save the file.

5. Run the simulation.

```
410 $PFLOTRAN_DIR/src/pflotran/pflotran -input_prefix reaction_sandbox_simple
```

subroutine SimpleEvaluate (...)

6. Plot the results with reaction\_sandbox\_simple.py.

python reaction\_sandbox\_simple.py

Code Block 8 illustrates a *commented* rate expression block for calculating the first order decay of species Aaq to daughter product Caq.

415 Code Block 8. Commented first-order rate expression block in SimpleEvaluate within reaction\_sandbox\_simple.F90.

```
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```





```
425 !uncomment: RateA = stoichA * Rate
425 !uncomment: RateC = stoichC * Rate
...
end subroutine SimpleEvaluate
```

The user deletes all "!uncomment:" prefixes in the codeblock as shown in Code Block 9, compiles PFLOTRAN, navigates 430 to the reaction\_sandbox folder, and runs the code.

Code Block 9. Uncommented first-order rate expression block in SimpleEvaluate within reaction\_sandbox\_simple.F90.

```
subroutine SimpleEvaluate (...)
...
! first-order (A -> C)
435
...
k = 1.26839d-9 ! [1/sec]
stoichA = -1.d0
stoichC = 1.d0
Rate = k * Aaq * L_water ! [mol/sec]
440
RateA = stoichA * Rate
RateC = stoichC * Rate
...
end subroutine SimpleEvaluate
```

Figure 5 illustrates the simulation results. Plotted to the left is concentration breakthrough at the observation point, and to the right is a snapshot of concentration profiles at 12.5 years. Species  $A_{aq}$  clearly decays into  $C_{aq}$  while the other species are transported without reaction. The user may evaluate the other reaction rate expressions in the subroutine with the same approach.

# 4.3 Reaction Sandbox Example

- 450 PFLOTRAN's reaction\_sandbox\_example.F90 serves as a template for implementing new Reaction Sandboxes. The sandbox implements the first-order decay of  $A_{aq}$  without daughter products, and the developer modifies the templated source code to incorporate new reactions. Comments within the source code enumerate steps for implementing new reactions beginning with the renaming of subroutines and variables and ending with the implementation of subroutine ExampleDestroy at the bottom of the file. In between, source code is modified to implement the new reaction(s).
- 455 Code Block 10 illustrates the first two steps embedded within comments in the source code near the top of the file.

Code Block 10. Source code with embedded comments from the top of reaction\_sandbox\_example.F90. module Reaction\_Sandbox\_Example\_class





#### **Reaction Sandbox Simple**





```
#include "petsc/finclude/petscsys.h"
      use petscsys
460
    ! 1. Change all references to "Example" as desired to rename the module and
         subroutines within the module.
    !
465
      use Reaction_Sandbox_Base_class
      use Global_Aux_module
      use Reactive_Transport_Aux_module
470
      use PFLOTRAN_Constants_module
      implicit none
      private
475
      2. Add module variables here. Note that one must use the PETSc data types
    !
         PetscInt, PetscReal, PetscBool to declare variables of type integer
    !
```





```
! float/real*8, and logical respectively. E.g.
!
480 ! PetscReal, parameter :: formula_weight_of_water = 18.01534d0
...
```

Comments within ExampleEvaluate provide a detailed description of the subroutine's arguments and local variables, including members of the reaction and reactive transport auxiliary variable classes (i.e. reaction\_rt\_type and reac-485 tive\_transport\_auxvar\_type, respectively) which may be used in rate expression calculations. The units of all variables and the Residual and Jacobian arrays are also provided. Code Block 11 shows several of these detailed comment blocks.

Code Block 11. Representative comments from subroutine ExampleEvaluate.

```
. . .
490
      ! rt_auxvar - Object holding chemistry information (e.g. concentrations,
          activity coefficients, mineral volume fractions, etc.).
      !
                                                                      See
      !
          reactive_transport_aux.F90.
      !
      !
          Useful variables:
495
      !
            rt_auxvar%total(:, iphase) - total component concentrations
                                          [mol/L water] for phase
      !
            rt auxvar%pri molal(:) - free ion concentrations [mol/kg water]
      !
            rt_auxvar%pri_act_coef(:) - activity coefficients for primary species
      1
      !
            rt_auxvar%aqueous%dtotal(:, iphase) - derivative of total component
      !
                         concentration with respect to free ion [kg water/L water]
500
      . . .
    ! 10. Add code for residual evaluation
      ! Unit of the residual must be in moles/second
505
      ! global_auxvar%sat(iphase) = saturation of cell
      ! 1.d3 converts m<sup>3</sup> water -> L water
      L_water = material_auxvar%porosity*global_auxvar%sat(iphase)* &
                 material_auxvar%volume *1.d3
      ! always subtract contribution from residual
510
      Residual(this%species_id) = Residual(this%species_id) - &
        (-1.d0) * & ! negative stoichiometry
        this%rate_constant * & ! 1/sec
```



515



```
L_water * & ! L water

! rt_auxvar%total(this%species_id, iphase) = species total component

! concentration

rt_auxvar%total(this%species_id, iphase) ! mol/L water

....
```

Once refactoring is complete, the developer must modify the corresponding class and subroutine names within reaction\_sandbox.F90 to match those in the newly refactored reaction\_sandbox\_example.F90. The source code may then be compiled without changing the reaction\_sandbox\_example.F90 filename. Should this filename be revised, the developer must update the corresonding filenames within the pflotran\_object\_files.txt and pflotran\_dependencies.txt files referenced by the makefile.

The best approach to compiling the code is to perform a make clean to remove all previously built modules and object 525 files and the pflotran executable, and then, make pflotran to compile the code. reaction\_sandbox\_example.in located in \$PFLOTRAN\_DIR/regression\_tests/default/reaction\_sandbox provides a representative input deck for this example. Note that the card EXAMPLE in the REACTION\_SANDBOX block of this input file must be updated to match the corresponding keyword added to reaction\_sandbox.F90.

# 5 Conclusions

530 Customization of biogeochemical reaction networks is often necessary in the development of reactive transport simulators employed to simulate problem-specific scenarios in the real world. For researchers in the natural sciences who are focused more on the biology, chemistry, and physics than computational science, modifying these codes to incorporate new scientific processes can be challenging. The PFLOTRAN Reaction Sandbox may help remedy this issue.

The Reaction Sandbox provides a modular environment for prototyping new kinetic rate expressions that do not exist within 535 PFLOTRAN. Within the Reaction Sandbox, novel reaction networks can evolve and mature over time; natural selection can run its course. Once vetted, these reactions may be incorporated more efficiently elsewhere within the permanent code base.

This work demonstrates the implementation of reactions within the Reaction Sandbox. Several new Reaction Sandbox classes are conceptualized and implemented based on existing rate expressions within PFLOTRAN for microbially-mediated biodegradation to improve the code's ability to match a hypothetical, empirical data and provide greater flexibility from the end

540 user perspective. Reaction Sandbox Simple is presented as a means of prototyping numerous common kinetic rate expressions within a preconfigured column experiment context. Reaction Sandbox Example provides a template for implementing and incorporating new reactions within PFLOTRAN.

*Code availability.* The source code, input files and results presented in this manuscript are based on commit 70beb16 of the PFLOTRAN Git repository. A snapshot of this commit is available at https://doi.org/10.5281/zenodo.5507599. The corresponding version of PETSc





545 is v3.13 and was configured on Ubuntu 20.04 with GCC 9.3 using the following config script: ./configure -CFLAGS='-O3' -CXXFLAGS='-O3' -FFLAGS='-O3' -with-debugging=no -download-mpich=yes -download-hdf5=yes -downloadfblaslapack=yes -download-metis=yes -download-parmetis=yes.

# **Appendix A: Reaction Sandbox Member Procedure Interfaces**

This section documents Fortran class reaction\_sandbox\_base\_type member procedure interfaces not covered in Section 3.2.1 550 (see reaction\_sandbox\_base.F90).

# A1 ReadInput => BaseReadInput

ReadInput provides a customizable interface for reading parameters associated with the reaction sandbox from a block in the input file. The input block is opened by a unique keyword associated with the child Reaction Sandbox class.

Code Block 12. BaseReadInput argument list.

```
555
      subroutine BaseReadInput(this, input, option)
        class(reaction_sandbox_base_type) :: this
        type(input_type), pointer :: input
        type(option_type) :: option
```

560

this: the reaction\_sandbox\_base\_type object

input : an object storing the input file object pointer and line buffers

option: an object storing run time options including process rank and an error messaging buffer

# A2 Setup => BaseSetup

Setup initializes the Reaction Sandbox class by allocating dynamic memory, mapping species IDs, assigning stoichiometries 565 and rate constants, etc. The degree to which a Reaction Sandbox class settings are customizable at run time is up to the developer and his or her creativity in implementing ReadInput and Setup.

Code Block 13. BaseSetup argument list.

```
subroutine BaseSetup(this, reaction, option)
  . . .
```

570

```
class(reaction_sandbox_base_type) :: this
class(reaction_rt_type) :: reaction
type(option_type) :: option
```





575 this: the reaction\_sandbox\_base\_type object

reaction : an object storing chemical species and general reaction information

option: an object storing run time options including process rank and an error messaging buffer

# A3 Evaluate => BaseEvaluate

Evaluate calculates kinetic rates for all reactions in the Reaction Sandbox class and adds the (kinetic rate) contributions to the residual and Jacobian arrays. This is the only subroutine that must be extended in child Reaction Sandbox classes.

```
Code Block 14. BaseEvaluate argument list.
```

	subroutine BaseEvaluate(this, Residual, Jacobian, compute_derivative, &
	rt_auxvar , global_auxvar , material_auxvar , &
	reaction, option)
585	
	<b>class</b> (reaction_sandbox_base_type) :: this
	<b>class</b> (reaction_rt_type) :: reaction
	! the following arrays must be declared after reaction
	PetscReal :: Residual(reaction%ncomp)
590	PetscReal :: Jacobian (reaction%ncomp, reaction%ncomp)
	PetscBool :: compute_derivative
	<b>type</b> (reactive_transport_auxvar_type) :: rt_auxvar
	<b>type</b> (global_auxvar_type) :: global_auxvar
	<b>class</b> (material_auxvar_type) :: material_auxvar
595	type(option_type) :: option

this: the reaction\_sandbox\_base\_type object

Residual: a 1D array of double precision numbers holding contributions to the residual equations at each grid cell

Jacobian : a 2D array of double precision numbers holding contributions to the Jacobian matrix at each grid cell

600 compute\_derivative : a logical flag toggling on the calculation of analytical derivatives for the Jacobian matrix when true

```
rt_auxvar: an object storing reactive transport state variables (e.g. concentrations, rates, etc.) at each grid cell
```

global\_auxvar : an object storing flow state variables (e.g. density, saturation, etc.) at each grid cell

material\_auxvar: an object storing material and cell properties (e.g. porosity, volume, etc.) at each grid cell





605 reaction: an object storing chemical species and general reaction information

option: an object storing run time options including process rank and an error messaging buffer

# A4 UpdateKineticState => BaseUpdateKineticState

UpdateKineticState updates state variables associated with the Reaction Sandbox class that are stored in rt auxvar and updated at the end of a time step based on rates calculated in the Reaction Sandbox (e.g. for mass balance calculations, updates to mineral volume fractions, etc.). 610

**Code Block 15.** BaseUpdateKineticState argument list.

```
subroutine BaseUpdateKineticState(this,rt_auxvar,global_auxvar, &
                                         material_auxvar, reaction, option)
         . . .
615
        class(reaction_sandbox_base_type) :: this
        type(reactive_transport_auxvar_type) :: rt_auxvar
        type(global_auxvar_type) :: global_auxvar
        class(material_auxvar_type) :: material_auxvar
        class(reaction_rt_type) :: reaction
        type(option_type) :: option
```

620

this: the reaction\_sandbox\_base\_type object

rt\_auxvar: an object storing reactive transport state variables (e.g. concentrations, rates, etc.) at each grid cell

global\_auxvar : an object storing flow state variables (e.g. density, saturation, etc.) at each grid cell

625 material\_auxvar: an object storing material and cell properties (e.g. porosity, volume, etc.) at each grid cell

reaction : an object storing chemical species and general reaction information

option: an object storing run time options including process rank and an error messaging buffer

# A5 AuxiliaryPlotVariables => BaseAuxiliaryPlotVariables

AuxiliaryPlotVariables appends Reaction Sandbox-specific state variables stored in rt\_auxvar to the list of output variables to be printed to observation and snapshot files. 630

**Code Block 16.** BaseAuxiliaryPlotVariables argument list.

```
subroutine BaseAuxiliaryPlotVariables(this, list, reaction, option)
```

. . .



635



```
class(reaction_sandbox_base_type) :: this
type(output_variable_list_type), pointer :: list
class(reaction_rt_type) :: reaction
type(option_type) :: option
```

this : the reaction\_sandbox\_base\_type object

640 output\_variable\_list\_type: a linked list of output variables to which new sandbox-specific output\_variable\_type objects will be added

reaction : an object storing chemical species and general reaction information

option : an object storing run time options including process rank and an error messaging buffer

# A6 Destroy => BaseDestroy

645 Destroy deallocates all dynamic memory in the Reaction Sandbox class at the end of a simulation.

```
Code Block 17. BaseDestroy argument list.
```

subroutine BaseDestroy(this)
...
class(reaction\_sandbox\_base\_type) :: this

650

this: the reaction\_sandbox\_base\_type object

Author contributions. GH implemented the Reaction Sandbox in PFLOTRAN, designed and implemented the example sandboxes, and prepared the manuscript.

Competing interests. The author declares that he has no conflicts of interest.

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