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# **NRAP-Open-IAM User's Guide**

***Release alpha 2.8.1-23.12.15***

**NRAP-Open-IAM Development Team**

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## OBTAINING NRAP-OPEN-IAM



### 1.1 Introduction

NRAP-Open-IAM is an open-source Integrated Assessment Model (IAM) for Phases II and III of the National Risk Assessment Partnership (NRAP). The goal of this software is to go beyond risk assessment into risk management and containment assurance. NRAP-Open-IAM is currently in active development and is available for testing and feedback only.

As this is a prototype of software being actively developed, we are seeking any feedback or bug reports. Feedback can be emailed to the NRAP-Open-IAM project at [NRAP@netl.doe.gov](mailto:NRAP@netl.doe.gov), or the current development team lead at [Nathaniel.Mitchell@netl.doe.gov](mailto:Nathaniel.Mitchell@netl.doe.gov), or any other member of the development team.

Issues can also be reported on GitLab issues page for NRAP-Open-IAM: [https://gitlab.com/NRAP/OpenIAM/-/issues?sort=created\\_date&state=opened](https://gitlab.com/NRAP/OpenIAM/-/issues?sort=created_date&state=opened)

If you have been given access to the code indirectly and would like to be notified when updates are available for testing, please contact the development team to be added to our email list.

### 1.2 Downloading NRAP-Open-IAM

NRAP-Open-IAM tool and examples can be downloaded from a public GitLab repository located at <https://gitlab.com/NRAP/OpenIAM>. If the NRAP-Open-IAM was downloaded from the GitLab repository, the folder name may have the repository's current hash appended to it. Feel free to rename the folder to something simple like *NRAPOpenIAM* to simplify the navigation.

In addition to that, the copy of the tool can be obtained through NETL's Energy Data eXchange website: <https://edx.netl.doe.gov/dataset/phase-iii-nrap-open-iam> by requesting an access through e-mail addressed to [NRAP@netl.doe.gov](mailto:NRAP@netl.doe.gov). The NRAP-Open-IAM is distributed as a zip file that can be extracted in the location specified by user.

## 1.3 Installing NRAP-Open-IAM

The NRAP-Open-IAM requires Python version 3.9 or greater to operate. If you need to install Python, we describe all steps of the installation process below.

### General Installation Guide:

- Extract the tool files from the provided/downloaded zip.
- Navigate to the *installers* folder within the recently unzipped directory.
- Navigate to the folder corresponding to the operating system that you are utilizing.
- Follow instructions file located in the folder for your operating system.

For Windows: The file *Installation\_Instructions\_Windows.txt* describes steps required to install needed Python packages for the proper work of NRAP-Open-IAM.

For macOS: The file *Installation\_Instructions\_macOS.txt* describes steps user needs to follow in order to install required Python packages.

For Linux OS: Linux users are assumed to know the installation commands for their specific version of Linux needed to install required tools. The file *Installation\_Instructions\_Linux.txt* specifies the needed software and package dependencies.

For alternative installation of Python the following packages are needed: NumPy, SciPy, PyYAML, Matplotlib, Pandas, TensorFlow (of version 2.6), Keras, scikit-learn, Pmw, pip, and six. In most cases (mainly dependent on the platform and Python distribution) the required libraries can be installed using pip or conda package managers. Additional libraries recommended to run Jupyter notebooks and scripts illustrating work of NRAP-Open-IAM are IPython and Jupyter.

On macOS and Linux machines the gfortran compiler needs to be present/installed to compile some of the NRAP-Open-IAM code (macOS users can find gfortran here: (<https://gcc.gnu.org/wiki/GFortranBinariesMacOS>)).

After the proper version of Python is installed, the NRAP-Open-IAM can be set up and tested. **Note: If Python was installed through Anaconda please use Anaconda prompt instead of command prompt for setup and tests.** In the NRAP-Open-IAM distribution folder find and open the sub-folder *setup*. Next, open a command prompt/Anaconda prompt in the *setup* folder (on Windows, this can be done by holding Shift and right clicking inside the folder when no file is selected, then selecting **Open command window here**; alternatively, one can navigate to the folder *setup*, type `cmd` in the address bar of the file browser and hit Enter to open the command prompt there). (On Windows, Anaconda prompt can be found in the programs menu under submenu **Anaconda3 (64-bit)**.) Run the setup script by entering the command:

```
python openiam_setup_tests.py
```

in the command prompt/Anaconda prompt. This will test the version of Python installed on the system. Next the setup script will test the versions of several Python libraries that the NRAP-Open-IAM depends on. The setup script will compile several Fortran libraries needed for some component models on Mac and Linux. Users of Windows OS will be provided with the compiled libraries. Finally, the setup script will run the test suite to see if the NRAP-Open-IAM has been installed correctly. If the results printed to the console indicate errors during the testing the errors have to be resolved before the NRAP-Open-IAM can be used. When contacting the developers to resolve problems please include all output from the setup script or test suite runs.

## 1.4 Testing installation

After setup the test suite can be run again by entering the NRAP-Open-IAM *test* directory in a terminal and typing:

```
python iam_test.py
```

Test results will be printed to the terminal. The setup script run during the installation process uses the same test suite after testing whether the necessary Python libraries are installed, and compiling the NRAP-Open-IAM libraries.

## 1.5 Contributors

During the Phase II and/or Phase III of the NRAP the following researchers contributed to the development of NRAP-Open-IAM (listed in alphabetical order with affiliation at the time of active contribution):

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## GETTING STARTED

This document provides guidance for the use of NRAP-Open-IAM. The document is quite large, but each section is bookmarked. Utilizing these bookmarks will aid with the navigation of this document.

The NRAP-Open-IAM has several ways for a user to build and run simulations, including a graphical user interface (GUI), text based control files (the control file interface, CFI), and python scripts. The simplest way to build and run simulations for NRAP-Open-IAM is the GUI. To launch the GUI, open a command prompt in the *source/GUI* directory and type:

```
python NRAP_OPENIAM.py
```

## 2.1 Conceptual Model and Overview

NRAP-Open-IAM is designed for simulating and analyzing the behavior of geologic carbon storage (GCS) system models. These system models can include representations of reservoir response to CO<sub>2</sub> injection, the resulting impacts on potential leakage pathways like wellbores, and any contaminant plumes that form in aquifers or the atmosphere due to brine or CO<sub>2</sub> leakage. Each part of this process (reservoir response, leakage pathway behavior, and impacts on the receptor of leakage) is represented by a particular component within the system model. Additionally, NRAP-Open-IAM can automate specific analyses, such as determining the area of review for a GCS site. The intended purpose of NRAP-Open-IAM is to provide quantitative metrics that aid in GCS project planning, permitting, and operational decision support. NRAP-Open-IAM can be run through the graphical use interface (GUI; section [GUI Operation](#)), the control file interface (section [Control File Interface](#)), or a script-based approach in Python.

### 2.1.1 System Model Design

Within NRAP-Open-IAM, the system model contains components that are connected to each other. For example, a reservoir component can provide pressures and CO<sub>2</sub> saturations to a wellbore component. The wellbore component then uses these inputs to calculate CO<sub>2</sub> and brine leakage rates to a specific aquifer. In its turn, an aquifer component can use these leakage rates to model the evolution of pH and TDS plumes within the aquifer. An aquifer component is a receptor type of component, which can also be represented by an atmosphere component. For example, a wellbore component can provide leakage rates to the atmosphere, and an atmosphere component can then be used to model the evolution of atmospheric contaminant plumes. [Fig. 2.1](#) is a conceptual model demonstrating the connection of reservoir components, wellbore components, and a receptor (aquifer or atmosphere components). Almost all NRAP-Open-IAM system models include a stratigraphy component. A stratigraphy component can be used to define the thicknesses and depths of the storage reservoir, aquifers, and aquitards in the model domain. Many NRAP-Open-IAM components are designed to link with the stratigraphy component in a way that automatically sets component parameters related to the stratigraphy.

The setup demonstrated in [Fig. 2.1](#) is the most common design of an NRAP-Open-IAM system model. Other designs not fitting the illustrated scheme are possible as well. For example, some of the components producing leakage rates

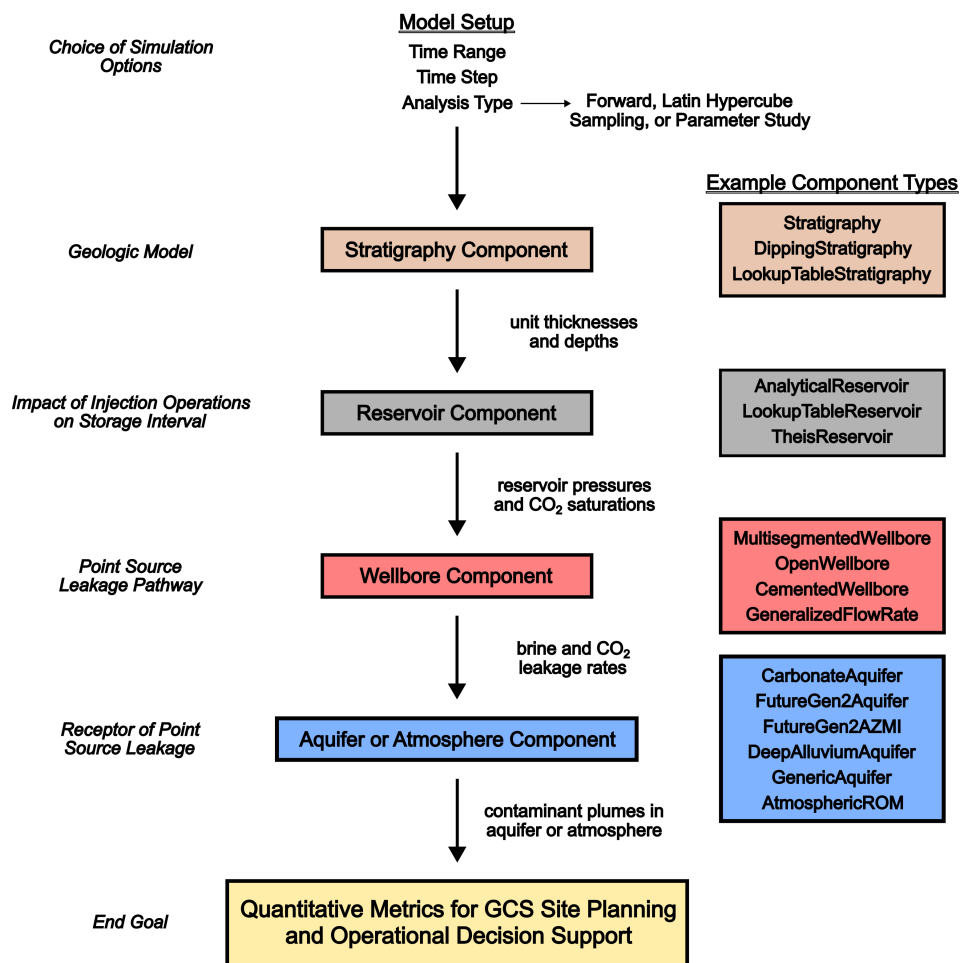


Fig. 2.1: Standard System Model Design

are not designed to be linked to an aquifer component that accepts leakage from a point source. The SealHorizon component can use pressures and CO<sub>2</sub> saturations from a reservoir component to model brine and CO<sub>2</sub> leakage rates through a fractured caprock overlying the storage reservoir, but the predicted leakage rates occur over an area of the caprock/reservoir interface. The aquifer components shown in Fig. 2.1 are designed to receive leakage rates from a point source, like a wellbore, and therefore should not be used with rates that apply over an area or a linear element (e.g., faults in the FaultFlow component). Such components can still be used to evaluate leakage risks, however, and Fig. 2.2 shows a conceptual model demonstrating the connection of a reservoir component with a non-local leakage pathway component (i.e., not a point source leakage pathway). Aquifer components accepting leakage rates that apply over areas or linear elements may be developed in the future.

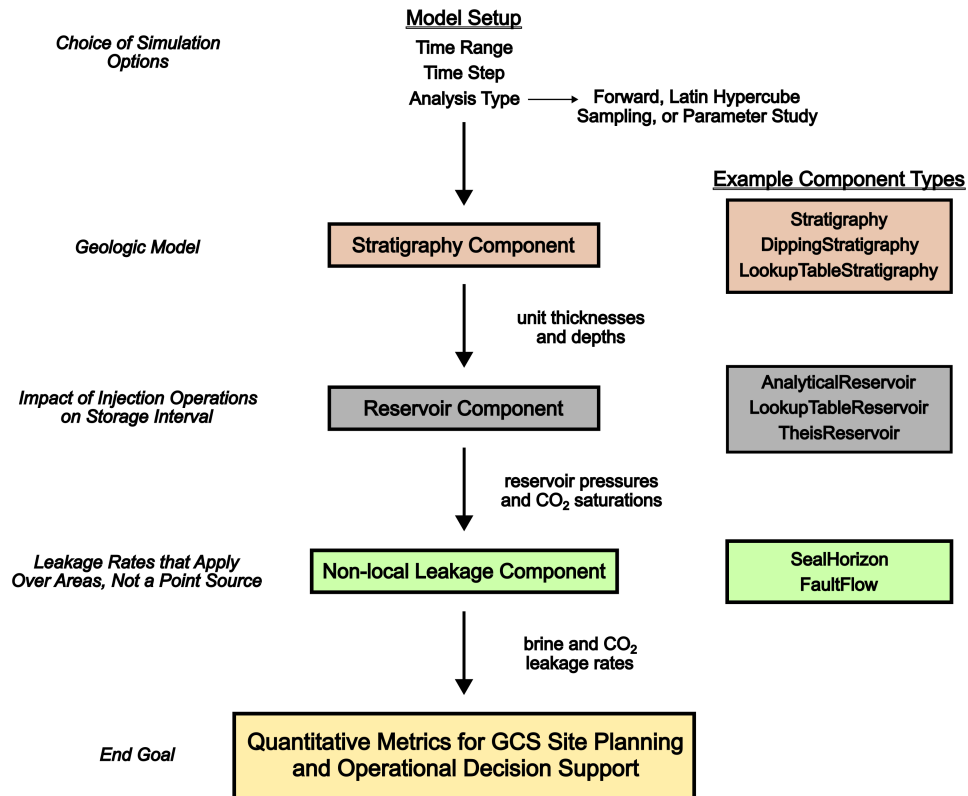


Fig. 2.2: Non-local Leakage System Model Design

NRAP-Open-IAM has several components that are not designed to be connected to other components. For example, the PlumeStability component does not connect to the stratigraphy or leakage pathway components. Instead, it uses .csv files containing reservoir conditions over time to evaluate the development and movement of plumes within the reservoir (e.g., plume areas, velocity of the plume centroid, and the direction in which the dispersion of the plume occurs). The ChemicalWellSealing component, which evaluates if and when a fracture will seal due to calcite precipitation, also does not connect with other components. This type of system model design is demonstrated in Fig. 2.3.

NRAP-Open-IAM is designed to accept a variety of component designs, so it can be used to create many different system model designs. Note that the same type of .csv files used by PlumeStability components can also be used to read reservoir conditions into a system model with LookupTableReservoir components (e.g., driving wellbore leakage components with output from high-fidelity reservoir simulations).

The support of one GCS site may require the use of multiple system models. For example, the ChemicalWellSealing component (Fig. 2.3) can help inform which wells are likely to self-seal and, therefore, might be excluded from being considered as a possible leakage pathway in a larger GCS system model (Fig. 2.1). The ChemicalWellSealing

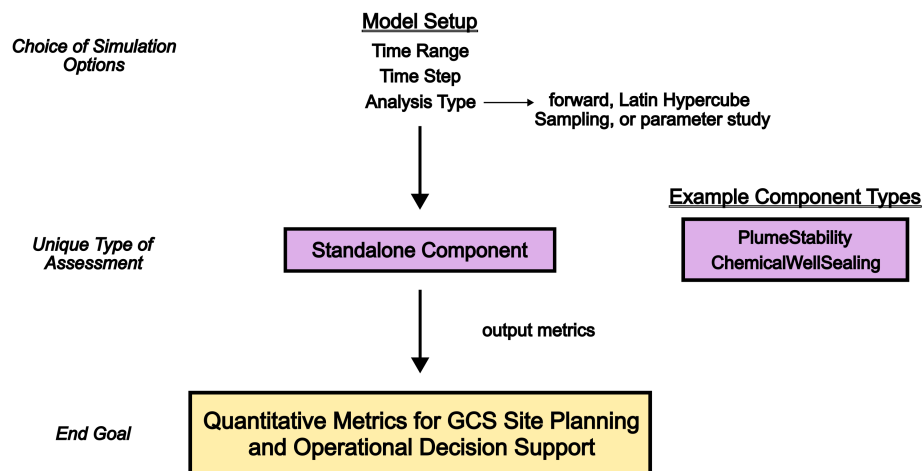


Fig. 2.3: Standalone Component System Model Design

component has a parameter related to reservoir pressure, so one should first constrain this parameter by running a reservoir simulation (either through a separate program or through an NRAP-Open-IAM simulation where the reservoir component is the final component in the chain). In this case, one might use three GCS system models (1) to constrain reservoir pressures, (2) to determine whether particular wells will self-seal (Fig. 2.3), and (3) to model the behavior of the whole GCS site (excluding wells that will self-seal; Fig. 2.1).

## 2.1.2 Component Parameters and Applications

Each component model has a variety of parameters, and these parameters have different limits or ranges. When setting up a simulation, any component parameter that is not given a value by the user will automatically be set to the parameter's default value.

Different components are meant to be used in different applications. For example, the `TheisReservoir` component was designed to handle multiple (brine) injection and/or extraction wells over a large area, with the pressures produced reflecting the interaction of multiple wells. In contrast, the `AnalyticalReservoir` component can simulate only one injection well. The `FutureGen2Aquifer` and `FutureGen2AZMI` components were designed to represent the specific geochemistry of the Future Gen site but may be applicable to similar aquifers with depths between 100 m and 700 m (`FutureGen2Aquifer`) and 700 m and 1600 m (`FutureGen2AZMI`). The `OpenWellbore` component is useful for evaluating worst-case scenarios where leakage through an unplugged wellbore enters an aquifer. In contrast, a `MultisegmentedWellbore` component simulates flow through impaired cement and can account for leakage into multiple aquifers overlying the injection zone. More details regarding the distinctions between different wellbore components and their intended applications are available in chapter [Component Comparison](#). Descriptions of NRAP-Open-IAM components and their parameters are provided in chapter [Components Description](#) below.

### 2.1.3 Analysis Types

NRAP-Open-IAM simulations can use one of three analysis types: Forward, Latin Hypercube Sampling (LHS), or parameter study (Parstudy) (Fig. 2.1 - Fig. 2.3). A Forward analysis (forward in the control file interface) is deterministic. Each parameter is fixed at a certain value, so the simulation has the same results each time it is run.

LHS simulations (lhs in the control file interface) are stochastic: the parameters are varied between minimum and maximum values. An LHS simulation consists of many separate realizations, where each realization has different parameter values. This approach helps to address parameter uncertainty. For example, by constraining parameter values within reasonable bounds, one can focus on the proportion of LHS realizations in which leakage occurs. This proportion can be used to estimate the likelihood of such a leakage event. Leakage occurring in 10% of realizations could be considered to correspond to a 10% probability of leakage, given the user's current understanding of the parameter values. Latin Hypercube Sampling is different from a purely random approach in that it evenly samples the range of values for each stochastic parameter, and runs the model for different combinations of parameter values. A purely random sampling of parameter values could fail to explore a large section of the parameter space. LHS simulations divide each parameter's range into subranges; within each subrange, a parameter value is randomly selected. Because LHS simulations need to evenly sample the parameter space through the assessment of multiple subranges, each LHS simulation requires a minimum number of realizations. If one attempts to run an LHS simulation without a sufficient number of realizations, the code will raise a `LinAlgError` saying "Matrix is not positive definite." If this error occurs, increase the total number of realizations. Additionally, if you attempt to run an LHS simulation without including any parameters that vary stochastically (i.e., by giving the parameter minimum and maximum limits), then the simulation will encounter an error. In this case, the simulation should instead use a Forward analysis type.

The Parstudy analysis type (parstudy in the control file interface) also evaluates different realizations of each simulation, where each realization has different parameter values. The Parstudy analysis type also divides a variable parameter's specified range into separate subranges and then selects a value from each subrange. While an LHS simulation has the user to specify how many realizations to evaluate, a Parstudy simulation has the user to specify how many parameter values to use for each stochastic parameter (i.e., number of subranges within the overall range). As a result, the number of realizations for Parstudy simulations increases exponentially with the number of stochastic parameters.

Overall, the Forward analysis type is intended to be used for decision-support GCS system models that are constrained in scope; in some cases, a deterministic simulation can demonstrate a finding in a more clear and concise manner. The LHS analysis type is intended to be used for decision-support GCS system models that address the uncertainty in parameter values. There can be substantial uncertainty in parameters like permeability, and producing results that effectively support an argument (e.g., that leakage risks are minimal at a site) can require the acknowledgement and representation of that uncertainty in GCS system models. Finally, the Parstudy analysis type is intended to be used to study the effects of certain parameters on model outputs (i.e., sensitivity analysis). Studying the effects of certain parameters is important for decision support. Understanding which parameters have the most significant impact can help inform the user about the parameters they should focus on constraining in their study area. The control file interface can also be used to automate sensitivity analysis within LHS simulations; see *ControlFile\_ex8a-ControlFile\_ex8d*.

### 2.1.4 Visualization Options

NRAP-Open-IAM includes a variety of visualization options. For example, there are multiple options for plotting a model domain's stratigraphy with 2-dimensional and 3-dimensional plots. The 3-dimensional Stratigraphy plot type can display features like injection sites and wellbores. Certain stratigraphy components can create spatial variations in unit thicknesses and depths, and the stratigraphy visualization options are provided to help users ensure that the model domain conforms to the intended design. Other plot types available with NRAP-Open-IAM include: time series of outputs; map-view figures showing the extent of reservoir and aquifers impacts (meant to inform an operation's area of review (AoR)); map-view figures showing when monitoring wells at specified locations can detect contaminant plumes in aquifers (time to first detection, TTFD); and map-view figures showing the extent of atmospheric CO<sub>2</sub> plumes.

Many of the plot types are not available through the post-processing stage of the GUI (section *GUI Operation*); this part of the GUI can only generate time series plots and maps of atmospheric CO<sub>2</sub> plumes. When using a workflow

(section *Workflows in NRAP-Open-IAM*) in the GUI, however, the workflow will automatically save figures to the output directory. For example, the AoR workflow will save AoR plots, while the TTFD workflow will save TTFD plots. The control file interface and script applications have access to all visualization options in NRAP-Open-IAM (see, e.g., section *Visualization Options in the CFI*). Simulation results are also saved in .csv files, however, and these files can be used to create figures in separate programs.

## 2.1.5 Workflow Types

NRAP-Open-IAM simulations can be set up manually, where the user selects every component and sets up the connections between components (e.g., a reservoir component providing pressures and CO<sub>2</sub> saturations to a wellbore component). Alternatively, the user can also set up a simulation by selecting a workflow type (section *Workflows in NRAP-Open-IAM*). A workflow considers a specific type of analysis (e.g., determining an area of review) and automatically configures aspects of the system model for that analysis type. For example, the AoR workflow requires reservoir, wellbore, and aquifer components that are connected and produce certain outputs. The AoR workflow automatically sets up the component connections and output types, preventing the errors that can occur if the system model and components are not set up correctly. A workflow generally serves as a blueprint for a system model (Fig. 2.1), allowing the user to perform a specific analysis more quickly and easily. Workflows are available in the control file interface and GUI.

## 2.2 GUI Operation

When the graphical user interface (GUI) is first opened, a disclaimer screen will be shown followed by the main interface.

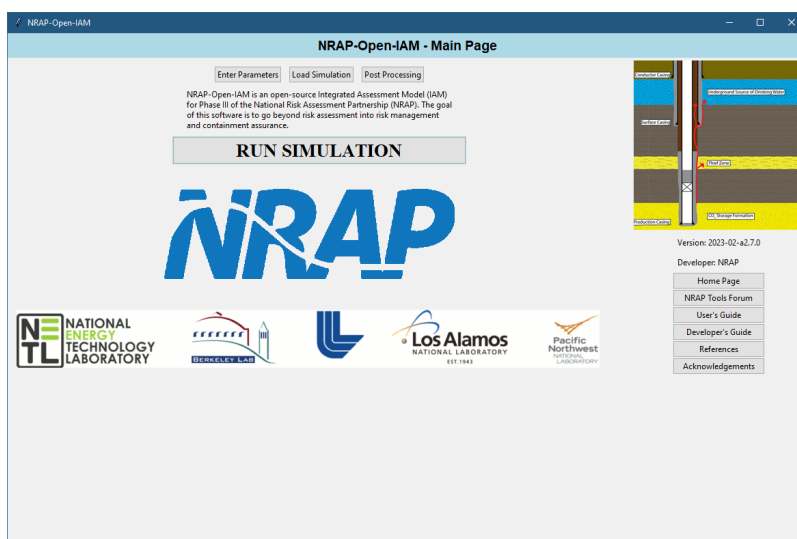


Fig. 2.4: Main NRAP-Open-IAM Interface

To begin building a model, click on the **Enter Parameters** button. The process of building a model consists of entering basic model parameters, defining the stratigraphy of the site, and then adding each component to be included in the system model. Therefore, the first tab that a user would see after clicking the **Enter Parameters** button is the model parameters view.

Start by defining a **Simulation** name for the model: the name also will be used as the name of the file containing details of the NRAP-Open-IAM simulation. Time should be entered in years. The **End** time is the total number of years that will be considered in simulation. The **Time step** entry specifies the uniform time step taken during the

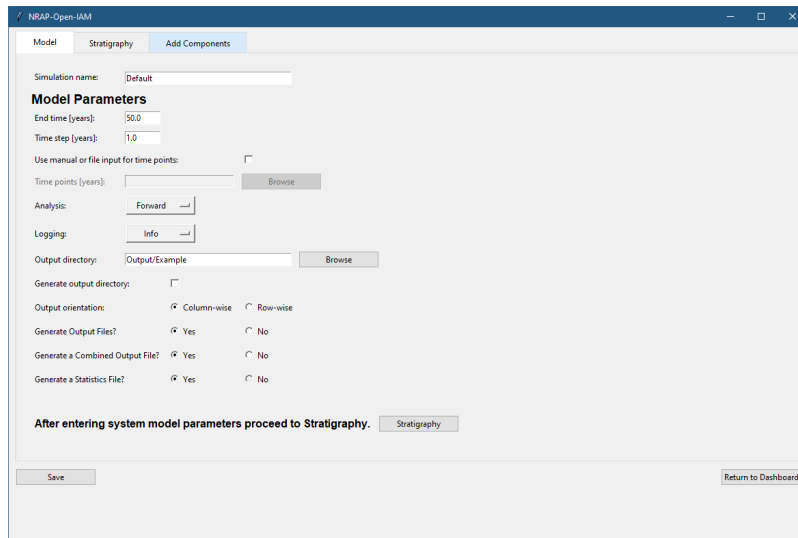


Fig. 2.5: Model Parameters View

simulation (typically 1 year time steps are used). NRAP-Open-IAM can run simulations with three different analysis types: Forward, LHS, and Parstudy. The Forward analysis type runs a single deterministic scenario, where parameter values are constant and the results will be the same each time the simulation is run. LHS (abbreviation for Latin Hypercube Sampling) is the analysis type used for stochastic simulations with random variations in parameter values. The Parstudy analysis type (short for parameter study analysis) evaluates a user-defined number of random values for each stochastic parameter. See section [Conceptual Model and Overview](#) for more details regarding these analysis types.

NRAP-Open-IAM creates a log file for each simulation: the level of information being logged can be set by the Logging entry. In general, the default level of Info would contain the most useful messages. A Debug (debugging) level of Logging will contain more information about component model connections, setup and calls, but will produce very large files and should be avoided for large simulations. Warn (warning) and Error levels can be used if log file sizes become an issue; only more important messages will be logged.

NRAP-Open-IAM will save all the simulation results to the specified Output directory. In the text field corresponding to Output directory, the user needs to enter a path to the folder where the output will be saved. In the case the entered path does not exist, the empty directory will be created if the Generate output directory box is checked. Additionally, if the provided path is not absolute, it is assumed that it starts in the NRAP-Open-IAM root folder. A {datetime} stamp can be added to the folder name so that each run of a particular simulation will be saved separately, otherwise results from a previous run will be overwritten by subsequent runs until the output folder is changed. After setting up the model parameters, proceed to the Stratigraphy tab.

In the Stratigraphy tab, model parameters related to the stratigraphy of the geologic carbon storage site are defined. All coordinate systems are assumed to have units of meters and are defined by the reservoir component used. Model parameters for the stratigraphy and appropriate components are defined by either assigning a fixed value or random distribution to vary over. For the LHS analysis type, parameters defined with a distribution will be sampled from that distribution. For the Forward analysis type, all parameters should be specified with a fixed value. See the [Stratigraphy Component](#) section of this document for a list of all available parameters and their definitions. Although there are other types of stratigraphy components in NRAP-Open-IAM (e.g., LookupTableStratigraphy), these other component types are not currently available in the GUI.

Parameter	Fixed Value	Value
Datum pressure [Pa]:	101325	
Shale 3 thickness [m]:	Fixed Value	100
Aquifer 2 thickness [m]:	Fixed Value	75
Shale 2 thickness [m]:	Fixed Value	100
Aquifer 1 thickness [m]:	Fixed Value	75
Shale 1 thickness [m]:	Fixed Value	100
Reservoir thickness [m]:	Fixed Value	50

Fig. 2.6: Stratigraphy View

### 2.2.1 Adding Component Models

NRAP-Open-IAM is designed so that only the components of interest need to be included in the system model. Generally, a simulation will be built upwards starting from the deepest component (e.g., first reservoir, then wellbore, then aquifer, then atmosphere). To add a component, first give it a name (each component must have a unique name). Next, select the type of component model to be used. When adding subsequent components, a connection to existing components can be specified.

Fig. 2.7: Adding a Component Model

Each component model has component-specific input parameters and outputs. Parameters can be specified to be sampled from different distributions. If a parameter value is left unchanged from its initial **Fixed Value** setting, the default value shown will be used. When running a **Forward** model, parameters should only be specified as fixed values. When running a **Parstudy** simulation, the parameters meant to vary should be specified as having a uniform distribution and minimum and maximum values. For **LHS** simulations, any distribution can be specified. Parameter and output definitions can be found in the chapter [Components Description](#).



**Simple Reservoir Component**

Parameter	Fixed Value	Value
Reservoir permeability ( $\log_{10} \text{ m}^2$ )	<input type="checkbox"/>	-12
Reservoir porosity [-]	<input type="checkbox"/>	0.3
Brine density ( $\text{kg/m}^3$ )	<input type="checkbox"/>	1000
CO <sub>2</sub> density ( $\text{kg/m}^3$ )	<input type="checkbox"/>	479
Brine viscosity (Pa·s)	<input type="checkbox"/>	0.002535
CO <sub>2</sub> viscosity (Pa·s)	<input type="checkbox"/>	3.95e-5
Brine saturation [-]	<input type="checkbox"/>	0.1
Compressibility ( $\text{Pa}^{-1}$ )	<input type="checkbox"/>	5.1e-11
CO <sub>2</sub> injection rate ( $\text{m}^3/\text{s}$ )	<input type="checkbox"/>	0.1

**Outputs**

☐ Pressure [Pa]    ☐ CO<sub>2</sub> saturation [-]    ☐ CO<sub>2</sub> mass [kg]

Fig. 2.8: Setup of Reservoir Component

When using a component that generally needs input from another component but that component is not to be part of the model (i.e., using a wellbore model without a reservoir model), dynamic parameters can be used for component model input. For dynamic parameters, a value must be specified for each time step in the simulation. Values can be entered manually (separated by a comma) or by providing the path to a file containing the data. Some components require specification of which layer in the stratigraphy they represent (such as an aquifer model).

**Add Component**

Component name:

Model type:

Connections:

Add each component model for the system to be simulated. After specifying all component models, save the model and return to Dashboard to run the simulation.

Fig. 2.9: Adding Second Component

After one component has been added to the system model, more components can be added. When all required components have been included, save the model and return to the dashboard. The system model can then be run using the **RUN SIMULATION** button on the main dashboard.

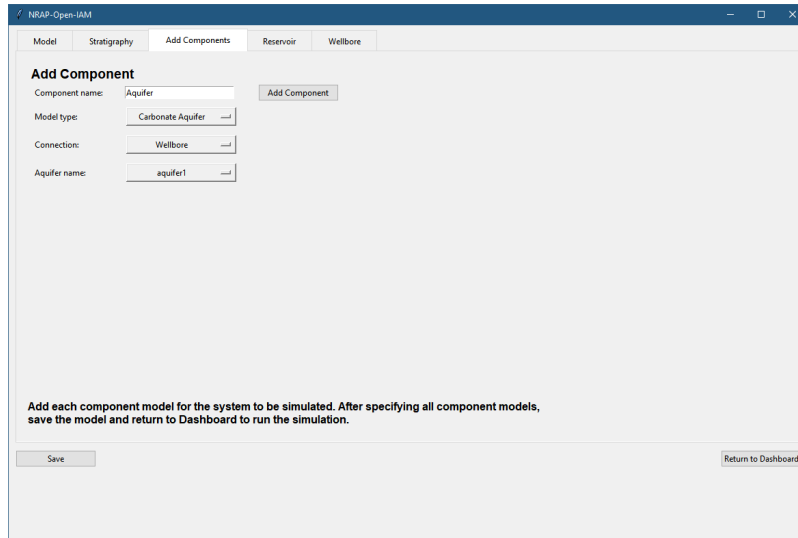


Fig. 2.10: Adding a Component Model with Connection and a Stratigraphy Selection

## 2.2.2 GUI Setup Examples

In the folder *examples*, there is a subfolder called *GUI\_Files* with example simulation files that can be loaded into the GUI and run by NRAP-Open-IAM. To run one of the provided examples select **Load Simulation** on the main dashboard of the GUI. In the file browser that appears, navigate to the *GUI\_Files* subfolder of the *examples* folder and select the first example file *01\_Forward\_AR\_CW.OpenIAM*. This example runs a simple forward model with an *AnalyticalReservoir* component providing inputs to a *CementedWellbore* component.

When the file is loaded into the GUI, the parameters of the simulation can be inspected. After the simulation is complete, the user can proceed to the post-processing step (by clicking **Post Processing** on the main dashboard of the GUI) to visualize and, for some scenarios, analyze the obtained results. The Post Processing tab has a folder selection button with which the user can select the folder containing simulation results. Note that the selection of the folder (and loading of results) might fail if the simulation did not finish successfully. In this case it is recommended to check file *IAM\_log.log* within the output folder containing useful (debug, info, warning or error) messages produced during the simulation. A text editor like Notepad can be used to open the *IAM\_log.log* file. The file name of each GUI example is made to show the components and analyses demonstrated by the example (e.g., AR for *AnalyticalReservoir* and OW for *OpenWellbore*).

The second example file *02\_LHS\_AR\_MSW.OpenIAM* is a stochastic simulation for a system model containing *AnalyticalReservoir* and *MultisegmentedWellbore* components. The example illustrates the use of the Latin Hypercube Sampling analysis type, with parameter values varying across 30 different realizations.

The third example file *03\_LHS\_LUT\_MSW.OpenIAM* illustrates the use of *LookupTableReservoir* and *MultisegmentedWellbore* components. The data set used for the *LookupTableReservoir* component is based on a simulation made for the Kimberlina oil field ([20]).

The fourth example file *04\_LHS\_DP\_MSW.OpenIAM* illustrates Latin Hypercube Sampling analysis type applied to a *MultisegmentedWellbore* component. The pressure and CO<sub>2</sub> saturation required as inputs of the component are provided in the form of arrays. This form of input arguments is called dynamic parameters (DP; i.e., component inputs that change over time).

The fifth example file *05\_LHS\_AR\_OW\_CA.OpenIAM* illustrates the application of three component models: *AnalyticalReservoir*, *OpenWellbore* and *CarbonateAquifer* components. The example uses the LHS analysis type to estimate the reservoir's response to CO<sub>2</sub> injection, the leakage of fluids through a wellbore, and the impact of this leakage on an aquifer overlying the storage reservoir.

## 2.3 Output, Plotting, and Analysis in the GUI

Output is written to the folder specified in the model definition with the `Output` directory. If the path to the output directory is not absolute (i.e., does not contain the drive letter) it is assumed to start from the NRAP-Open-IAM root folder containing the tool distribution.

For each component model of the system, Outputs can be specified. When an output is specified for a forward model the values of that output are written to a file (*output\_name.txt*) in the `Output` Directory. For a stochastic model (LHS or Parstudy), the parameters used and outputs produced are saved to the files *parameter\_realizations.csv* and *output\_realizations.csv*, respectively. A statistical summary of the input parameters and output observations are also saved to the files *parameter\_stats.csv* and *output\_stats.csv*, respectively. Additionally, the GUI input provided is reformatted into a *.yaml* control file, and this file is saved in the `Output` Directory folder.

After a simulation has run, the post processing section of the GUI can be used to generate plots of the results and run sensitivity analyses. When the post processing page is first opened, it asks for a folder specification. This folder is the output folder containing the results you want to analyze. Navigate to that output folder using the **Browse** button.

### 2.3.1 Plotting in the GUI

Users can access the post-processing capabilities of GUI by clicking on the **Post Processing** button on the main page. The **Post Processor** window will appear.

After a folder containing the results of the simulation is selected, different plotting options appear. There are several types of plots that can be created depending on the type of simulation the components used. The simplest plot is a `Time Series` plot, where the output is plotted against time. If the simulation is stochastic (LHS or Parstudy analysis types), each realization will be plotted as a separate line. When evaluating a stochastic simulation, the `Time Series Stats` and `Time Series And Stats` plot types can also be selected. A `Time Series Stats` plot shows basic statistics of the results over time (mean and median values as well as the four quartiles). A `Time Series And Stats` plot will show these statistics as well as separate lines for each realization. Use the checkboxes to select the output types to display in the plot (e.g., **pressure** or **CO2saturation**). The checkboxes will only reflect the outputs saved by the simulation. The `Use subplots` checkbox specifies whether each result should be shown in its own subplot. Finally, the user can set the plot title and filename with the `Title` and `Filename` entries, respectively.

If an `AtmosphericROM` component was included in the simulation, map-view plots of the atmospheric plume can also be generated.

### 2.3.2 Sensitivity Analysis in the GUI

If the simulation results are from an LHS simulation, the `Processing` menu on the `Post Processing` window will have options for several types of sensitivity analysis. Note that while a sensitivity analysis can be run on simulations with a small number of realizations, the results will most likely be inaccurate. If the sensitivity coefficients do not sum to one, or if they vary largely through time, the number of realizations might need to be increased. Generally, 500 to 1000 realizations are needed for a sensitivity analysis. This number might vary, however, depending on the complexity of the simulation. Each type of sensitivity analysis will produce plots and/or text file output in the output directory.

The `Correlation Coefficients` option produces a plot matrix of either Pearson or Spearman correlation coefficients. Any system model observation can be excluded from the analysis if needed, although no exclusions need to be made.

The `Sensitivity Coefficients` option calculates the sensitivity coefficients for each selected output to all inputs. Selecting multiple outputs will run the sensitivity coefficient calculation multiple times. The capture point is the index for the point in time at which the sensitivity coefficient are to be calculated. A capture point of 0 will correspond with year 0 of the simulation. The analysis produces a bar chart.

The `Multiple Sensitivity Coefficients` option calculates the impact of input parameters on multiple outputs. Multiple outputs should be selected here. The capture point is the index for point in time at which the sensitivity coefficient are to be calculated. The analysis will produce a bar chart.

The `Time Series Sensitivity` option will produce a line graph illustrating how the impact from input parameters changes over time with respect to an output value. Selecting multiple output values will run the analysis multiple times. The capture point determines the time at which the sensitivity coefficients are compared and then ordered based on the comparison.

## 2.4 Control File Interface

Control files are a method of providing user input to NRAP-Open-IAM. These files use a YAML format (extension `.yaml`), and they can be opened with a text editor (e.g., Notepad). Any line in the control file starting with a pound sign (`#`) is a comment that will be ignored by the program. The basic format of the control file is the name of an entry (e.g., parameter name) followed by a colon, a space, and a value. Some entries are not directly followed by a value. Instead, the colon is the last character on the line and then other entries are indented beneath the entry.

The use of NRAP-Open-IAM with a control file is called the control file interface (CFI). In this section, the use of the CFI is demonstrated with excerpts from *ControlFile\_ex1a.yaml*:

```
1  #-----
2  # NRAP-Open-IAM Control File Example 1a
3  #-----
4  ModelParams:
5      EndTime: 50
6      TimeStep: 1.0
7      Analysis: forward
8      Components: [AnalyticalReservoir1,
9                  CementedWellbore1]
10     OutputDirectory: ../../output/output_ex1a_{datetime}
11     Logging: Debug
12     GenerateOutputFiles: True
13     GenerateCombOutputFile: False
14     GenerateStatFiles: False
```

Here, the first three lines are comments that are not read by the code. The fourth line defines the keyword `ModelParams` which describes parameters of the system model. The subsequent lines contain parameters of `ModelParams`. A `ModelParams` section is required in all NRAP-Open-IAM control files. The `EndTime` keyword defines the ending time for the simulation in years (50 years in this example). The `TimeStep` parameter defines the length of a time step (1 year in this example). The type of analysis being run is a `forward` (deterministic) simulation. Other possible options for `Analysis` parameter are `lhs` for Latin Hypercube Sampling analysis and `parstudy` for a parameter study. See section [Conceptual Model and Overview](#) for more details regarding these analysis types.

The `Components` parameter is a required entry that contains a list of component model names that are defined later in the file. The component list will always begin with a square bracket '[' followed by each of the component names that make up the system separated by a comma ',' and closed by a square bracket ']'. The names of the components are listed in the order they are supposed to be run. This order can be important, for example, when a wellbore component needs to run after the reservoir component it is connected to.

The next keyword `OutputDirectory` defines a directory for the output to be written into. The output directory can be appended with a keyword `{datetime}`. When a simulation is run, the `{datetime}` keyword will be replaced with the date and time of the simulation run. Note that the `{datetime}` keyword is optional: if it is omitted subsequent runs of the simulation will overwrite past results. That is, if there is a need to keep all results from re-running an NRAP-Open-IAM case, the `{datetime}` keyword will easily facilitate this; if re-running an NRAP-Open-IAM case should

overwrite previous results, the {datetime} keyword should be omitted. In the output folder, NRAP-Open-IAM places a copy of the input file, a folder ("csv\_files") containing all outputs from the component models written to .csv files, and files for all of the plots created.

The keyword `Logging` defines what level of logging information is written out to the logging files. Options for `Logging` levels are `Debug`, `Info`, `Warning`, `Error`, and `Critical`. `Info` is the default level (if no `Logging` keyword is given, the logging will be set to `Info`), and it provides valuable information about when parameters go outside of permitted ranges and when there are problems in the program. `Debug` is a good option if there are problems with the simulation and more information is required to explore the causes. A logging level of `Warning` will limit the information to only messages deemed more important (warning messages). Choosing logging levels of `Error` or `Critical` will limit the messages to only more serious problems, with `Critical` messages having the highest importance.

The `GenerateOutputFiles`, `GenerateCombOutputFile`, and `GenerateStatFiles` control the types of output files saved. Each of these entries can be set to `True` or `False`; if any of these entries are not provided, the default setting is `True`. If `GenerateOutputFiles` is `True`, the simulation will save results to .csv files. Each component output will be saved to its own .csv file. If `GenerateCombOutputFile` is `True`, all simulation results will be saved to one file. If the analysis type is `forward`, the file is called `simulation_results.csv`. If the analysis type is `lhs` or `parstudy`, the file is called `output_realizations.csv` and the parameter values used are saved to another file called `parameter_realizations.csv`. If `GenerateStatFiles` is `True` and the analysis type is `lhs` or `parstudy`, the simulation will save two files detailing the statistics of the output (`output_stats.csv`) and the parameter values used (`parameter_stats.csv`). The statistics shown are the minimum, maximum, and mean values as well as the standard deviation, variance, and different percentiles (2.5%, 5%, 25%, 50%, 75%, 95%, and 97.5%; in the files, the 2.5% and 97.5% values are labelled as the 25th per mille and 975th per mille, respectively, as percentiles do not include decimals). If the analysis type is `forward`, the statistics files will not be saved.

```

1  #-----
2  Stratigraphy:
3      numberOfShaleLayers:
4          vary: False
5          value: 3
6      # Thickness is in meters
7      shale1Thickness:
8          min: 500.0
9          max: 550.0
10         value: 525.0
11      shale2Thickness:
12          min: 450.0
13          max: 500.0
14          value: 475.0
15      shale3Thickness:
16          vary: False
17          value: 11.2
18      aquifer1Thickness:
19          vary: False
20          value: 22.4
21      aquifer2Thickness:
22          vary: False
23          value: 19.2
24      reservoirThickness:
25          vary: False
26          value: 51.2

```

The next section of the file is the `Stratigraphy` section. This section defines any model parameters related to the stratigraphy of the CO<sub>2</sub> storage site. See the [Stratigraphy Component](#) section of this document for a list of all available parameters for the `Stratigraphy` component. Note that while this example uses a

Stratigraphy component, there are other types of stratigraphy components available (e.g., `DippingStratigraphy` and `LookupTableStratigraphy`). A control file must have a section for one of the stratigraphy component types.

Any parameters for the `Stratigraphy` component are defined here with either a deterministic value or a range to vary over. A fixed value of any given parameter can be specified with the `vary: False` and `value: ###` specification shown here or simply `parameterName: ###` (where `###` is the value). If the parameter is meant to vary across different realizations, the `min` and `max` entries should be provided under the parameter to represent the minimum and maximum parameter limits, respectively.

The next sections of the input file define every component model contained in the component model list specified earlier in the control file (under `ModelParams`). The first component in the `Components` list is `AnalyticalReservoir1`, and the component settings are defined in the following section:

```

37 #-----
38 # AnalyticalReservoir1 is a user defined name for component;
39 # the type AnalyticalReservoir is the ROM model name
40 #-----
41 AnalyticalReservoir1:
42     Type: AnalyticalReservoir
43     InjectionWell:
44         coordx: 10
45         coordy: 20
46     Parameters:
47         injRate: 0.1
48     Outputs: [pressure,
49              CO2saturation]
```

The name `AnalyticalReservoir1` can be replaced with any other name defined by user (e.g., `ARes1`), but the component will only be included in the system model if its name is in the `Components` list described in the previous section `ModelParams` (the names used must match). The `Type` is a keyword that defines the component model to be used and must match up with one of the component models currently available in NRAP-Open-IAM. The `InjectionWell` entry specifies the location of the injection well. The `x` and `y` coordinates of the injection well are set with the `coordx` and `coordy` values under `InjectionWell`, respectively. If these inputs are not provided, the default injection well location for an `AnalyticalReservoir` is  $x = 0\text{ m}$ ,  $y = 0\text{ m}$ . The `Parameters` section defines parameters of the component model. Descriptions of the parameters available for the user to specify can be found in the [Components Description](#) chapter of the current documentation. The component model parameters are specified in the same fashion as the `Stratigraphy` parameters shown above. The `Outputs` entry specifies the observations of the component model that will be output from the simulation. Please refer to the [Components Description](#) chapter of this document to see which parameters and outputs are available for user specification in the CFI.

Generally, dynamic (time-varying) input to component models comes from the output of other connected component models (e.g., the pressure and saturation as an input to a wellbore leakage model comes from a reservoir model). In some instances, there may be a need to study a component model without another attached component models feeding the input. In this case dynamic input can be specified with the `DynamicParameters` keyword. Under the `DynamicParameters` section each input name is specified followed by a list of values (enclosed in square brackets, “[ ]”) of the same length as the number of time points (a value for each time point, including an initial value). See files `ControlFile_ex7a.yaml` and `ControlFile_ex7b.yaml` for example of control files utilizing dynamic input for some components.

The next section of the input file is similar to the previous section and defines the next component model `CementedWellbore1`.

```

47 #-----
48 CementedWellbore1:
49     Type: CementedWellbore
```

(continues on next page)

(continued from previous page)

```

50 Connection: AnalyticalReservoir11
51 Number: 4
52 Locations:
53   coordx: [100, 540]
54   coordy: [100, 630]
55 RandomLocDomain:
56   xmin: 150
57   xmax: 250
58   ymin: 200
59   ymax: 300
60 Parameters:
61   logWellPerm:
62     min: -14.0
63     max: -12.0
64     value: -13.0
65 Outputs: [CO2_aquifer1,
66           CO2_aquifer2,
67           CO2_atm,
68           brine_aquifer1,
69           brine_aquifer2]

```

This part of the example sets up the `CementedWellbore` component model named `CementedWellbore1`. There are four wellbores of this type being added with `Number: 4`: two of the locations are given in the `Locations` part and the other two are generated randomly within the domain specified in the `RandomLocDomain` entry. All coordinate systems are assumed to have units of meters.

Known wellbore coordinates are entered as a comma separated list (i.e., contained in brackets, “[ ]”). The x and y coordinates are entered with the `coordx` and `coordy` entries, respectively. There must be a comma between each coordinate (e.g., `coordx: [100, 540]`).

Unknown wellbore locations can be generated by specifying more wellbores (with `Number: ```) than the number of known wellbore locations. To control the locations of randomly placed wells, a ```RandomLocDomain` section needs to be used as:

```

55 RandomLocDomain:
56   xmin: 150
57   xmax: 250
58   ymin: 200
59   ymax: 300

```

This specification will limit the x-coordinate of random wells to be between 150 and 250, and the y-coordinate to be between 200 and 300. Sampling will be from a uniform distribution on the domain defined by `xmin` and `xmax` (or `ymin` and `ymax`). Known wells will be placed first; after all known well coordinates are used, wells will be placed within the random wells domain. See *ControlFile\_ex3.yaml* for an example using random well placement and *ControlFile\_ex4a.yaml* for example using only known well locations.

Note that when using a `LookupTableReservoir` component, the wellbore locations must occur in the domain contained within the files used for the Lookup Table Reservoir component. If any wellbore locations fall outside of the x and y values covered in that file, the simulation will encounter an error.

Different components can take different entries in a control file. For example, the `InjectionWell` entry works with an `AnalyticalReservoir` component, but not a `LookupTableReservoir` component (for that component, the injection well locations are set by the reservoir simulations used to create the files used). The entries that can be provided for each component type are demonstrated in different control file examples; see chapter [Components Description](#) to find control file and script examples for each component.



The last section of the input file is used to specify the plots to be produced.

```

70 #-----
71 # Plot setup part of the control file
72 #-----
73 Plots:
74     CO2_Leakage1:
75         TimeSeries: [CO2_aquifer1]
76         Subplot:
77             NumCols: 2
78             Use: True
79     CO2_Leakage2:
80         TimeSeries: [CO2_aquifer2]
81         Subplot:
82             NumCols: 2
83             Use: True
84     Pressure_plot:
85         TimeSeries: [pressure]
86         Subplot:
87             NumCols: 2
88             Use: True
89         AnalyticalReservoir1_000.pressure: 'Pressure at well #1'
90         AnalyticalReservoir1_001.pressure: 'Pressure at well #2'
91         AnalyticalReservoir1_002.pressure: 'Pressure at well #3'
92         AnalyticalReservoir1_003.pressure: 'Pressure at well #4'
93     Title: Reservoir Pressures at Wellbore Locations

```

Here, three plots are being requested (*CO2\_Leakage1*, *CO2\_Leakage2*, and *Pressure\_plot*). The first two plots will illustrate the CO<sub>2</sub> leakage to the shallow aquifer and the thief zone aquifer; the third plot will illustrate the pressures in the reservoir for the four wellbore locations specified earlier in the control file. *CO2\_Leakage1*, *CO2\_Leakage2*, and *Pressure\_plot* are the user-defined names of the three plots to be created; these names will also be used as the filenames of the figures saved in the output directory. *TimeSeries* is a keyword that instructs the program to plot the observation data as a time series plot. The values to be plotted in each of the three plots (**CO2\_aquifer1**, **CO2\_aquifer2**, and **pressure**) have to be defined in the control file as outputs from one of the specified component models. Each plot will have a title corresponding to the values plotted. A user-defined title can be specified with the *Title* keyword (as illustrated for the *Pressure\_plot*) in the given plot section. For each aquifer, the CO<sub>2</sub> leakage rates for all wells will be plotted on the same figure but on different subplots. If each observation is to be plotted on a separate subplot, the *Subplot* keyword with *Use* set to *True* must be specified, as illustrated in the example setup. Additionally, the *NumCols* keyword (under the *Subplot* section) can be used to set the number of subplot columns to use. The number of rows is controlled by the number of different values (observations) to plot over the number of columns. Each subplot will be given a default title based on the variable portrayed in the subplot. The subplot title names can be replaced with the user defined ones by using the full observation name as a key (e.g., *AnalyticalReservoir1\_000.pressure* for the pressure at the first location) and the desired title as the value under the *Subplot* section as shown in the setup of *Pressure\_plot*. Note that these titles apply to specific subplots, while the *Title* keyword discussed above applies to the larger figure.

The example file described here can be found in the *examples/Control\_Files* directory with the filename *ControlFile\_ex1a.yaml*. To run this example, open a command prompt in the *examples/Control\_Files* directory, activate the environment created for NRAP-Open-IAM, and run the command:

```
python ../../source/openiam/openiam_cf.py --file ControlFile_ex1a.yaml
```

Note: use \ on Windows and / on Mac and Linux.

Other example control files can be found in the same directory. These examples can be run by replacing the file name



in the above command with the user specified one. Different control file examples are used to demonstrate all of the visualization options available in NRAP-Open-IAM. For more details regarding the visualization options available, see section *Visualization Options in the CFI*.

In a control file, an entry situated above the other entries will be read into Python as a dictionary. For example, the Plots section shown above will be converted into the following format in Python:

```

47 Plots = {
48     'CO2_Leakage1':
49         {'TimeSeries': ['CO2_aquifer1'],
50          'Subplot':
51              {'NumCols': 2,
52               'Use': True
53              }
54         },
55     'CO2_Leakage2':
56         {'TimeSeries': ['CO2_aquifer2'],
57          'Subplot':
58              {'NumCols': 2,
59               'Use': True
60              }
61         },
62     'Pressure_plot':
63         {'TimeSeries': ['pressure'],
64          'Subplot':
65              {'NumCols': 2,
66               'Use': True,
67               'AnalyticalReservoir1_000.pressure': 'Pressure at well #1',
68               'AnalyticalReservoir1_001.pressure': 'Pressure at well #2',
69               'AnalyticalReservoir1_002.pressure': 'Pressure at well #3',
70               'AnalyticalReservoir1_003.pressure': 'Pressure at well #4'
71              },
72          'Title': 'Reservoir Pressures at Wellbore Locations'
73         }
74     }

```

In Python, the '{' and '}' characters mark the start and end of a dictionary. The entire control file is read into *openiam\_cf.py* as a dictionary called *yaml\_data*. If a user only wants to run simulations with the CFI, then it is not necessary for the user to understand this conversion or to know Python. If a user wants to access parts of the CFI through a script-based approach (e.g., using the NRAP-Open-IAM plot types in a user-created script), however, then understanding this conversion is important. There are script examples demonstrating the use of NRAP-Open-IAM plot types developed for the CFI, including the examples *iam\_sys\_reservoir\_mswell\_4aquifers\_timeseries.py*, *iam\_sys\_reservoir\_mswell\_stratplot\_dipping\_strata.py*, and *iam\_sys\_reservoir\_mswell\_futuregen\_ttfplot\_dipping\_strata.py*.

## 2.5 Visualization Options in the CFI

The plot types available within NRAP-Open-IAM are: `TimeSeries`, `TimeSeriesStats`, `TimeSeriesAndStats`, `StratigraphicColumn`, `Stratigraphy`, `AoR`, `TTFD`, `GriddedMetric`, `GriddedRadialMetric`, `AtmPlumeSingle`, and `AtmPlumeEnsemble`. This section review the process of creating these plots in the control file interface (CFI) and then discusses each plot type separately.

To create a figure using a simulation run with a `.yaml` control file, the the figure must be set up in the `Plots` section. Within the `Plots` section, the user can have multiple entries that each represent a different plot to be created. The names of these entries are defined by the user, and the names are generally used as the file name for the corresponding figure. An exception occurs with some plots. For example, with the `TTFD` plot type generates a large number of plots, and the final file names will depend on the input used (e.g., `TDS_Plume_Timings_Realization10.png`). The plot name provided can have extensions appended that specify the file type of the resulting figure file (e.g., `.png`, `.tiff`, or `.eps`). Below, we show two examples of `TimeSeries` plot entries in a `.yaml` control file.

```
1 Plots:
2   Pressure_Figure:
3     TimeSeries: [pressure]
4   CO2_Sat_Figure.tiff:
5     TimeSeries: [CO2saturation]
```

Since the first plot entry (`Pressure_Figure`) does not have an extension (e.g., `.png` or `.tiff`) appended at the end (e.g., `Pressure_Figure.tiff`), the produced figure will be of the default `.png` type. The second plot entry (`CO2_Sat_Figure.tiff`), however, includes `.tiff` at the end of the name; the resulting figure file will be of `.tiff` type. Note that if an extension is provided when using a plot type that generates names (e.g., the `TTFD` plot type), the generated names will still use the extension provided.

When we refer to an entry as being indented beneath another entry, we mean that in a `.yaml` file the indented entry is on a lower line and preceeded by additional four spaces. The indentation of one entry carries over to all entries contained within that entry. In the example above, for example, `Pressure_Figure:` is indented beneath `Plots:`, and `TimeSeries: [pressure]` is indented beneath `Pressure_Figure:`. `TimeSeries` is preceeded by eight spaces, while `Pressure_Figure` is preceeded by four spaces. Additionally, the name of each entry is followed by a colon (`:`). When entries have other entries indented beneath them, the colon will be the last character on that line (e.g., `Plots:` or `Pressure_Figure:`). When entries do not have other entries indented beneath them and instead take an input value or list, then the colon is followed by that input (e.g., `TimeSeries: [pressure]`). Note that for the rest of this section, we will not include the colon when discussing an entry; it is assumed that each entry is followed by a colon.

All plot types have certain entries that are either required or optional. Some of these optional entries are used by multiple plot types. To avoid repeating the definitions of these entries, we first present the optional entries used by multiple plot types. Then, we review each plot type.

Each plot type has the optional entries `FigureDPI` and `FigureSize`.

- **FigureDPI** - the dots-per-inch (DPI) of the resulting figure(s) (default is 100). Most figure types produce only one figure file, but plot types like `Stratigraphy` and `TTFD` can produce multiple figures from one entry. Larger DPIs will create high-resolution, high-quality figures, but the file sizes are also larger. File size and quality are also influenced by the extension used (e.g., `.png` or `.tiff`). Recommended `FigureDPI` values are between 100 and 300.
- **FigureSize** - the width and height of the figure in inches. The width and height must be entered as a list of length two (e.g., `FigureSize: [15, 8]`), with the width as the first number and the height as the second. The default values for the plot types are: `[13, 8]` for `TimeSeries`, `TimeSeriesStats`, and `TimeSeriesAndStats`; `[6, 10]` for `StratigraphicColumn` plots; `[12, 10]` for `Stratigraphy` plots; `[10, 8]` for `AoR`, `TTFD`, `GriddedMetric`, and `GriddedRadialMetric` plots; and `[15, 10]` for `AtmPlumeSingle` and `AtmPlumeEnsemble` plots. This entry can also be provided as `figsize` instead of `FigureSize`.

The TimeSeries, TimeSeriesStats, TimeSeriesAndStats, AoR, StratigraphicColumn, and Stratigraphy plot types have the optional entry Title.

- **Title** - the text placed in bold at the top of the figure. If the Title entry is given for StratigraphicColumn or Stratigraphy plots, it replaces the default titles. For the remaining plot types, by default, there is no title for the overall figure, only titles for each individual subplot generated based on the output shown (with AoR plots having one subplot). Note that if one wants to have CO<sub>2</sub> in a title, one should enter it as 'CO\$\_2\$' (the \$\$ symbols aid in the proper formatting).

Below , we show examples of FigureDPI, FigureSize, and Title entries used in a *.yaml* control file.

```

1 Plots:
2   Strat_Col:
3     StratigraphicColumn:
4       Title: Stratigraphy
5       FigureDPI: 200
6       FigureSize: [8, 12]
7   Stratigraphy_Figure.tiff:
8     Stratigraphy:
9       Title: Stratigraphy
10      FigureDPI: 300
11      FigureSize: [13, 11]
12   Pressure_Figure:
13     TimeSeriesStats: [pressure]
14     FigureDPI: 100
15     FigureSize: [12, 8]
16     Title: Pressure Results
17   TDS_Volume_AoR.tiff:
18     AoR: [Dissolved_salt_volume]
19     FigureDPI: 200
20     FigureSize: [12, 12]
21     Title: AoR Results, Dissolved Salt Impact
22   TTFD_Figures:
23     TTFD:
24       PlumeType: Dissolved_salt
25       ComponentNameList: [GenericAquifer1, GenericAquifer2]
26       FigureDPI: 200
27       FigureSize: [12, 11]
28   Gridded_Brine_Aquifer:
29     GriddedMetric:
30       ComponentNameList: [FaultFlow1]
31       MetricName: brine_aquifer
32       FigureDPI: 200
33       FigureSize: [9, 9]
34   Gridded_Salt_Mass_Frac:
35     GriddedRadialMetric:
36       ComponentNameList: [GenericAquifer1]
37       MetricName: Dissolved_salt_mass_fraction
38       FigureDPI: 200
39       FigureSize: [11, 9]
40   Atmospheric_Plume_Single:
41     AtmPlumeSingle:
42       FigureDPI: 200
43       FigureSize: [16, 12]

```

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```

44 Atmospheric_Plume_Probability.tiff:
45     AtmPlumeEnsemble:
46         FigureDPI: 300
47         FigureSize: [14, 11]

```

Notice that the `FigureDPI` and `FigureSize` entries for the `StratigraphicColumn`, `Stratigraphy`, `TTFD`, `GriddedMetric`, `GriddedRadialMetric`, `AtmPlumeSingle`, and `AtmPlumeEnsemble` plots are indented under the plot type. In contrast, the `FigureDPI`, `FigureSize`, and `Title` entries for the `TimeSeriesStats` and `AoR` plots are not indented beneath the plot type. This discrepancy occurs because the `TimeSeriesStats` and `AoR` entries are followed by a metric (e.g., `[pressure]`), while the other plot type entries are not.

The `StratigraphicColumn` and `Stratigraphy` plot types both have the optional entries `ReservoirColor`, `ShaleColor`, `AquiferColor`, `ReservoirAlpha`, `ShaleAlpha`, `AquiferAlpha`, `ReservoirLabel`, `Shale#Label`, and `Aquifer#Label` (where `#` is a particular unit number). Note that the color and alpha entries containing `Shale` and `Aquifer` can be used with a number specifying a certain shale or aquifer (e.g., `Shale2Color` or `Aquifer1Alpha`). Without a specific number, these entries will apply to all shales or aquifers (excluding units that have their own, separate entry of the same type). Note that the color entries can be a string (e.g., `ReservoirColor: orange` or `Aquifer2Color: g`) or a list of length three representing fractions of red, green, and blue (`Aquifer2Color: [0.25, 0.25, 1]`). The entries containing `Alpha` set the alpha values used in the plot. Alpha values range from 0 to 1 and control transparency, with 1 being fully opaque and values approaching 0 becoming more transparent. For examples showing the use of these entries, see *ControlFile\_ex33b*. To prevent a label from being shown, a label entry can be given as `'` (e.g., `Shale2Label: ''`). One might not want a label if the setup causes overlap between different labels.

- `ReservoirColor` - the color used when plotting the reservoir. The default is `[0.33, 0.33, 0.33]`.
- `ShaleColor` - the color used when plotting all shales (or a specific shale, if given as `Shale#Color`, where `#` is an appropriate unit number). The default is red.
- `AquiferColor` - the color used when plotting all aquifers (or a specific aquifer, if given as `Aquifer#Color`, where `#` is an appropriate unit number). The default is blue.
- `ReservoirAlpha` - the alpha value used when plotting the reservoir. The default value is 0.5.
- `ShaleAlpha` - the alpha value used when plotting all shales (or a specific shale, if given as `Shale#Alpha`, where `#` is an appropriate unit number). The default value is 0.25.
- `AquiferAlpha` - the alpha value used when plotting all aquifers (or a specific aquifer, if given as `Aquifer#Alpha`, where `#` is an appropriate unit number). The default value is 0.25.
- `ReservoirLabel` - the label displayed for the reservoir. In `StratigraphicColumn` plots, the default label is "Reservoir Thickness: H m," where H is the unit thickness. In `Stratigraphy` plots, the default label is "Reservoir."
- `Shale#Label` - the label displayed a specific shale, where `#` is an appropriate unit number. In `StratigraphicColumn` plots, the default label is "Shale # Thickness: H m," where H is the unit thickness. In `Stratigraphy` plots, the default label is "Shale #."
- `Aquifer#Label` - the label displayed a specific aquifer, where `#` is an appropriate unit number. In `StratigraphicColumn` plots, the default label is "Aquifer # Thickness: H m," where H is the unit thickness. In `Stratigraphy` plots, the default label is "Aquifer #."

The `Stratigraphy`, `TTFD`, `GriddedMetric`, `GriddedRadialMetric`, `AtmPlumeSingle`, and `AtmPlumeEnsemble` plot types all have the optional entries `PlotInjectionSites`, `InjectionCoordx`, `InjectionCoordy`, `SpecifyXandYLims`, and `SaveCSVFiles`.

- `PlotInjectionSites` - an option to plot injection sites (default is `False`). The only acceptable values are `True` or `False`.

- **InjectionCoordx** - value or list of values for the x coordinate(s) of injection site(s) (default is None). The values are given in meters. This entry must be provided when using a `LookupTableReservoir`, as that component type does not have an `.injX` attribute. Other reservoir types like `AnalyticalReservoir` can be displayed without an `InjectionCoordx` entry.
- **InjectionCoordy** - value or list of values for the y coordinate(s) of injection site(s) (default is None). The values are given in meters. This entry must be provided when using a `LookupTableReservoir`, as that component type does not have an `.injY` attribute. Other reservoir types like `AnalyticalReservoir` can be displayed without an `InjectionCoordy` entry.
- **SaveCSVFiles** - an option to save results in `.csv` files. The only acceptable values are `True` or `False`. The default value for `AoR`, `TTFD`, `GriddedMetric`, `GriddedRadialMetric`, `AtmPlumeSingle`, and `AtmPlumeEnsemble` plots is `True`, while the default value for `Stratigraphy` plots is `False`. For `Stratigraphy` plots, the `.csv` files contain unit thicknesses and depths across the domain.

If set up, `SpecifyXandYLims` is a dictionary containing two entries: `xLims` and `yLims` (i.e., `xLims` and `yLims` are indented beneath ``SpecifyXandYLims`` in a `.yaml` file).

- **SpecifyXandYLims** - a dictionary containing two optional entries related to the limits of the figure's x and y axes (default is None). Within this dictionary are the entries `xLims` and `yLims`.
- **xLims** - an entry under `SpecifyXandYLims` containing a list of length two that represents the x-axis limits (e.g., `xLims: [0, 1000]`; default is None). The values are given in meters. The first and second values in the list are the lower and upper limits, respectively. If `xLims` is not provided or provided incorrectly, the figure will use the default approach for setting the x-axis limits.
- **yLims** - an entry under `SpecifyXandYLims` containing a list of length two that represents the y-axis limits (e.g., `yLims: [0, 1000]`; default is None). The values are given in meters. The first and second values in the list are the lower and upper limits, respectively. If `yLims` is not provided or provided incorrectly, the figure will use the default approach for setting the y-axis limits.

The `Stratigraphy`, `TTFD`, and `AtmPlumeEnsemble` plots also have the optional entry `SpecifyXandYGridLims`, which is a dictionary containing the `gridXLims` and `gridYLims` entries. `AoR` plots do not have grid entries because the x and y values used are those of the wellbore components. `GriddedRadialMetric` plots use the radial grids produced by a component (e.g., a `GenericAquifer` component), while `GriddedMetric` plots use only the locations corresponding with the output.

- **SpecifyXandYGridLims** - a dictionary containing two optional entries related to the x and y limits for the gridded data evaluated (default is None). In `Stratigraphy` plots, the gridded data are the three-dimensional planes depicting the the top of each unit. For `TTFD` and `AtmPlumeEnsemble` plots, the gridded data are the color-labelled values. Within this dictionary are the entries `gridXLims` and `gridYLims`.
- **gridXLims** - an entry under `SpecifyXandYGridLims` containing a list of length two that represents the x-axis limits for the grid used to evaluate results (e.g., `gridXLims: [100, 900]`; default is None). The values for `gridXLims` are in meters. The first and second values in the list are the lower and upper limits, respectively. If `gridXLims` is not provided or provided incorrectly, the figure will use the default approach for creating the gridded values.
- **gridYLims** - an entry under `SpecifyXandYGridLims` containing a list of length two that represents the y-axis limits for the grid used to evaluate results (e.g., `gridYLims: [100, 900]`; default is None). The values for `gridYLims` are in meters. The first and second values in the list are the lower and upper limits, respectively. If `gridYLims` is not provided or provided incorrectly, the figure will use the default approach for creating the gridded values.

The `Stratigraphy`, `TTFD`, and `AtmPlumeEnsemble` plot types can all use the optional entries `xGridSpacing` and `yGridSpacing`:

- **xGridSpacing** - a horizontal distance ( $m$ ) used as the interval between the grid points in the x-direction (default is None). If this entry is not provided, the x-coordinates of the grid points are defined using a default approach (1/100th of the range in x-values).

- `yGridSpacing` - a horizontal distance ( $m$ ) used as the interval between the grid points in the y-direction (default is None). If this entry is not provided, the y-coordinates of the grid points are defined using a default approach (1/100th of the range in y-values).

The AoR, GriddedMetric, and GriddedRadialMetric plot types have the optional entry `TimeList`:

- `TimeList` - a list specifying the times (in years) for which to create separate figures (e.g., `TimeList: [1, 5, 10]`). Otherwise, one figure can be created for each timestep by having `TimeList: All`. If `TimeList` is not entered for an AoR plot, the figures created will show the maximum values for all locations across all model times. If `TimeList` is not entered for a GriddedMetric or GriddedRadialMetric plot, the default setting is `TimeList: All`.

The TTFD, GriddedMetric, and GriddedRadialMetric plot types all have the required entry `ComponentNameList`:

- `ComponentNameList` - a list containing the names provided for each of the components producing output to be used for the creation of the figures (e.g., `ComponentNameList: [FutureGen2AZMI1, FutureGen2AZMI2]` in `ControlFile_ex40.yaml`). Below, we show a section of the `.yaml` file for `ControlFile_ex40.yaml`. This section demonstrates where the name is provided for the `FutureGen2AZMI2` component. Below the excerpt is an example of how component names are set when using NRAP-Open-IAM in a script application.

Excerpt from `ControlFile_ex40` demonstrating how an aquifer component is given the name `FutureGen2AZMI2`:

```

1 FutureGen2AZMI2:
2   Type: FutureGen2AZMI
3   Connection: MultisegmentedWellbore1
4   AquiferName: aquifer3
5   Parameters:
6     por: 0.132
7     log_permh: -12.48
8     log_aniso: 0.3
9     rel_vol_frac_calcite: 0.1
10  Outputs: [pH_volume, TDS_volume, Dissolved_CO2_volume,
11            Dissolved_CO2_dx, Dissolved_CO2_dy, Dissolved_CO2_dz]
```

Example of setting the component name (`FutureGen2AZMI2`) in a script application:

```

1 fga = sm.add_component_model_object(FutureGen2AZMI(name='FutureGen2AZMI2', parent=sm))
```

The GriddedMetric and GriddedRadialMetric plot types both have the required entry `MetricName`:

- `MetricName` - the name of the metric to plot. For a GriddedMetric plot, the SealHorizon and FaultFlow outputs **CO2\_aquifer**, **brine\_aquifer**, **mass\_CO2\_aquifer**, or **mass\_brine\_aquifer** can be provided for `MetricName`. For a GriddedRadialMetric plot, the GenericAquifer outputs **Dissolved\_CO2\_mass\_fraction** or **Dissolved\_salt\_mass\_fraction** can be provided for `MetricName`. When plotting these GenericAquifer metrics, the component used must also produce **r\_coordinate** and **z\_coordinate** outputs.

The GriddedMetric, GriddedRadialMetric, and AtmPlumeSingle plot types have the optional entry `Realization`:

- `Realization` - the realization number for which to display results (default is 0). Note that this optional input is only used in simulations using Latin Hypercube Sampling (`lhs`) and Parameter Study (`parstudy`) analysis types. This input uses the indexing rules in Python, where 0 represents the first realization and ( $N - 1$ ) represents the last (where  $N$  is the number of realizations).

The GriddedMetric and GriddedRadialMetric plot types both have the optional entry `EqualAxes`:

- `EqualAxes` - the option to force the x and y axes to cover the same distances (for an equal aspect ratio). The acceptable values are `True` or `False`, and the default value is `True`. If set to `True`, the axes limits given with

`xLims` and `yLims` will not be used.

Examples of setting up each plot type in a `.yaml` file are shown in the sections below.

### 2.5.1 TimeSeries, TimeSeriesStats, and TimeSeriesAndStats

The `TimeSeries`, `TimeSeriesStats`, and `TimeSeriesAndStats` plot types are used to display results varying over time. Although this section covers three plot types, these plot types are different variations of the same type of plot.

`TimeSeries` plots are line plots of results varying over time. The number of lines in the resulting figure depends on the setup of the scenario. For example, components and associated locations entered in the `.yaml` file can define the number of curves shown in the figure but only the components that produce the metric being plotted (e.g., **pressure** or **brine\_aquifer1**) influence the number of lines created for that particular metric.

`TimeSeriesStats` and `TimeSeriesAndStats` plots can only be produced for simulations using the Latin Hypercube Sampling (LHS, `lhs` in the CFI) or Parameter Study (`parstudy` in the CFI) analysis types (not the `forward` analysis type). Simulations using the `lhs` and `parstudy` analysis types create separate simulations (i.e., different realizations) that explore the parameter space. The parameters varied are those entered with minimum and maximum values, which are meant to model a uniform distribution. Consider, for example, a `TimeSeriesStats` plot set up for an LHS run with 30 realizations. The `ModelParams` section of the `.yaml` file would be similar to this excerpt from `ControlFile_ex4a.yaml`:

```

1  ModelParams:
2      EndTime: 10
3      TimeStep: 1.0
4      Analysis:
5          Type: lhs
6          siz: 30
7      Components: [AnalyticalReservoir1,
8                  OpenWellbore1,
9                  CarbonateAquifer1]
10     OutputDirectory: output/output_ex4a_{datetime}
11     Logging: Debug

```

The entries `Type: lhs` and `siz: 30` under `Analysis` specify the run as an LHS simulation with 30 realizations. Each realization will use different values for the parameters that are set up to vary. In a `TimeSeries` plot, the outputs for each realization will be represented by separate lines.

If an `lhs` or `parstudy` simulation uses many realizations and many component locations, a `TimeSeries` plot could become visually unclear. To avoid a lack of visual clarity, `TimeSeriesStats` plots show basic information about the distribution of results over time. The plot produces lines representing mean and median values as well as shaded regions showing the four quartiles of the distribution varying over time (0th to 25th, 25th to 50th, 50th to 75th and 75th to 100th percentiles).

`TimeSeriesAndStats` plots combine the approaches of `TimeSeries` and `TimeSeriesStats` plots. The mean, median, and quartiles are shown along with line graphs for each realization.

`TimeSeries`, `TimeSeriesStats`, and `TimeSeriesAndStats` plots can have the following optional entries: `UseMarkers`, `VaryLineStyles`, `UseLines`, `Subplot`, `Title`, `FigureDPI`, and `FigureSize` (the latter three are described above). Note that `Subplot` is a dictionary containing the optional entries `Use` and `NumCols` (i.e., the `Use` and `NumCols` options are on a line beneath `Subplots` and preceded by four more spaces).

- `UseMarkers` - an option to show results with values annotated with markers like circles and squares (default is `False`). The only acceptable values are `True` or `False`. If markers are used, the colors of markers and lines will vary in the normal manner (i.e., a rotation through the default matplotlib color order).



- **VaryLineStyles** - an option to vary the line styles used (default is `False`). The only acceptable values are `True` or `False`. The matplotlib line styles used are 'solid', 'dotted', 'dashed', and 'dashdot'. Line colors will still vary in the normal manner.
- **UseLines** - an option to show results with lines (default is `True`). The only acceptable values are `True` or `False`. If neither markers nor lines are used, the plot will not show any results. One should only set **UseLines** to `False` if **UseMarkers** is set to `True`. If **UseLines** is set to `False`, **VaryLineStyles** will automatically be set to `False`, regardless of the entry provided in the `.yaml` file.
- **Subplot** - a dictionary containing the optional entries **Use** and **NumCols**. This entry can also be provided as **subplot**.
- **Use** - the option to use multiple subplots (`True`) or not (`False`). The default value is `True`. The different subplots can show the results for different locations and/or the results for different metrics (e.g., **pressure** in one subplot, **CO2saturation** in another). If different types of output (e.g., **pressure** and **CO2saturation**) are included in a **TimeSeries** plot but **Use** is set to `False`, the y-axis label will only reflect one of the two output types. This entry can also be provided as **use**.
- **NumCols** - the number of columns used to set up the subplots, if **Use** is set to `True`. If the plot includes 3 or fewer metrics (influenced by the output type(s) and locations used), the default value is 1. If the plot includes 4 or more metrics, the default value is 2. This entry can also be given as **ncols**. The number of rows is taken as the number required to plot the metrics used by the plot (given the number of columns). The number of metrics is set by the number of output types given (e.g., two for **TimeSeries**: [**pressure**, **CO2saturation**]) and the number of locations for those output types. For example, if **NumCols** is two, then the number of rows will be half the number of metrics (rounding up to the next integer). Note that **TimeSeries**, **TimeSeriesStats**, and **TimeSeriesAndStats** plots will automatically adjust the font sizes used to seek to avoid text overlapping with other features. Accomplishing a specific subplot configuration without the text becoming too small, for example, can be aided by also changing the **FigureSize** entry (see above).

These optional entries are not indented under **TimeSeries** or **TimeSeriesAndStats** in a `.yaml` file, but are instead indented under the figure name. If **UseMarkers**, **VaryLineStyles**, or **UseLines** are provided for a **TimeSeriesStats** plot, the entries will have no effect (i.e., they do not influence the mean and median lines or the shaded quartiles).

The titles for individual subplots are generated based on the metric shown in the subplot (i.e., output type and location the output was produced for). One can specify the subplot title that will correspond to a specific output, however, by entering the output name under **Subplot**. The output name depends on the corresponding component, the output type, and the location index. For example, an **AnalyticalReservoir** component named **AnalyticalReservoir1** producing **pressure** at location 0 will result in an output name of **AnalyticalReservoir1\_000.pressure**. The component name is followed by an underscore, then the location index (starting at 0 and expressed with three digits), then a period, and finally the output name (e.g., **pressure**). The text used for the title is given after the output name as **ComponentName\_000.OutputName: Text Used for Title**. For an example of this approach, see *ControlFile\_ex1a*.

Below, we show examples of **TimeSeries** and **TimeSeriesAndStats** plots in a `.yaml` control file.

```

1 Plots:
2   Pressure_and_Sat:
3     TimeSeries: [pressure, CO2saturation]
4     UseMarkers: False
5     UseLines: True
6     VaryLineStyles: True
7     FigureDPI: 150
8     Subplot:
9       Use: True
10      NumCols: 2
11      AnalyticalReservoir1_000.pressure: 'Pressure at Well 0'
12      AnalyticalReservoir1_001.pressure: 'Pressure at Well 1'

```

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```

13     AnalyticalReservoir1_000.CO2saturation: 'CO$_2$ Saturation at Well 0'
14     AnalyticalReservoir1_001.CO2saturation: 'CO$_2$ Saturation at Well 1'
15     Pressure_Stats:
16         TimeSeriesAndStats: [pressure]
17         UseMarkers: True
18         UseLines: False
19         VaryLineStyles: False
20         FigureDPI: 400

```

For examples of TimeSeries plots, see control file examples 1a, 1b, 2, 3, 7a, 7b, and 14. For examples of TimeSeriesStats plots, see control file examples 4a, 4b, 6, 8, 15, and 39. For examples of TimeSeriesAndStats plots, see control file examples 4a, 14, and 40. For script examples using the TimeSeries plot, see *iam\_sys\_reservoir\_mswell\_4aquifers\_timeseries.py* and *iam\_sys\_reservoir\_mswell\_4aquifers\_timeseries\_2locs.py*.

## 2.5.2 StratigraphicColumn

StratigraphicColumn plots show unit thicknesses and depths at one location. If the simulation does not use spatially variable stratigraphy (e.g., the Stratigraphy component), then the unit thicknesses at the one location used are representative of the entire domain. If the simulation does use spatially variable stratigraphy (e.g., the DippingStratigraphy or LookupTableStratigraphy components), then the plot shows the calculated unit thicknesses for the location used.

The StratigraphicColumn plot type has the following optional entries: XValue, YValue, DepthText, ReservoirColor, ShaleColor, AquiferColor, ReservoirAlpha, ShaleAlpha, AquiferAlpha, ReservoirLabel, ShaleLabel, AquiferLabel, FigureDPI, FigureSize, and Title. All of these entries but XValue, YValue, and DepthText are described above.

- XValue - the x-coordinate (*m*) of the location used for the plot. The default value is 0 m.
- YValue - the y-coordinate (*m*) of the location used for the plot. The default value is 0 m.
- DepthText - option specifying whether to show depths at each unit interface (True) or not (False). The default value is True. One may want to disable the depth text if, for example, certain units are so thin that the text for different units plot on top of each other.

Two examples of StratigraphicColumn plots in a *.yaml* control file are shown below. One plot does not include any optional entries and, therefore, uses the default options. The other plot includes a variety of optional entries.

```

1     Plots:
2         Strat_Col_Default:
3             StratigraphicColumn:
4         Strat_Col_With_Options.tiff:
5             StratigraphicColumn:
6                 Title: Stratigraphy
7                 Shale1Label: Lower Aquitard
8                 Shale2Label: Middle Aquitard
9                 Shale3Label: Upper Aquitard
10                Aquifer1Label: AZMI
11                Aquifer2Label: Freshwater Aquifer
12                ReservoirLabel: Storage Reservoir
13                ReservoirColor: darkmagenta
14                Shale1Color: [0.33, 0.33, 0.33]
15                Shale2Color: olive
16                Shale3Color: rosybrown

```

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```

17     Aquifer1Color: deeppink
18     Aquifer2Color: mediumturquoise
19     Shale1Alpha: 0.3
20     Shale2Alpha: 0.3
21     Shale3Alpha: 0.45
22     Aquifer1Alpha: 0.3
23     Aquifer2Alpha: 0.45
24     ReservoirAlpha: 0.45
25     FigureDPI: 300
26     XValue: 2000
27     YValue: 2000
28     DepthText: False

```

For more examples of StratigraphicColumn plots, see control file examples *ControlFile\_ex33a-ControlFile\_ex38c*.

### 2.5.3 Stratigraphy

Stratigraphy plots are three-dimensional plots showing the specified stratigraphy as well as features like wellbores and injection sites. These plots work with all stratigraphy component types (Stratigraphy, DippingStratigraphy, and LookupTableStratigraphy).

Stratigraphy plots can have the following optional entries: PlotWellbores, PlotWellLabels, WellLabel, PlotInjectionSites, PlotInjectionSiteLabels, InjectionCoordx, InjectionCoordy, PlotStratComponents, StrikeAndDipSymbol, SpecifyXandYLims, SpecifyXandYGridLims, xGridSpacing, yGridSpacing, View, SaveCSVFiles, ReservoirColor, ShaleColor, AquiferColor, ReservoirAlpha, ShaleAlpha`, AquiferAlpha`, ReservoirLabel, Shale#Label, Aquifer#Label, FigureDPI, FigureSize, and Title. Four of these entries (StrikeAndDipSymbol, SpecifyXandYLims, SpecifyXandYGridLims, and View) are dictionaries containing additional entries (i.e., more entries indented beneath them in a *.yaml* file). The entries SpecifyXandYLims, SpecifyXandYGridLims, xGridSpacing, yGridSpacing, SaveCSVFiles, PlotInjectionSites, InjectionCoordx, InjectionCoordy, ReservoirColor, ShaleColor, AquiferColor, ReservoirAlpha, ShaleAlpha`, AquiferAlpha`, ReservoirLabel, Shale#Label, Aquifer#Label, FigureDPI, and FigureSize are described above.

- **PlotWellbores** - an option to plot wellbores as vertical lines (default is True). The only acceptable values are True or False.
- **PlotWellLabels** - an option to show text labels specifying wellbore types and numbers (default is True). If WellLabel is not entered, labels will be set according to the wellbore component type. For example, the labels could be “Open Wellbore 1” for an Open Wellbore, “M.S. Wellbore 1” for a MultiSegmented Wellbore, or “Cemented Wellbore 1” for a Cemented Wellbore. If WellLabel is entered, the text provided will be used. The only acceptable values are True or False.
- **WellLabel** - the label used for wellbores if PlotWellLabels` is set to ``True. If the text given includes empty brackets (*{}*), then the location index will be inserted in that position. If this entry was given as WellLabel: Legacy Well {}, for example, then the labels would range from “Legacy Well 0” to “Legacy Well (N - 1),” where N is the maximum location index for the wellbore components (location indices use the python indexing). If WellLabel is given without brackets, then the same text will be displayed for each wellbore component (e.g., WellLabel: Well). if PlotWellLabels` is set to ``True but WellLabel is not entered, labels will be set using the default approach.
- **PlotInjectionSiteLabels** - an option to show a text label for the injection site(s) (default is False).
- **PlotStratComponents** - the option to plot squares along each wellbore at the depths at which the wellbore intersects the top of a unit (default is False). The tops of shales are shown with red squares, while the tops of aquifers are shown with blue squares. The only acceptable values are True or False.

- **StrikeAndDipSymbol** - a dictionary containing four optional entries related to the strike and dip symbol shown in the figure (default is None). Within this dictionary are the entries **PlotSymbol**, **coordx**, **coordy**, and **length**.
- **PlotSymbol** - an entry under **StrikeAndDipSymbol** that specifies whether to show the strike and dip symbol (default is True). The only acceptable values are True or False.
- **coordx** - an entry under **StrikeAndDipSymbol** that specifies the x-coordinate at which to plot the strike and dip symbol (default is None). If **coordx** is not provided, the graph will use a default location (which depends on the domain).
- **coordy** - an entry under **StrikeAndDipSymbol** that specifies the y-coordinate at which to plot the strike and dip symbol (default is None). If **coordy** is not provided, the graph will use a default location (which depends on the domain).
- **length** - an entry under **StrikeAndDipSymbol** that specifies the length scale (*m*) of the strike and dip symbol (default is None). For flat-lying units, the length is the diameter of the circular symbol used. For dipping units, the length applies to the line going in direction of strike (not the line in the dip direction). If **length** is not provided, the graph will use a calculated length (which depends on the domain).
- **View** - a dictionary containing two optional entries related to the perspective of the three-dimensional graph (default is None). Within this dictionary are the entries **ViewAngleElevation** and **ViewAngleAzimuth**. A separate version of the figure is created for each combination of the **ViewAngleElevation** and **ViewAngleAzimuth** entries, where the first values in the keywords list are used for the same graph and so on.
- **ViewAngleElevation** - an entry under **View** containing a list of the elevation angles (in degrees) to use in the Stratigraphy plot(s) (default is [10, 30]). Values must be between -90 and 90. See the matplotlib documentation regarding view angles. This list must have the same length as the **ViewAngleAzimuth** list.
- **ViewAngleAzimuth** - an entry under **View** containing a list of the azimuth angles (in degrees) to use in the Stratigraphy plot(s) (default is [10, 30]). Values must be between 0 and 360. See the matplotlib documentation regarding view angles. This list must have the same length as the **ViewAngleElevation** list.

Two examples of .yaml entries for Stratigraphy plots are shown below. The first entry uses the default settings, while the second entry specifies each option. Since the simulation uses a **LookupTableReservoir**, the entry has to include **InjectionCoordx** and **InjectionCoordy**. **InjectionCoordx** and **InjectionCoordy** are not required when using another type of reservoir component with option **PlotInjectionSites: True**.

```

1 Plots:
2   Strat_Plot_Default_Settings:
3     Stratigraphy:
4   Strat_Plot.tiff:
5     Stratigraphy:
6       Title: Proposed GCS Site
7       FigureDPI: 500
8       PlotInjectionSites: True
9       PlotInjectionSiteLabels: True
10      InjectionCoordx: 200
11      InjectionCoordy: 200
12      PlotWellbores: True
13      PlotWellLabels: True
14      PlotStratComponents: True
15      SaveCSVFiles: False
16      SpecifyXandYLims:
17        xLims: [0, 400]
18        yLims: [0, 400]
19      SpecifyXandYGridLims:
20        gridXLims: [25, 375]
```

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```

21         gridYLims: [25, 375]
22     StrikeAndDipSymbol:
23         PlotSymbol: True
24         coordx: 100
25         coordy: 300
26         length: 75
27     View:
28         ViewAngleElevation: [5, 10, 5, 10]
29         ViewAngleAzimuth: [300, 300, 310, 310]

```

For examples of Stratigraphy plots, see examples *ControlFile\_ex33a.yaml-ControlFile\_ex38c.yaml*. For examples of using Stratigraphy plots in a script application, see the files *iam\_sys\_reservoir\_mswell\_stratplot\_dipping\_strata.py* and *iam\_sys\_reservoir\_mswell\_stratplot\_no\_dip.py*.

## 2.5.4 AoR

Area of Review (AoR) plots are developed to estimate the AoR needed for a geologic carbon storage project based on the spatial extent of reservoir impacts (pressure and CO<sub>2</sub> saturation) and potential aquifer impacts (dissolved salt and dissolved CO<sub>2</sub> plume volumes). The potential extent is found by distributing wellbore components across the domain. We recommend setting wellbore locations using the grid placement option (see examples *ControlFile\_ex31a.yaml* to *ControlFile\_ex31d.yaml*). The wellbores are hypothetical and used to consider the aquifer impacts that could occur if a leakage pathway (extending from the reservoir to the aquifer being considered) was available at each wellbore location considered. The approach used for AoR plots is based on the work [1].

AoR plots can be made while using any type of wellbore component, but we recommend using the `OpenWellbore` component. The `OpenWellbore` component represents a worst-case scenario, where the wellbore is uncemented. Additionally, this component can use a critical pressure in leakage calculations. One might want to use another wellbore type (e.g., `MultisegmentedWellbore` or `CementedWellbore`), however, if there are reliable constraints on the effective permeabilities of legacy wells in the study area. Without such constraints, it can be more responsible to evaluate the implications of a worst-case scenario with uncemented legacy wells.

Note that the AoR plot type is meant to be used only for one aquifer at a time, with that aquifer being represented by only one type of aquifer component (e.g., representing contaminant spread in aquifer 2 with a `FutureGen2Aquifer` component). For example, file *ControlFile\_ex31a.yaml* has `AnalyticalReservoir` components that provide the input for `OpenWellbore` components, and the `OpenWellbore` components provide input to `FutureGen2Aquifer` components. The `FutureGen2Aquifer` components are set up to represent aquifer 2. If the user added an entry to the *.yaml* file for a `FutureGen2AZMI` aquifer component representing aquifer 1, the AoR plot could not make plots representing the impacts on both aquifers 1 and 2. In this case, one would need to create a separate *.yaml* file that creates AoR plots just for aquifer 1.

AoR plots can be created for four types of outputs: reservoir pressures (**pressure**), reservoir CO<sub>2</sub> saturations (**CO2saturation**), aquifer contaminant plume volumes from CO<sub>2</sub> leakage (**pH\_volume** and **Dissolved\_CO2\_volume**), and aquifer contaminant plume volumes from brine leakage (**TDS\_volume** or **Dissolved\_salt\_volume**). The type of plume volume output depends on the aquifer component used (e.g., `GenericAquifer` vs. `FutureGen2Aquifer`).

The AoR plot type examines these metrics at each location in the domain (i.e., each hypothetical wellbore location) and displays the maximum value over time (across all times or at specific times, depending on the `TimeList` entry provided; this entry is discussed below). For LHS simulations, the AoR plot displays the maximum values over time at each location from all LHS realizations. This approach is meant to depict how severe the reservoir and aquifer impacts could become.

Note that model run times can increase dramatically with the number of wellbore locations. Additionally, some aquifer components generally require longer model run times (e.g., `GenericAquifer`) in comparison with other aquifer components (e.g., `FutureGen2Aquifer`). Also note that `FutureGen2Aquifer` is meant to be set up for aquifers with

bottom depths  $\leq 700$  m, while FutureGen2AZMI is meant to be set up for aquifers with bottom depths  $\geq 700$  m.

The AoR plot type can be used with `AnalyticalReservoir`, `LookupTableReservoir`, and `TheisReservoir` components. When using a `TheisReservoir` component, however, the system model should not include wellbore or aquifer components (see `ControlFile_ex47.yaml`). The `TheisReservoir` can represent multiple injection and/or extraction wells, but the component only produces **pressure** output (if **CO2saturation** is requested from the component, the values are always zero). When using a `TheisReservoir`, the AoR plot type should only be used on the **pressure** output. While an AoR plot usually focuses on the locations used for the wellbore component, when using a `TheisReservoir` component the locations used are those entered directly for the `TheisReservoir` (`ControlFile_ex47.yaml`).

Using the AoR plot type leads to the creation of `.csv` files containing the values shown in the AoR plots. When using the AoR plot type, we recommend setting `GenerateOutputFiles` and `GenerateCombOutputFile` to `False` in the `ModelParams` section of the `.yaml` file. The large number of wellbore locations commonly used for AoR plots causes a large number of output files. A reservoir and aquifer component is created for each wellbore location, and every component will have its output saved if those options are set to `True`. The `.csv` files created for the AoR plots contain all of the necessary information and these files are much smaller in size.

AoR plots can have nine optional entries: `PlotInjectionSites`, `InjectionCoordx`, `InjectionCoordy`, `SaveCSVFiles`, `FigureDPI`, `FigureSize`, `TimeList`, `CriticalPressureMPa`, and `BrineDensity`. All of these entries except for `CriticalPressureMPa` and `BrineDensity` are described above.

- **CriticalPressureMPa** - this entry controls how critical pressure is handled in an AoR plot evaluating the *pressure* metric. The entry can either be given as `Calculated` or as a number representing a critical pressure in MPa (e.g., `CriticalPressureMPa: 20.5` for 20.5 MPa). Note that this plot entry only controls how reservoir pressures are evaluated in an AoR plot; this plot entry will not impact the behavior of an `OpenWellbore` component using a critical pressure. The critical pressure for an `OpenWellbore` component must be handled in the set up of the component itself.
- **BrineDensity** - When the critical pressure is calculated and the wellbore component has a brine density parameter (e.g., the `brineDensity` parameter of `OpenWellbore` and `MultisegmentedWellbore` components), that brine density parameter will be used when calculating the critical pressure. When the wellbore component does not have a brine density parameter (e.g., `CementedWellbore` components) but critical pressure is calculated, then a brine density in  $kg/m^3$  can be provided with the `BrineDensity` entry (e.g., `BrineDensity: 1100` for a density of  $1100 kg/m^3$ ). If needed but not provided, the default value is  $1012 kg/m^3$ . If this entry would not be used, given the wellbore component type and `CriticalPressureMPa` entry, then any input provided for `BrineDensity` will be ignored.

If the `CriticalPressureMPa` entry is given for an AoR plot evaluating the **pressure** metric, the figure will highlight all locations with pressures exceeding the critical pressure used. Without the inclusion of the `CriticalPressureMPa` entry, an AoR plot examining **pressure** will only display the reservoir pressures across the domain (it will not highlight a particular area of the domain).

For AoR plots that evaluate the other metrics (**CO2saturation**, **pH\_volume**, **TDS\_volume**, **Dissolved\_CO2\_volume**, and **Dissolved\_salt\_volume**), the figure will highlight any wellbore location that has a nonzero result (i.e., a nonzero CO<sub>2</sub> saturation or a nonzero plume volume).

The AoR should, however, extend to the grid points closest to the injection site(s) that do not satisfy these criteria (nonzero **CO2saturation** values, nonzero plume volumes, or **pressure** values exceeding the critical pressure). In other words, consider a situation where there is a point with nonzero plume volumes is next to a point that never had plume volumes. If another point was placed between these two points, this new point may also have nonzero plume volumes. Therefore, excluding this area from the AoR may prevent the detection of leakage events. No such exclusion will occur if the boundaries of the AoR include points that never met the criteria discussed above.

If the `TimeList` entry is not provided for an AoR plot, the figure will show the maximum values at each location across all model times. If `TimeList` is provided as a list of times in years (e.g., `TimeList: [1, 5, 10]` or `TimeList: [10]`), then the figures created will represent the maximum values at each location at the specified time(s). Otherwise, an AoR figure can be made for every model time by providing `TimeList: All`. Evaluating how the potential impacts of a project change over time can inform, for example, how the required extents of surveying efforts change over time

(i.e., discovering and effectively plugging legacy wells at larger distances from the injection site).

Below is an example of two AoR plot entries in a `.yaml` file. The first entry uses the default settings, while the second specifies all available options. Since the simulation uses a `LookupTableReservoir` this example includes `InjectionCoordx` and `InjectionCoordy`. These inputs are not required for other reservoir component types.

```
1 Plots:
2   AoR_pH_Default_Settings:
3     AoR: [pH_volume]
4   AoR_TDS.tiff:
5     AoR: [TDS_volume]
6     PlotInjectionSites: True
7     InjectionCoordx: 2.37e5
8     InjectionCoordy: 4.41e6
9     FigureDPI: 300
10    SaveCSVFiles: False
11    TimeList: All
```

For examples of AoR plots, see *ControlFile\_ex31a.yaml* to *ControlFile\_ex32c.yaml* and *ControlFile\_ex47.yaml*.

## 2.5.5 TTFD

The time to first detection (TTFD) plot type uses contaminant plume output from aquifer components to simulate when a monitoring well would be able to detect the plume in the aquifer(s) considered. If the TTFD plot type is run without monitoring locations provided, it still produces maps showing the spread of contaminant plumes across the domain. These figures (and the `.csv` files that can be saved) could then be used to decide where to place monitoring sensors.

The TTFD plot type can produce three types of figures: maps of earliest plume timings across the domain (i.e., the earliest time at which the plume type occurs in each part of the aquifer(s) considered), maps showing the TTFD provided by the entered monitoring locations, and maps of the probability of plume occurrence in the aquifer(s) considered. The figures with the TTFD from monitoring locations are only created if monitoring locations are entered. The maps of plume probabilities are only created if the analysis type is Latin Hypercube Sampling (`lhs`) or Parameter Study (`parstudy`). Note that plume probabilities are calculated as the number of realizations in which a plume occurred at each location divided by the total number of realizations.

The TTFD plot type requires the use of at least one of the following aquifer component types (with the component(s) set up to represent the aquifer(s) considered): `CarbonateAquifer`, `FutureGen2Aquifer`, `FutureGen2AZMI`, `GenericAquifer`, `DeepAlluviumAquifer`, or `DeepAlluviumAquiferML`. Note that the `FutureGen2Aquifer` component is used for aquifers with bottom depths  $\leq 700$  m, while the `FutureGen2AZMI` component is used for aquifers with bottom depths  $\geq 700$  m. The aquifer component(s) must also produce the plume dimension metrics associated with the plume type considered (e.g., `TDS_dx`, `TDS_dy`, and `TDS_dz` for TDS plumes). Note that `CarbonateAquifer` components do not produce plume dimension outputs for different plume types, so the required outputs when using `CarbonateAquifer` are `dx` and `dy` (which represent the lengths of the impacted aquifer volume in the x- and y-directions, respectively).

The plume timing and plume probability figures made with the TTFD plot type show four subplots. Each subplot contains a quarter of the depth range from the top of the reservoir to the surface. Each subplot contains the results for sections of aquifers within the corresponding depth range. If monitoring sensor locations are provided, each subplot will also show any sensors with depth (`z`) values in the subplot's depth range as black triangles. Because there are multiple `z` grid points within each subplot, there can be different layers of results displayed. The code is set up to make the top layer shown be the layer with the lowest plume timing or highest plume probability (for the corresponding figure types). The matplotlib function used to display results by color (`contourf`) can fail to display results when there are very few points with results in a layer. To address such situations, if there are fewer than 25 points with results we display each value as a color-labelled circle.



While the plume timing plots show the earliest plume timings at each grid location across the domain, the monitoring TTFD plots only display plume timings that are sufficiently close to the sensor location(s) provided. The purpose of such graphs is to show when the sensors used could warn site operators that an aquifer has been impacted. If the chosen sensor *x*, *y*, and *z* values do not provide any warning of plumes in an aquifer, and there are plumes in that aquifer, then the monitoring locations should be changed. The distance over which sensors can detect a plume are controlled by the `VerticalWindow` and `HorizontalWindow` entries, which are discussed below. Note that the TTFD plot type can produce output for the DREAM tool (Design for Risk Evaluation And Management) if `WriteDreamOutput` is set to `True` (see below). DREAM is designed to optimize the placement of monitoring sensors.

Unlike most other plot types, the TTFD plot type has two required entries: `PlumeType` and `ComponentNameList`. TTFD plots will not be produced without appropriate input for these entries. `ComponentNameList` is discussed above.

- `PlumeType` - the type of plume metric being considered. Acceptable values are *Pressure*, *pH*, *TDS*, *Dissolved\_CO2*, *Dissolved\_salt*, and *CarbonateAquifer*. The *dx*, *dy*, and *dz* metrics (e.g., **Dissolved\_CO2\_dz**) for the `PlumeType` used must be produced by the aquifer components listed in `ComponentNameList`. The *dz* metrics are not required when using *CarbonateAquifer* components, however, as these components do not produce a *dz* plume metric. Additionally, when using `PlumeType: CarbonateAquifer` the plume timing and plume probability figures do not have different subplots for different depth ranges.

The TTFD plot type can have the following optional entries: `MonitoringLocations`, `SaveCSVFiles`, `WriteDreamOutput`, `SpecifyXandYLims`, `NumZPointsWithinAquifers`, `NumZPointsWithinShales`, `xGridSpacing`, `yGridSpacing`, `SpecifyXandYGridLims`, `PlotInjectionSites`, `InjectionCoordx`, `InjectionCoordy`, `FigureDPI`, and `FigureSize`. Three of these entries (`MonitoringLocations`, `SpecifyXandYLims`, and `SpecifyXandYGridLims`) are dictionaries containing additional entries (i.e., entries indented beneath mentioned keywords in a *.yaml* file). All of these entries except for `MonitoringLocations`, `WriteDreamOutput`, `NumZPointsWithinAquifers`, and `NumZPointsWithinShales` are described above.

The `NumZPointsWithinAquifers`, `NumZPointsWithinShales`, `xGridSpacing`, `yGridSpacing`, and `SpecifyXandYGridLims` entries all relate to the *x*-, *y*-, and *z*-coordinates of the grids used to evaluate plume extents and timings. The *dx*, *dy*, and *dz* plume dimension metrics (e.g., *pH\_dy* or *TDS\_dz*) are used to evaluate whether each (*x*, *y*, *z*) of a grid is within a plume area for each model timestep. Note that `NumZPointsWithinAquifers` and `NumZPointsWithinShales` do not have an effect when `PlumeType` is given as *CarbonateAquifer* because a *CarbonateAquifer* component does not produce a *dz* plume metric.

- `MonitoringLocations` - a dictionary containing five optional entries related to the sensors used to detect aquifer impacts. The five optional entries are `coordx`, `coordy`, `coordz`, `HorizontalWindow`, and `VerticalWindow`. Note that the lists provided for `coordx`, `coordy`, and `coordz` must all have the same length (although `coordz` is not used with option `PlumeType: CarbonateAquifer`).
- `coordx` - an entry under `MonitoringLocations` that specifies the *x*-coordinate(s) (*m*) of monitoring sensor(s), if any sensors are used. This entry must be provided as a list, even if only one location is used (e.g., [100] or [100, 200]).
- `coordy` - an entry under `MonitoringLocations` that specifies the *y*-coordinate(s) (*m*) of monitoring sensor(s), if any sensors are used. This entry must be provided as a list, even if only one location is used (e.g., [100] or [100, 200]).
- `coordz` - an entry under `MonitoringLocations` that specifies the depth(s) (*z*-coordinate(s), (*m*)) of monitoring sensor(s), if any sensors are used. Note that for this entry, depths beneath the surface are taken as negative values. This entry must be provided as a list, even if only one location is used (e.g., [-500] or [-500, -400]). The `coordz` entry is not required when using an option `plumeType: CarbonateAquifer`, as the *CarbonateAquifer* component does not produce a *dz* plume metric.
- `HorizontalWindow` - a (maximum) horizontal distance (*m*) from which monitoring sensor(s) will detect plumes (default is 1). For example, if the `HorizontalWindow` is 5 m, then the sensor will detect any plume at grid locations within 5 m of the sensor's `coordx` and `coordy` values (if the plume is also within `VerticalWindow` of the sensor's `coordz` value). This entry is meant to represent the sensitivity of a sensor, but that consideration must also involve the threshold used for the plume type considered (if the aquifer component has a user-

defined threshold for plume detection). For example, **Dissolved\_salt** plumes from the `GenericAquifer` are influenced by the **dissolved\_salt\_threshold** parameter. In contrast, the `FutureGen2Aquifer` component defines TDS plumes where the relative change in TDS is  $> 10\%$  (i.e., no user-defined threshold). The inclusion of plumes at nearby grid points is also dependent on the spacing of grid points; the x- and y-spacings are controlled by `xGridSpacing` and `yGridSpacing`, while the z-spacing is controlled by `NumZPointsWithinAquifers` and `NumZPointsWithinShales`. Note that the grid is made to include the x-, y-, and z-coordinates for monitoring locations, so there will always be a grid point for each monitoring sensor.

- **VerticalWindow** - a (maximum) vertical distance ( $m$ ) from which monitoring sensor(s) will detect plumes (default is 1). For example, if the **VerticalWindow** is 5 m, then the sensor will detect any plume within 5 m of the sensor's `coordz` values (if the plume is also within **HorizontalWindow** of the sensor's `coordx` and `coordy` value). This entry is meant to represent the sensitivity of a sensor, but that consideration must also involve the threshold used for the plume type considered (if the aquifer component has a user-defined threshold for plume detection). For example, **Dissolved\_CO2** plumes from the `GenericAquifer` are influenced by the **dissolved\_co2\_threshold** parameter. In contrast, the `FutureGen2Aquifer` component defines pH plumes where the absolute change in pH is  $> 0.2$  (i.e., no user-defined threshold). The inclusion of plumes at nearby grid points is dependent on the spacing of grid points; the x- and y-spacings are controlled by `xGridSpacing` and `yGridSpacing`, while the z-spacing is controlled by `NumZPointsWithinAquifers` and `NumZPointsWithinShales`. Note that the grid is made to include the x-, y-, and z-coordinates for monitoring locations, so there will always be a grid point for each monitoring sensor.
- **WriteDreamOutput** - the option to create `.iam` files containing plume timing results (default is `False`). These `.iam` files are the input for the DREAM program. DREAM is the Design for Risk Evaluation And Management tool, which was also developed by NRAP. The only acceptable values are `True` or `False`.
- **NumZPointsWithinAquifers** - the number of z-grid points extending from the bottom to the top of each aquifer (default is 10). The points are equally spaced.
- **NumZPointsWithinShales** - the number of z-grid points extending from the bottom to the top of each shale (default is 3). The points are equally spaced. Note that the top of an aquifer is also the bottom of a shale, and the same location is not entered twice. In other words, with the default values for **NumZPointsWithinAquifers** (10) and **NumZPointsWithinShales** (3) a z-grid will have ten points from the bottom to the top of an aquifer, then a point in the middle of the overlying shale (point 2 of 3 across the shale), and then ten points from the bottom to the top of the overlying aquifer (etc.). In this example, including points 1 and 3 for the shale would be redundant because those points are included for the aquifers below and above the shale.

Below, we show two examples of TTFD plot entries in the `Plots`` section of a `*.yaml*` file. The first plot (`*pH_Minimum_Input*`) has only the entries required to set up the TTFD plot type: `PlumeType` and `ComponentNameList`. The second plot (`TDS_All_Options_Specified.tiff`) includes all optional entries for the TTFD plot type. Although there are only two plot entries included, each entry can result in the creation of multiple figures (e.g., earliest plume timings, TTFD from monitoring locations, and plume probabilities for each model realization). Note that all entries for the TTFD plot type are indented under `TTFD` which is indented under the figure name.

```

1 Plots:
2   pH_Minimum_Input:
3     TTFD:
4       PlumeType: pH
5       ComponentNameList: [FutureGen2AZMI1, FutureGen2Aquifer1]
6   TDS_All_Options_Specified.tiff:
7     TTFD:
8       PlumeType: TDS
9       ComponentNameList: [FutureGen2AZMI1, FutureGen2Aquifer1]
10      FigureDPI: 300
11      MonitoringLocations:
12        coordx: [100, 200]
```

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```

13         coordy: [100, 200]
14         coordz: [-407.5, -407.5]
15         HorizontalWindow: 1
16         VerticalWindow: 5
17     PlotInjectionSites: True
18     InjectionCoordx: 50
19     InjectionCoordy: 50
20     SpecifyXandYLims:
21         xLims: [-25, 700]
22         yLims: [-25, 700]
23     NumZPointsWithinAquifers: 10
24     NumZPointsWithinShales: 3
25     xGridSpacing: 5
26     yGridSpacing: 5
27     SpecifyXandYGridLims:
28         gridXLims: [25, 650]
29         gridYLims: [25, 650]
30     WriteDreamOutput: False
31     SaveCSVFiles: True

```

For examples of TTFD plots, see *ControlFile\_ex39a.yaml* to *ControlFile\_ex43.yaml*. For script examples, see *iam\_sys\_reservoir\_mswell\_futuregen\_ttfplot\_no\_dip.py*, *iam\_sys\_reservoir\_mswell\_futuregen\_ttfplot\_no\_dip\_lhs.py*, and *iam\_sys\_reservoir\_mswell\_futuregen\_ttfplot\_dipping\_strata.py*.

## 2.5.6 GriddedMetric

The GriddedMetric plot type produces map view images of a gridded metric. While the radial metrics shown by the GriddedRadialMetric plot type are defined in relation to radius and depth values, the metrics shown by the GriddedMetric plot type are defined relative to x-coordinates and y-coordinates. For example, the GriddedMetric plot type can display the gridded output produced by SealHorizon and FaultFlow components.

**The GriddedMetric plot type has two required entries: ComponentNameList and MetricName.** Both are described above.

The GriddedMetric plot type has the following optional entries: Realization, TimeList, PlotInjectionSites, InjectionCoordx, InjectionCoordy, SpecifyXandYLims, SaveCSVFiles, EqualAxes, FigureDPI, and FigureSize. All of these entries are discussed above.

Below, we show two examples of setting up GriddedMetric plots in a .yaml control file. The first plot (*Plot\_Default\_Settings*) includes only the required entries, while the second (*Plot\_With\_Options*) includes all optional entries.

```

1 Plots:
2   Plot_Default_Settings:
3     GriddedMetric:
4       ComponentNameList: [Fault1]
5       MetricName: mass_brine_aquifer
6   Plot_With_Options:
7     GriddedMetric:
8       ComponentNameList: [Fault1]
9       MetricName: CO2_aquifer
10      Realization: 0
11      FigureDPI: 300

```

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```

12      TimeList: [1, 5, 10, 25, 50]
13      SaveCSVFiles: False
14      PlotInjectionSites: False
15      InjectionCoordx: 4.68e+04
16      InjectionCoordy: 5.11e+04
17      SpecifyXandYLims:
18          xLims: [38750, 40500]
19          yLims: [48266, 48400]
20      EqualAxes: False

```

For examples of GriddedMetric plots, see *ControlFile\_ex18.yaml*, *ControlFile\_ex19.yaml*, and *ControlFile\_ex23.yaml*.

### 2.5.7 GriddedRadialMetric

The GriddedRadialMetric plot type produces map view images of a gridded radial metric. The GenericAquifer produces four kinds of gridded radial metrics: **r\_coordinate**, **z\_coordinate**, **Dissolved\_CO2\_mass\_fraction**, and **Dissolved\_salt\_mass\_fraction**. Regions of an aquifer with dissolved CO<sub>2</sub> and dissolved salt mass fractions exceeding the corresponding mass fraction threshold are included in the plume volumes for the corresponding plume type. Those plume volumes can be visualized with the TTFD plot type. The GriddedRadialMetric plot type, however, can show more general changes in dissolved CO<sub>2</sub> and salt mass fractions (e.g., seeing changes in mass fractions below the plume definition thresholds).

The GriddedRadialMetric plot type has three required entries: **ComponentNameList**, **ZList**, and **MetricName**. **ComponentNameList** and **MetricName** are discussed above.

- **ZList** - the depths (*m*) at which to evaluate the radial metric output. The depth in the radial grid (e.g., **z\_coordinate** from GenericAquifer) that is closest to each value entered will be used. String inputs representing the bottom of a unit can also be provided. For example, the bottom and top depths of aquifer 2 can be set up by entering **ZList**: [aquifer2Depth, shale3Depth]. Shale 3 is on top of aquifer 2, so the bottom depth of shale 3 is the top depth of aquifer 2. Note that numeric values given for **ZList** (not string inputs like shale2Depth) are taken as being negative when they represent a depth beneath the surface (e.g., **ZList**: [-500, -400] for depths of 500 *m* and 400 *m*).

The GriddedRadialMetric plot type has 11 optional entries: **MinValue**, **DegreeInterval**, **Realization**, **TimeList**, **PlotInjectionSites**, **InjectionCoordx**, **InjectionCoordy**, **SpecifyXandYLims**, **SaveCSVFiles**, **EqualAxes**, **FigureDPI**, and **FigureSize**. All of these entries except for **MinValue** and **DegreeInterval** are discussed above.

- **MinValue** - the minimum value used for the colorbar on the figures. Any values beneath this minimum will not be displayed graphically, but the entire range of values is still displayed in the title of each figure. This parameter has a significant impact on GriddedRadialMetric figures. For example, the **Dissolved\_CO2\_mass\_fraction** and **Dissolved\_salt\_mass\_fraction** outputs saved by a GenericAquifer for a time of 0 years will all have values of zero. The outputs saved at other times, however, can have very low but nonzero values. The **Dissolved\_CO2\_mass\_fraction** values can be as low as 5.0e-3 at the highest radii, while the **Dissolved\_salt\_mass\_fraction** values can be as low as 1.0e-30. If the **MinValue** provided is zero, then the figures created will be zoomed out to encompass such low values at the highest radii evaluated (about 77.5 km). These large extents will make the figures visually unclear. For these figures to be useful, one should specify a **MinValue** that is high enough to enable the figure to focus on the area of interest (i.e., near the component's location) but low enough to not exclude too much of the output data. We recommend using a **MinValue** of 0.002 when evaluating **Dissolved\_salt\_mass\_fraction** (10 times lower than the default **dissolved\_salt\_threshold** of 0.02) and 0.01 when evaluating **Dissolved\_CO2\_mass\_fraction** (equal to the default **dissolved\_salt\_threshold** of 0.01). If **MinValue** is not entered, these values will be used as the defaults for the corresponding output type (dissolved salt or dissolved CO<sub>2</sub>). If all of the times evaluated only have values less or equal to **MinValue**, then

one figure will be made. This figure has a title that includes 'All Model Times.' Note that the .csv files saved when SaveCSVFiles is set to True will only include values above MinValue.

- **DegreeInterval** - the interval (degrees) used to create a map-view image from the radial output. The accepted values are 1, 5, 10, 15, 30, and 45. If DegreeInterval is not entered, the default values is 15 degrees.

Note that although the Realization entry for the GriddedRadialMetric plot type follows the indexing conventions of Python (i.e., Realization: 0 for the first realization), the figure files and .csv files saved by the GriddedRadialMetric plot type will present the simulation number as ranging from one to the total number of realizations (e.g., Simulation 1 instead of Simulation 0).

Below, we show two examples of GriddedRadialMetric plot entries in a control file. The first entry (*Min\_Input\_Dissolved\_Salt*) uses the minimum input required for the GriddedRadialMetric plot type. The second entry (*All\_Input\_Dissolved\_Salt*) uses all entries available for the GriddedRadialMetric plot type.

```

1  Plots:
2      Min_Input_Dissolved_Salt:
3          GriddedRadialMetric:
4              ComponentNameList: [GenericAquifer1]
5              MetricName: Dissolved_salt_mass_fraction
6              ZList: [aquifer2Depth]
7      All_Input_Dissolved_Salt:
8          GriddedRadialMetric:
9              ComponentNameList: [GenericAquifer1]
10             MetricName: Dissolved_salt_mass_fraction
11             ZList: [aquifer2Depth, shale3Depth]
12             TimeList: [1, 5, 10, 15, 20]
13             MinValue: 0.002
14             FigureDPI: 300
15             PlotInjectionSites: True
16             InjectionCoordx: 100
17             InjectionCoordy: 100
18             DegreeInterval: 1
19             Realization: 0
20             EqualAxes: True
21             SaveCSVFiles: True
22             SpecifyXandYLims:
23                 xLims: [-200, 400]
24                 yLims: [-200, 400]

```

For examples of GriddedRadialMetric plots, see *ControlFile\_ex53a.yaml* to *ControlFile\_ex53d.yaml*

## 2.5.8 AtmPlumeSingle

The AtmPlumeSingle plot type produces map view images depicting how CO<sub>2</sub> leakage at the surface creates atmospheric CO<sub>2</sub> plumes. These images are created for each time step during one realization of a simulation. Note that simulations using the Latin Hypercube Sampling (lhs) or Parameter Study (parstudy) analysis types have many realizations, while a simulation using a forward analysis type only has one realization. For the AtmPlumeSingle plot type with lhs or parstudy simulations, the visualization corresponding to the realization of interest can be specified with the Realization entry in the .yaml file (discussed above). Note that using the AtmPlumeSingle plot type requires the use of an AtmosphericROM component.

Here is an example of the ModelParams section from *ControlFile\_ex40.yaml*, where the number of LHS realizations is set as siz: 30.

```

1 ModelParams:
2   EndTime: 15.
3   TimeStep: 1
4   Analysis:
5     type: lhs
6     siz: 30
7   Components: [LookupTableReservoir1, MultisegmentedWellbore1,
8               FutureGen2AZMI1, FutureGen2AZMI2]
9   OutputDirectory: output/output_ex40_{datetime}
10  Logging: Info

```

The produced figures show the source of the CO<sub>2</sub> leak as a red circle and the plume as a blue circle. The source location(s) are set by the x and y coordinate(s) of the component that the AtmosphericROM is connected to. For example, in *ControlFile\_ex9a.yaml*, the AtmosphericROM component is connected to an OpenWellbore component and the OpenWellbore component has its locations entered with coordx and coordy. The coordx and coordy values serve as the coordinates of sources for the AtmosphericROM component. In the AtmPlumeSingle figures, the coordx and coordy values are shown as the CO<sub>2</sub> sources. In the final figures the plumes are labeled as *Critical Areas* because the area is defined as being within the **critical\_distance** output (from an AtmosphericROM) from the corresponding source. The critical areas are, therefore, the areas in which the CO<sub>2</sub> concentrations exceed the value defined by the parameter **CO\_critical**. The **critical\_distance** is the radius of each plume circle shown in AtmPlumeSingle plots, and this **critical\_distance** is also displayed on the figure with text.

Note that when multiple atmospheric plumes overlap enough, they will be displayed as one plume. The source shown will be between the sources of each individual plume.

AtmosphericROM components can be provided with receptor locations, which are meant to represent home or business locations where people will be present. If receptors are provided and the *.yaml* input for the AtmPlumeSingle includes the entry PlotReceptors: True, then receptor locations will be shown.

The AtmPlumeSingle plot type can have the following optional entries Realization, PlotReceptors, PlotInjectionSites, InjectionCoordx, InjectionCoordy, SpecifyXandYLims, FigureDPI, and FigureSize. All of these entries except for PlotReceptors are described above.

- PlotReceptors - option to plot receptor locations (default is False). The only acceptable values are True or False. If the receptors are far away from the source location(s) and/or the injection site, plotting the receptors may cause the x and y limits to be spread too far. The plumes may then be difficult to see.

Below is an example of the AtmPlumeSingle plot input in a *.yaml* control file. Note that InjectionCoordx and InjectionCoordy only have to be provided when using a LookupTableReservoir and setting PlotInjectionSites: True.

```

1 Plots:
2   ATM_single:
3     AtmPlumeSingle:
4       Realization: 10
5       FigureDPI: 300
6       PlotInjectionSites: True
7       InjectionCoordx: 3.68e4
8       InjectionCoordy: 4.83e4
9       PlotReceptors: True
10      SpecifyXandYLims:
11        xLims: [3.58e4, 3.78e4]
12        yLims: [4.73e4, 4.93e4]

```

For examples of AtmPlumeSingle plots, see *ControlFile\_ex9a.yaml* to *ControlFile\_ex9c.yaml*.

## 2.5.9 AtmPlumeEnsemble

The `AtmPlumeEnsemble` plot type can only be used in simulations with Latin Hypercube Sampling (`lhs`) or Parameter Study (`parstudy`) analysis types. This plot type involves concepts similar to those as those of the `AtmPlumeSingle` plot type. While the `AtmPlumeSingle` plot type displays the critical areas for one realization, the `AtmPlumeEnsemble` plot type displays the probability of critical areas occurring in the domain. These probabilities are calculated with the results from all realizations of the `lhs` or `parstudy` simulation. The probabilities specifically represent the likelihood of CO<sub>2</sub> plume concentrations exceeding the threshold set with the `CO_critical` parameter for `AtmosphericROM` components. The probabilities are shown as gridded data. The `AtmPlumeEnsemble` plot type is available only when the simulation includes an `AtmosphericROM` component.

The `AtmPlumeEnsemble` plot type has the optional entries `PlotReceptors`, `PlotInjectionSites`, `InjectionCoordx`, `InjectionCoordy`, `SpecifyXandYGridLims`, `xGridSpacing`, `yGridSpacing`, `SpecifyXandYLims`, `FigureDPI`, and `FigureSize`. All of these entries are described above.

Below is an example of a `AtmPlumeEnsemble` plot entry in a `.yaml` file:

```

1 Plots:
2   ATM_Ensemble.tiff:
3     AtmPlumeEnsemble:
4       FigureDPI: 300
5       PlotInjectionSites: True
6       InjectionCoordx: 200
7       InjectionCoordy: 200
8       PlotReceptors: False
9       xGridSpacing: 1
10      yGridSpacing: 1
11      SpecifyXandYGridLims:
12        gridXLims: [-100, 300]
13        gridYLims: [-100, 300]
14      SpecifyXandYLims:
15        xLims: [-125, 325]
16        yLims: [-125, 325]
```

For examples of `AtmPlumeEnsemble` plots, see `ControlFile_ex9a.yaml` and `ControlFile_ex9c.yaml`.

## 2.6 Analysis Options in the CFI

NRAP-Open-IAM uses the Model Analysis ToolKit (MATK) [9] for the basis of its probabilistic framework. More information about MATK can be found here: <http://dharp.github.io/matk/>. The MATK code repository can be found here: <https://github.com/dharp/matk>.

Parameter input and output interactions can be explored using the *Analysis* section of a `.yaml` control file. Correlation coefficients can be calculated using the `CorrelationCoeff` keyword. Parameter sensitivity coefficients for any output simulation value can be calculated using a Random-Balanced-Design Fourier Amplitude Sensitivity Test (RBD-Fast) technique. The control file keywords `SensitivityCoeff`, `MultiSensitivities`, `TimeSeriesSensitivity` can be used to access different sensitivity coefficient outputs. See control file examples `ControlFile_ex8a.yaml` to `ControlFile_ex8d.yaml` for more details regarding the use of the analysis section.

The Sensitivity Analysis is done with the SALib package [12]. For more information on the RBD-Fast technique see [30] and [27]. While not accessible through the control files, a scripting interface is provided to a Sobol sensitivity analysis [29].

## 2.7 Workflows in NRAP-Open-IAM

NRAP-Open-IAM simulations in the control file interface or GUI can be set up manually, with the user handling all details for the selected components (e.g., parameters, component connections, and outputs). Alternatively, the user can set up a simulation by selecting a workflow type. Each workflow type is made for a specific type of analysis. An analysis type can require certain types of components, and a workflow will automatically add and configure the required components. Additionally, the workflow will automatically set up the plots required for the analysis. By building a simulation from the blueprint of a workflow, the system model does not have to start from a blank slate. This approach is meant to streamline the analyses that are frequently conducted for geologic carbon storage sites.

For example, the system model created by the AoR workflow will always include a reservoir component, a wellbore component, and an aquifer component because those component types are required for an AoR analysis. The user does not need to specify that the wellbore component is connected to the reservoir component (i.e., it uses pressures and CO<sub>2</sub> saturations from the reservoir component), because that arrangement is assumed within the AoR workflow. By automatically handling such considerations, the AoR workflow allows AoR analyses to be conducted more quickly and easily.

### 2.7.1 Workflow types

The workflows currently available are `LeakageAssessment`, `AoR`, and `TTFD`. We discuss each of these workflow types below.

### 2.7.2 LeakageAssessment Workflow

The `LeakageAssessment` workflow is used to assess CO<sub>2</sub> and brine leakage rates given output from a reservoir component and the presence of (a) wellbore(s). For example, the wellbore component used could represent a legacy well. This workflow automatically adds and sets up a reservoir component and a wellbore component - no aquifer component is included. The `LeakageAssessment` workflow then makes four time series plots. Two of these plots show reservoir pressure and CO<sub>2</sub> saturation at the wellbore location(s) given. The other two plots show the leakage rates of CO<sub>2</sub> and brine through the wellbore(s) and to specific aquifer. Although this system model configuration is relatively simple, the `LeakageAssessment` workflow is meant to provide convenience while also preventing the errors that can occur when manually setting up a system model (e.g., mistyping a component's name).

### 2.7.3 AoR Workflow

The AoR workflow conducts an area of review analysis for a geologic carbon storage site. An AoR analysis evaluates the maximum potential impacts on the reservoir and an aquifer. The reservoir impacts considered are pressures and CO<sub>2</sub> saturations. The aquifer impacts considered are the contaminant plume volumes (e.g., pH and TDS plumes) that could arise if a leakage pathway was located at a particular location. Many hypothetical leakage pathways (wells) are considered, with their positions distributed across the entire study area. The AoR is determined as the area in which the hypothetical wells could cause the aquifer to be contaminated. In other words, the AoR includes any well location that, in the simulation, had a nonzero plume volume in the aquifer, a nonzero CO<sub>2</sub> saturation in the reservoir, or a reservoir pressure exceeding the critical pressure. The critical pressure is calculated as the pressure required to drive flow from the reservoir into the aquifer being considered. A critical pressure can be automatically calculated based on the reservoir depth, aquifer depth, and a specified brine density, or it can be given as a specific value.

Note that the AoR workflow is not the same as the AoR plot type (discussed in section [Visualization Options in the CFI](#)). While the AoR plot type only shows the results for one metric (e.g., reservoir pressure), the AoR workflow assesses all four metrics considered (four separate AoR plots) and creates an overall AoR that reflects all of these metrics.

AoR analyses can be conducted with any type of wellbore component, but we recommend using the `OpenWellbore` component. This component portrays an uncemented wellbore, which represents a worst-case scenario for leakage risks.



When an AoR might contain undocumented legacy wells, it can be best to assume a conservatively high portrayal of leakage risks. For example, if the undocumented legacy wells were assumed to be cemented, but some are uncemented or some have a higher effective permeability than assumed in the simulations, then the leakage risks would be underestimated and the resulting AoR would be too small. Additionally, the `OpenWellbore` component can utilize a critical pressure in leakage calculations. Other wellbore components (e.g., `MultisegmentedWellbore` and `CementedWellbore`) do not use a critical pressure. When using one of these wellbore components, the critical pressure would still be shown in the pressure AoR plot (i.e., one of the five map-view .figures created by the AoR workflow - this figure only shows results from the reservoir component). The critical pressure would not, however, be reflected in the pH and TDS plume volume AoR plots (two of the five map-view figures). In other words, these plume volumes result from CO<sub>2</sub> and brine leakage into the aquifer, and this leakage is calculated by an `OpenWellbore`, `MultisegmentedWellbore`, or `CementedWellbore` component. The `MultisegmentedWellbore` and `CementedWellbore` components will calculate leakage in their default manner, without a user-specified critical pressure. Nonetheless, the user may want to use one of these components if they have constraints on the effective permeabilities of legacy wells in the area.

If the critical pressure ( $P_{crit}$ ) is not given as a specific value, it is calculated with the approach shown in section [Equations](#). When using the AoR workflow, the critical pressure input provided is used in both (1) the map-view figure of reservoir pressures, which impacts the overall AoR figure, and (2) the leakage calculations of `OpenWellbore` components (if one is used). If the AoR workflow is using a wellbore component with a brine density parameter (e.g., **brineDensity** parameter of the `OpenWellbore` and `MultisegmentedWellbore` components), then that parameter will be used for the brine density value in the critical pressure equation. If the wellbore component does not have a brine density parameter (e.g., `CementedWellbore`), then a brine density can be specified with the `BrineDensity` entry (discussed further below).

While the AoR plot type can be used with a `TheisReservoir`, the AoR workflow cannot be used with a `TheisReservoir`. When using a `TheisReservoir`, AoR plots can only be made for the **pressure** output (see *ControlFile\_ex47*). The AoR workflow is meant to incorporate data from all of the metrics that can be used with the AoR plot type (i.e., plume volumes and **CO2saturation**), so the AoR workflow cannot be used with a `TheisReservoir`.

## 2.7.4 TTFD Workflow

The TTFD workflow conducts a time to first detection analysis. This analysis type examines the spread of a contaminant plume through an aquifer, with multiple map-view figures showing how the plume spreads over time. The time to first detection is obtained by providing the locations and depths of monitoring wells. The time at which a plume first reaches a monitoring well is taken as the TTFD. If the monitoring well locations used do not detect a plume, then the locations should be changed. This workflow can also produce the files used as input for DREAM (Designs for Risk Evaluation and Management), a tool designed to optimize monitoring network design at geological carbon storage sites.

The TTFD plot type, which creates map-view images of aquifer plume timings across the domain, exists separately from the TTFD workflow. The TTFD workflow is meant to offer convenience by automatically setting up the components, component connections, and plot entries (in a .yaml file) required for use of the TTFD plot type.

## 2.7.5 Setup of a Workflow in the Control File Interface

In a standard control file, there are separate sections for stratigraphy, model parameters, each component, and the plots to be produced. A control file using a workflow, however, can accomplish complex analyses while only using three sections: a stratigraphy section, a model parameters section, and a workflow section. We show an example of this setup below.

```

1 #-----
2 ModelParams:
3     EndTime: 20
4     TimeStep: 1.0
5     Analysis: forward

```

(continues on next page)

(continued from previous page)

```

6   Components: []
7   OutputDirectory: 'output/output_ex55a_{datetime}'
8   GenerateOutputFiles: False
9   GenerateCombOutputFile: False
10  Logging: Debug
11  #-----
12  Stratigraphy:
13    numberOfShaleLayers:
14      value: 5
15      vary: False
16    shale1Thickness:
17      value: 198.7
18      vary: False
19    shale2Thickness:
20      value: 74.4
21      vary: False
22    shale3Thickness:
23      value: 110.3
24      vary: False
25    aquifer1Thickness:
26      value: 33.2
27      vary: False
28    aquifer2Thickness:
29      value: 84.1
30      vary: False
31    reservoirThickness:
32      value: 7.0
33      vary: False
34  #-----
35  Workflow:
36    Type: LeakageAssessment
37    Options:
38      ReservoirComponentType: AnalyticalReservoir
39      WellboreComponentType: OpenWellbore
40      ReservoirOptions:
41        InjectionWell:
42          coordx: 10
43          coordy: 50
44        Parameters:
45          injRate: 0.1
46      WellboreOptions:
47        Locations:
48          coordx: [500]
49          coordy: [500]
50        Parameters:
51          wellRadius: 0.05
52      AquiferName: aquifer2
53      FigureDPI: 300
54  #-----

```

The ModelParams and Stratigraphy sections are set up in the same manner as a normal control file (see section [Control File Interface](#)). The Workflow section contains two entries: Type and Options. The Type entry is followed by one of the available workflow types. Here, the Type is LeakageAssessment. The Options entry is a dictionary



containing more entries within it (i.e., in a *.yaml* control file, the contained entries are on a line beneath `Options` and preceded by four additional spaces). The remaining details regarding the simulation are entered under `Options` (e.g., well locations, component parameters, and plotting options). Any options that are not given will be set at their default settings. For the sake of brevity, the example above does not show all of the entries that can be provided under `Options`. Some of the `Options` entries available depend on the workflow type, and the entries specific to each workflow are discussed further below.

The `Options` entries that are applicable across multiple workflow types are:

- `ReservoirComponentType` - the type of reservoir component to use. Acceptable inputs include `LookupTableReservoir` and `AnalyticalReservoir`. Any analysis conducted for a real site should use a `LookupTableReservoir` to incorporate results from high-fidelity reservoir simulations; the remaining reservoir component types are largely intended for testing purposes.
- `WellboreComponentType` - the type of wellbore component to use. Acceptable inputs include `OpenWellbore`, `MultisegmentedWellbore`, and `CementedWellbore`.
- `AquiferComponentType` - the type of aquifer component to use. Acceptable inputs include `GenericAquifer`, `FutureGen2Aquifer`, `FutureGen2AZMI`, and `DeepAlluviumAquifer`.
- `ReservoirOptions` - an entry containing other entries related to the reservoir component. All of the input provided in the reservoir component's section of a standard control file are included here (e.g., the `Parameters` and `InjectionWell` entries).
- `WellboreOptions` - an entry containing other entries related to the wellbore component. All of the input provided in the wellbore component's section of a standard control file are included here (e.g., the `Locations`, `LeakTo`, `Controls`, `ThiefZone`, and `Parameters` entries). For examples of how to set up entries like `Locations` and `Parameters`, see section [Control File Interface](#). When using an `OpenWellbore` component in a workflow, the default approach is to include the `Controls` section with `critPressureApproach` set to `True` and `enforceCritPressure` set to `False`. With these settings, the `OpenWellbore` component will automatically use a critical pressure calculated with the `brineDensity` parameter, the bottom depth of the aquifer receiving leakage, and the top depth of the reservoir.
- `AquiferOptions` - an entry containing other entries related to the aquifer component. All of the input provided in the aquifer component's section of a standard control file are included here (e.g., the `Parameters` entry).
- `AquiferName` - the name of the aquifer being considered in the analysis, formatted as `aquifer2` (where 2 is replaced by the desired aquifer number). If this entry is provided under the `AquiferOptions` entry, then that input will overwrite any input given for `AquiferName` under the `Options` entry. If not provided, the default setting is the highest aquifer.
- `PlotInjectionSites` - the option to plot injection sites on any map-view figures created. The only acceptable values are `True` or `False`. The default setting is `False`.
- `InjectionCoordx` - value or list of values for the x coordinate(s), or easting distance(s), of the injection site(s) (default is `None`). The values are given in meters. This entry must be provided when using a `LookupTableReservoir`, as that component type does not have an `.injX` attribute. Other reservoir types like `AnalyticalReservoir` can be displayed without an `InjectionCoordx` entry.
- `InjectionCoordy` - value or list of values for the y coordinate(s), or northing distance(s), of the injection site(s) (default is `None`). The values are given in meters. This entry must be provided when using a `LookupTableReservoir`, as that component type does not have an `.injY` attribute. Other reservoir types like `AnalyticalReservoir` can be displayed without an `InjectionCoordy` entry.
- `FigureDPI` - the dots-per-inch (DPI) of the figure(s) produced (default is 100). Larger DPIs will create high-resolution, high-quality figures, but the file sizes are also larger. File size and quality are also influenced by the extension used (e.g., `.png` or `.tiff`). Recommended `FigureDPI` values are between 100 and 300.
- `FigureSize` - the width and height of the figures produced in inches as a list of length two (e.g., `FigureSize: [14, 8]`), where the first number is the width and the second number is the height.

- **FigureName** - the name of the figure to be produced. If the name is given with an extension, then the figure will be saved using that file extension (e.g., `FigureName: Overall_AoR_Plot.tiff` for a .tiff file named `Overall_AoR_Plot`). If this entry is not provided, the default figure names will be used. If this entry is provided for a workflow that produces multiple figures (e.g., the `LeakageAssessment` and `TTFD` workflows), the name will not be used but any extension included will be used.
- **AutomateResCompSetup** - the option specifying whether the workflow should automatically set up the section for the reservoir component in the .yaml file. The only acceptable values are `True` or `False`, and the default setting is `True`.
- **AutomateWellCompSetup** - the option specifying whether the workflow should automatically set up the section for the wellbore component in the .yaml file. The only acceptable values are `True` or `False`, and the default setting is `True`.
- **AutomateAqCompSetup** - the option specifying whether the workflow should automatically set up the section for the aquifer component in the .yaml file. The only acceptable values are `True` or `False`, and the default setting is `True`.
- **AutomatePlotsSetup** - the option specifying whether the workflow should automatically set up the `Plots` section of the .yaml file. The only acceptable values are `True` or `False`, and the default setting is `True`. If set to `False`, then the `Plots` section of the .yaml control file must be set up by the user.

## 2.7.6 Setup of the LeakageAssessment Workflow in the Control File Interface

To use the `LeakageAssessment` workflow, set the `Type` entry of the `Workflow` section to `LeakageAssessment`.

The `LeakageAssessment` workflow requires the `Locations` input under `WellboreOptions`:

```

1  #-----
2  Workflow:
3      Type: LeakageAssessment
4      Options:
5          PlotType: TimeSeriesStats
6          FigureDPI: 300
7          UseMarkers: True
8          Subplot:
9              Use: True
10             NumCols: 2
11          FigureSize: [14, 8]
12          FigureName: Name.tiff  # Only the .tiff extension from this will be used
13          ReservoirComponentType: AnalyticalReservoir
14          ReservoirOptions:
15              InjectionWell:
16                  coordx: 0
17                  coordy: 0
18              Parameters:
19                  injRate: 0.5
20                  reservoirRadius: 2000
21                  logResPerm: -13.5
22                  brineDensity: 1100
23          WellboreComponentType: OpenWellbore
24          WellboreOptions:
25              Locations:
26                  coordx: [100, 200]
27                  coordy: [100, 200]

```

Here, the `Locations` entry is given `coordx` and `coordy` lists. Any valid input for `Locations` can be given under `WellboreOptions` (e.g., the `file`, `grid`, or `range` options; see *ControlFile\_ex30.yaml*).

In addition to the optional entries that are applicable to multiple workflows (discussed above), the `LeakageAssessment` workflow has the optional entries `PlotType`, `Subplot`, `UseMarkers`, `UseLines`, `VaryLineStyle`, and `FigureSize` (all of which are entered under `Options`). All of these entries except for `PlotType` are options for the `TimeSeries` plot type that control the appearance of a time series plot (e.g., specifying whether to use multiple subplots). For more information about those entries, see section *Visualization Options in the CFI*.

- `PlotType` - option specifying the type of time series plot. The options are `TimeSeries`, `TimeSeriesStats`, and `TimeSeriesAndStats`. For more details regarding these plot types, see section *Visualization Options in the CFI*.

The figures produced will focus on the leakage received by the aquifer specified through the `AquiferName` entry discussed above. If that entry is not provided, then the default setting is the highest aquifer.

For examples of the `LeakageAssessment` workflow in the control file interface, see *ControlFile\_ex58a.yaml* to *ControlFile\_ex58c.yaml*.

## 2.7.7 Setup the AoR Workflow in the Control File Interface

To use the AoR workflow, set the `Type` entry of the `Workflow` section to `AoR`.

When using the AoR workflow, we recommend setting `GenerateOutputFiles` and `GenerateCombOutputFile` to `False` in the `ModelParams` section of the `.yaml` file. The large number of wellbore locations commonly used for AoR plots causes a large number of output files. A reservoir and aquifer component is created for each wellbore location, and every component will have its output saved. The `.csv` files created by the AoR workflow contain all of the necessary information and these files are much smaller in size.

In addition to the optional entries that are applicable to multiple workflows (discussed above), the `AoR Workflow` has the optional entries `TimeList`, `CriticalPressureMPa`, and `BrineDensity`. These entries can be provided under `Options`. The `TimeList` and `BrineDensity` entries have the same impact on both the AoR plot type and the AoR workflow; for more details on these entries, refer to the section *Visualization Options in the CFI*. The `CriticalPressureMPa` entry has a different impact when used for an AoR plot entry or for the AoR workflow, however, so we describe the `CriticalPressureMPa` entry for the AoR workflow below.

- `CriticalPressureMPa` - this entry controls how critical pressure is handled in the AoR analysis. The entry can either be given as `Calculated` or as a number representing a critical pressure in MPa (e.g., `CriticalPressureMPa: 20.5` for 20.5 MPa). The AoR plot type also has an entry called `CriticalPressureMPa`, but when that entry is used for an AoR plot it only has an impact if the AoR plot analyzes the *pressure* metric (i.e., no impact on plots evaluating  $\text{CO}_2$  saturations or aquifer plume volumes). When `CriticalPressureMPa` is given for the AoR workflow, however, this setting impacts both the critical pressures used by an `OpenWellbore` component (if one is used) and the critical pressures used in AoR plots evaluating the *pressure* metric.

Below, we show an example of an AoR workflow in the control file interface. Only the `Workflow` section is shown (the `ModelParams` and `Stratigraphy` sections are not shown).

```

1  #-----
2  Workflow:
3      Type: AoR
4      Options:
5          FigureName: Overall_AoR_Plot.tiff
6          CriticalPressureMPa: Calculated
7          ReservoirComponentType: LookupTableReservoir
8          ReservoirOptions:

```

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```

9       FileDirectory: source/components/reservoir/lookuptables/FutureGen2/1008_sims
10      TimeFile: time_points.csv
11      ParameterFilename: parameters_and_filenames_trunc.csv
12      Interpolation2D: False
13      Parameters:
14          index: 1
15      WellboreComponentType: OpenWellbore
16      WellboreOptions:
17          Locations:
18              file: source/components/reservoir/lookuptables/FutureGen2/1008_sims/fg1.csv
19              read_z_values: True
20              thin_point_density: True
21              min_x_spacing: 10000
22              min_y_spacing: 10000
23          Parameters:
24              wellRadius: 0.05
25              logReservoirTransmissivity: -10.0
26              logAquiferTransmissivity: -10.0
27              brineSalinity: 0.0475
28              brineDensity: 1075
29      AquiferName: aquifer4
30      AquiferComponentType: FutureGen2Aquifer
31      AquiferOptions:
32          Parameters:
33              por: 0.18
34              log_permh: -11.92
35              log_aniso: 0.3

```

Unlike the LeakageAssessment and TTFD workflows, the AoR workflow can be run without explicitly setting the locations of wellbores. Specifically, the user does not have to enter x and y values through the Locations entry under WellboreOptions, as discussed above. The user can enter that information, but if that input is not provided then there are default settings for the AoR workflow. When using a LookupTableReservoir, the default x and y values for the wellbore components are taken directly from the .csv files tied to the LookupTableReservoir component (i.e., by using the file entry under Locations; see *ControlFile\_ex30.yaml*). Because the point densities of reservoir models generally become quite high near an injection site, the default approach is to thin the point density by enforcing a minimum point spacing of 20 km in the x and y directions (the min\_x\_spacing and min\_y\_spacing entries shown above). This large spacing is the default setting because these simulations can have long run times. Reducing the point density of wellbore locations reduces the number of reservoir, wellbore, and aquifer components used. If the reservoir component is not a LookupTableReservoir, then the default wellbore locations are arranged in a six by six grid (total of 36 points) with x and y values ranging from -50 km to 50 km.

To manually define wellbore locations with a grid, use the grid entry under Locations. The grid entry contains the entries xmin, xmax, xsize, ymin, ymax, and ysize. The minimum x and y values (in meters) are defined by xmin and ymin, respectively. The maximum x and y values (in meters) are defined by xmax and ymax, respectively. Finally, the number of grid points in the x and y directions are defined by xsize and ysize, respectively.

```

1      WellboreOptions:
2          Locations:
3              grid:
4                  xmin: -7500
5                  xmax: 7500
6                  xsize: 14
7                  ymin: -7500

```

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```

ymax: 7500
ysize: 14

```

In a normal control file (i.e., one that does not use a workflow), an `OpenWellbore` component can only use a specific critical pressure (as opposed to a calculated one) if the user provides the `critPressure` parameter and sets both the `critPressureApproach` and `enforceCritPressure` entries under `Controls` to `True`. When using the `AoR` workflow that includes an `OpenWellbore` component, setting the `CriticalPressureMPa` entry to a specific value (e.g., `CriticalPressureMPa: 22.07` for 22.07 MPa) also sets the critical pressure for the `OpenWellbore` component. In this case, you do not need to manually set the `critPressure` parameter or the entries under `Controls` for the `OpenWellbore` component. The workflow automatically changes these settings, with the `CriticalPressureMPa` value being converted to *Pa* for the `critPressure` parameter. See *ControlFile\_ex56c.yaml* for an example where the critical pressure is set to a specific value.

Note that the `CriticalPressureMPa` input will not impact the leakage calculations of wellbore components like `MultisegmentedWellbore` or `CementedWellbore`, which do not use a critical pressure.

For examples of the `AoR` workflow in the control file interface, see *ControlFile\_ex55a.yaml* to *ControlFile\_ex56g.yaml*.

## 2.7.8 Setup the TTFD Workflow in the Control File Interface

To use the TTFD workflow, set the `Type` entry of the `Workflow` ``section to ``TTFD.

The TTFD Workflow requires the input of `Locations` data under `WellboreOptions`. Any valid input for `Locations` can be given under `WellboreOptions` (e.g., the `file`, `grid`, or `range` options; see *ControlFile\_ex30.yaml*).

The `PlumeType` entry plays an important role in the TTFD workflow.

- `PlumeType` - the type of plume metric being considered. Acceptable values are *Pressure*, *pH*, *TDS*, *Dissolved\_CO2*, *Dissolved\_salt*, and *CarbonateAquifer*. The plume type used must be compatible with the type of aquifer component used. For example, the default aquifer component is `GenericAquifer`, and that component only produces *Dissolved\_CO2* and *Dissolved\_salt* plumes. Using a plume type of *pH* would cause an error when using a `GenericAquifer`. The default plume type is *pH* when using a `FutureGen2Aquifer`, `FutureGen2AZMI`, `DeepAlluviumAquifer`, or `DeepAlluviumAquiferML` component. The default plume type is *Dissolved\_CO2* when using a `GenericAquifer` component. And the default plume type is *CarbonateAquifer* when using a `CarbonateAquifer` component.

Note that certain components define plume extents with fixed thresholds. For example, `FutureGen2Aquifer` and `FutureGen2AZMI` components define *pH* plumes as areas with changes in pH of greater than 0.2. In contrast, certain components allow the user to set the thresholds for plume definition. For example, the `GenericAquifer` defines *Dissolved\_CO2* plumes as areas with CO<sub>2</sub> mass fractions exceeding the `dissolved_co2_threshold` parameter. The `GenericAquifer` also allows the user to set the initial salinity of the aquifer, which impacts how easily the `dissolved_salt_threshold` parameter is exceeded and how large the *Dissolved\_salt* plumes become.

The TTFD workflow also has all of the optional entries that apply to the TTFD plot type (`MonitoringLocations`, `WriteDreamOutput`, `SaveCSVFiles`, `SpecifyXandYLims`, ``SpecifyXandYGridLims, `xGridSpacing`, `yGridSpacing`, `NumZPointsWithinAquifers`, and `NumZPointsWithinShales`). For the TTFD workflow, all of these entries are provided under `Options` in the `Workflow` section of a `.yaml` control file. For descriptions of all of these options, see the description of the TTFD plot type in section *Visualization Options in the CFI*.

The input provided under `MonitoringLocations` controls the positions and depths of the monitoring wells used. If `MonitoringLocations` is not provided, then the TTFD workflow will evaluate the evolution of contaminant plumes but not show any detection times. If `WriteDreamOutput` is set to `True`, then the TTFD workflow will save *.iam* files containing the plume timing results. These *.iam* files can be used as input to the DREAM tool (Design for Risk Evaluation and Management), which was also developed by NRAP.

Below, we show an example of the `Workflow` section of a `.yaml` control file using the TTFD workflow.

```

1  #-----
2  Workflow:
3      Type: TTFD
4      Options:
5          MonitoringLocations:
6              coordx: [200, 800, 200, 800]
7              coordy: [800, 200, 800, 200]
8              coordz: [-1015, -715, -1015, -715]
9          WriteDreadOutput: False
10         SaveCSVFiles: False
11         FigureDPI: 300
12         FigureName: Name.tiff           # specifies .tiff extension
13         PlumeType: Dissolved_salt
14         PlotInjectionSites: True
15         ReservoirComponentType: AnalyticalReservoir
16         ReservoirOptions:
17             InjectionWell:
18                 coordx: 500
19                 coordy: 500
20             Parameters:
21                 injRate: 0.1
22                 logResPerm: -13
23                 reservoirRadius: 5000
24         WellboreComponentType: OpenWellbore
25         WellboreOptions:
26             Controls:
27                 critPressureApproach: True # Both set to True, so the critPressure
28                 enforceCritPressure: True # parameter will be used
29             Locations:
30                 coordx: [400, 600]
31                 coordy: [600, 400]
32             Parameters:
33                 wellRadius: 0.05
34                 critPressure: 2.207e+7     # equivalent to 22.07 MPa
35         AquiferName: aquifer2
36         AquiferComponentType: GenericAquifer
37         AquiferOptions:
38             Parameters:
39                 por: 0.118
40                 log_permh: -13.39
41                 log_aniso: 0.3

```

For examples of the TTFD workflow in the control file interface, see *ControlFile\_ex57a.yaml* to *ControlFile\_ex57c.yaml*.



## 2.8 Equations

In this section, we present equations related to geologic carbon storage. While all component models have their own governing equations, information about each component can be found in chapter [Components Description](#). This section is only meant to address equations that are frequently referenced in this document.

### 2.8.1 Critical Pressure

Critical pressure can be used in AoR plots as well as the leakage calculations of the `OpenWellbore` component. If given a critical pressure, an AoR plot evaluating reservoir pressures will highlight any points where the pressures exceed the critical pressure. In this case, the AoR plot is only considering the output of a reservoir component (not a wellbore or aquifer component).

If the critical pressure ( $P_{crit}$ ) is not given as a specific value, it is calculated as [7]:

$$P_{crit} = (\rho_w \times g \times d_{aq}) + (\rho_{br} \times g \times (d_{res} - d_{aq})),$$

where  $\rho_w$  and  $\rho_{br}$  are the densities of water ( $1000 \text{ kg/m}^3$ ) and brine, respectively,  $g$  is gravitational acceleration ( $9.81 \text{ m/s}^2$ ),  $d_{aq}$  is the depth to the bottom of the aquifer impacted by leakage ( $m$ ), and  $d_{res}$  is the depth to the top of the reservoir ( $m$ ). Higher brine densities generally produce lower leakage rates; it is more difficult to lift brine from the reservoir to an aquifer when the brine has a higher density. In an AoR plot evaluating reservoir pressure,  $d_{aq}$  reflects the aquifer being considered in the AoR analysis. For an `OpenWellbore` component using a critical pressure,  $d_{aq}$  should reflect the aquifer receiving leakage from the wellbore (i.e.,  $d_{aq}$  is equal to the **wellTop** parameter) and  $\rho_{br}$  is set by the **brineDensity** parameter.

If an AoR plot is made for a simulation using a wellbore component that has a brine density parameter (e.g., **brineDensity** parameter of the `OpenWellbore` and `MultisegmentedWellbore` components), then that parameter will be used for the  $\rho_{br}$  value. If the wellbore component does not have a brine density parameter (e.g., `CementedWellbore`), then a brine density can be specified with the **BrineDensity** entry for the AoR plot type. This entry can also be used with the AoR workflow. Otherwise, a default brine density value of  $1045 \text{ kg/m}^3$  will be used.

## 2.9 Units

Data passed between models need to have consistent units. Here is a list of the units used by NRAP-Open-IAM.

- Pressure is assumed to be in units of Pascals ( $Pa$ ).
- Time is assumed to be in days.
- Distance, width, length, height, depth are assumed to be in units of meters ( $m$ ).
- Flow rates are assumed to be in units of kilograms per second ( $kg/s$ ).
- Mass is assumed to be in units of kilograms ( $kg$ ).
- Viscosities are assumed to be in units of Pascal seconds ( $Pa \cdot s$ ).
- Permeability is assumed to be in units of meters squared ( $m^2$ ).

Note that these rules apply to the exchange of data between component models in NRAP-Open-IAM during a simulation. There are input options in the CFI and GUI that take values in different units. For example, the AoR plot and AoR workflow have the input **CriticalPressureMPa**, which takes a pressure value in  $MPa$ . While NRAP-Open-IAM simulations use time in days, many temporal inputs provided through the CFI or GUI take time in years (before starting the simulation, the conversion to days is handled automatically). Any such distinctions are clearly noted in this document.



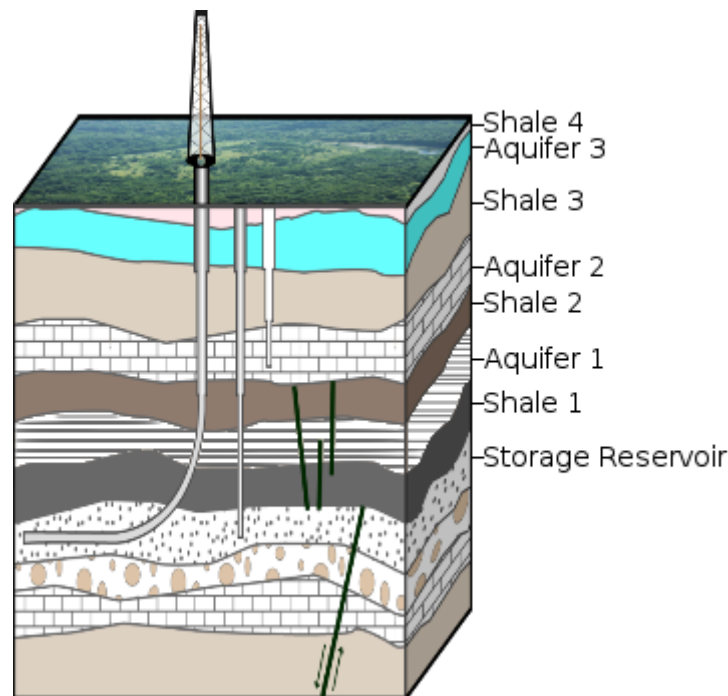


## COMPONENTS DESCRIPTION

This section of the document will describe each of the component models available for use in NRAP-Open-IAM. For each component model, all parameters that can be specified are described along with the parameters' units and acceptable ranges. Descriptions of the outputs that can be returned by each component are also provided.

### 3.1 Stratigraphy Component

The Stratigraphy component is a component containing parameters describing the structure of the storage reservoir system. The stratigraphy component allows the number of shale (or aquitard) layers to be set, thus, setting the total number of layers in the system. Each shale or aquifer layer can take on the default thickness for that layer type or be assigned a user defined value with `shale#Thickness` or `aquifer#Thickness` keywords where # is replaced by an index of the layer (e.g., `shale1Thickness`). Layers are numbered from the bottom upward: shale 1 is the layer above the storage reservoir, and, with N shale layers total, shale N is the surface layer.



This component represents flat-lying stratigraphy; when using this component, unit thicknesses and depths do not vary across the domain.

Descriptions of the component's parameters are provided below.

- **numberOfShaleLayers** [-] (3 to 30) - number of shale layers in the system (default: 3). The shale units must be separated by an aquifer.
- **shaleThickness** [*m*] (1 to 1600) - thickness of shale layers (default 250). Thickness of shale layer 1, for example, can be defined by **shale1Thickness**; otherwise, shale layers for which the thickness is not defined will be assigned a default thickness.
- **aquiferThickness** [*m*] (1 to 1600) - thickness of aquifers (default: 100). Thickness of aquifer 1, for example, can be defined by **aquifer1Thickness**; otherwise, aquifers for which the thickness is not defined will be assigned a default thickness.
- **reservoirThickness** [*m*] (1 to 1600) - thickness of reservoir (default: 50)
- **datumPressure** [*Pa*] (80,000 to 300,000) - pressure at the top of the system (default: 101,325)
- **depth** [*m*] (5 to 30,000) - depth to the top of reservoir (default: 950). The depth to the top of the reservoir can also be obtained through the **reservoirTopDepth** or **shale1Depth** composite parameters (the bottom of shale 1 is also the top of the reservoir).

The depths to the bottom, middle, and top of each unit are produced as composite parameters (i.e., calculated as functions of the thickness parameters used):

- **shale#Depth** [*m*] (boundaries depend on the user input) - depth to the base of the shale layer with index # (default value is not defined)
- **aquifer#Depth** [*m*] (boundaries depend on the user input) - depth to the base of the aquifer layer with index # (default value is not defined)
- **reservoirDepth** [*m*] (boundaries depend on the user input) - depth to the base of the reservoir (default value is not defined).
- **shale#MidDepth** [*m*] (boundaries depend on the user input) - depth to the middle of the shale layer with index # (default value is not defined)
- **aquifer#MidDepth** [*m*] (boundaries depend on the user input) - depth to the middle of the aquifer layer with index # (default value is not defined)
- **reservoirMidDepth** [*m*] (boundaries depend on the user input) - depth to the middle of the reservoir (default value is not defined).
- **shale#TopDepth** [*m*] (boundaries depend on the user input) - depth to the top of the shale layer with index # (default value is not defined)
- **aquifer#TopDepth** [*m*] (boundaries depend on the user input) - depth to the top of the aquifer layer with index # (default value is not defined)
- **reservoirTopDepth** [*m*] (boundaries depend on the user input) - depth to the top of the reservoir (default value is not defined).

Although these composite parameters are automatically added in the control file interface and graphical user interface, they must be added explicitly as composite parameters in script applications. For examples of such applications, see the script examples `iam_sys_strata.py`, `iam_sys_strata_reservoir_openwell_genericaquifer.py`, or `iam_sys_strata_reservoir_openwell_genericaquifer_5locs.py`.

## 3.2 Dipping Stratigraphy Component

The Dipping Stratigraphy component is similar to the Stratigraphy component, but it portrays dipping units. This component takes strike and dip values as well as unit thicknesses for a reference location. It then uses that input to produce unit thicknesses and depths at for the given output location(s). This component

The arrangement of geologic units for this component is the same as the that for the Stratigraphy component. The reservoir is the lowest unit. Alternating shale and aquifer layers overlie the reservoir. These units each have an index, with the index number increasing closer to the surface. Shale 1 is above the reservoir, aquifer 1 is above shale 1, and shale 2 is above aquifer 1. This pattern continues until shale N, where N is the number of shale units used (the *numberOfShaleLayers* parameter). There are N - 1 aquifers. The *numberOfShaleLayers* parameter cannot vary across the domain.

The *dipDirection* keyword can be used to specify the direction of dip. The *dipDirection* can be given as a cardinal direction (N, E, S, W, NE, SE, SW, and NW). In the control file interface, the *dipDirection* can be specified under the Controls entry, which is entered directly under DippingStratigraphy. For example, the *strike* and *dip* parameters could be given as 45 ° and 3 °, respectively. Because the strike is to the northwest, the *dipDirection* could be to the southeast or northwest (SE or NW). In this example, providing a *dipDirection* to the south or north (S or N) will also be taken as a dip to the southeast or northeast, respectively (i.e., a southwards component of dip vs. a northwards component). If the *dipDirection* keyword is not specified, then the *dipDirection* will be taken as 90 ° clockwise from the strike direction, if viewed in a map-view image (i.e., the right-hand rule in geology; when facing in the direction of strike, the dip direction is to your right).

In the NRAP-Open-IAM control file interface, the type name for the Dipping Stratigraphy component is *DippingStratigraphy*.

Descriptions of the component's parameters are provided below.

- **strike** [-] (0 to 360) - strike of the units in degrees (default: 315). In a map-view image, this parameter is taken as increasing from zero with rotation clockwise from north. For example, *strike* values of 0, 90, 180, and 270 would represent units striking to the north, east, south, and west, respectively.
- **dip** [-] (0.1 to 89.9) - dip of the units in degrees (default: 5). If the dip causes any unit thicknesses in the domain to fall outside the range of 1 m to 1600 m, that unit thickness will be set to the closest limit.
- **numberOfShaleLayers** [-] (3 to 30) - number of shale layers in the system (default: 3). The shale units must be separated by an aquifer.
- **shaleThickness** [m] (1 to 1600) - thickness of shale layers at the reference location (default: 250). Thickness of shale layer 1, for example, can be defined by **shale1Thickness**; otherwise, shale layers for which the thickness is not defined will be assigned a default thickness.
- **aquiferThickness** [m] (1 to 1600) - thickness of aquifers at the reference location (default: 100). Thickness of aquifer 1, for example, can be defined by **aquifer1Thickness**; otherwise, aquifers for which the thickness is not defined will be assigned a default thickness.
- **reservoirThickness** [m] (1 to 1600) - thickness of the reservoir at the reference location (default: 50)
- **datumPressure** [Pa] (80,000 to 300,000) - pressure at the top of the system (default: 101,325)

Unit thicknesses and the depths to the bottom, middle, and top of each unit at specific locations (x and y values) are produced as observations:

The observations from the Lookup Table Stratigraphy component are:

- **shale#Thickness** [m] - thickness of shale # at the output location(s), where # is an index ranging from one to *numberOfShaleLayers*.
- **aquifer#Thickness** [m] - thickness of aquifer # at the output location(s), where # is an index ranging from one to (*numberOfShaleLayers* - 1).

- **reservoirThickness** [*m*] - reservoir thickness at the output location(s).
- **shale#Depth** [*m*] - depth to the bottom of the shale unit with index # at the output location(s).
- **aquifer#Depth** [*m*] - depth to the bottom of aquifer unit with index # at the output location(s).
- **reservoirDepth** [*m*] - depth to the base of the reservoir at the output location(s).
- **shale#MidDepth** [*m*] - depth to the middle of the shale layer with index # at the output location(s).
- **aquifer#MidDepth** [*m*] - depth to the middle of the aquifer layer with index # at the output location(s).
- **reservoirMidDepth** [*m*] - depth to the middle of the reservoir at the output location(s).
- **shale#TopDepth** [*m*] - depth to the top of the shale unit with index # at the output location(s).
- **aquifer#TopDepth** [*m*] - depth to the top of the aquifer unit with index # at the output location(s).
- **reservoirTopDepth** [*m*] - depth to the top of the reservoir at the output location(s).

If the component produces a unit thickness for a shale, aquifer, or reservoir that falls outside the range of 1 *m* to 1600 *m*, then that thickness will be set to the lower or upper limit (whichever is closer). For example, if the unit thickness would be 1700 *m*, based on the parameters and output locations used, then that thickness observation will instead be set to 1600 *m*.

For control file examples using the `DippingStratigraphy` component, see `ControlFile_ex33a.yaml` to `ControlFile_ex35.yaml`, and `ControlFile_ex39b.yaml`. For script examples, see `iam_sys_dippingstrata.py`, `iam_sys_dippingstrata_gridded.py`, `iam_sys_reservoir_mswell_stratplot_dipping_strata.py`, and `iam_sys_reservoir_mswell_futuregen_ttfplot_dipping_strata.py`

### 3.3 LookupTableStratigraphy Component

The Lookup Table Stratigraphy component can be used to read stratigraphy data from a `.csv` or `.hdf5` file and then utilize this stratigraphy data in a simulation. Unit thicknesses are specified for different coordinates in the file, and spatial interpolation is used to produce unit thicknesses at different locations within the domain. This approach is meant to allow greater flexibility, but it also requires the user to be careful when setting up the data file used. The output from the Lookup Table Stratigraphy component can be checked with the `Stratigraphy` and `StratigraphicColumn` plot types in the control file interface.

In the NRAP-Open-IAM control file interface, the type name for the Lookup Table Stratigraphy component is `LookupTableStratigraphy`.

The arrangement of geologic units for this component is the same as the that for the `Stratigraphy` component. The lowest unit considered is the reservoir. There are alternating shales and aquifers overlying the reservoir. The lowest shale, shale 1, is immediately above the reservoir. Aquifer 1 is above shale 1, and shale 2 is above aquifer 1. This pattern continues until shale *N*, which extends to the surface, where *N* is the number of shale units used. There are *N* - 1 aquifers, and the number of shale layers is specified with the `numberOfShaleLayers` parameter. The `numberOfShaleLayers` cannot vary across the domain. If one wants a unit to effectively pinch out in certain areas, however, its thickness values for that area in the file can be decreased to to the minimum thickness of 1 *m*.

In the NRAP-Open-IAM control file interface, the `FileDirectory` and `FileName` keywords must be specified. The `FileDirectory` keyword indicates the directory where the data files for the lookup tables are located. The `FileName` keyword is the name of the `.csv` or `.hdf5` file that stores the unit thicknesses. The name should be given with the extension included (e.g., `stratigraphy.csv`). The first two columns in the file should be labeled as *x* and *y*, with these two columns containing easting and northing distances, respectively. The other columns in the file should be labeled with names that correspond with shale units (`shale#Thickness`), aquifer units (`aquifer#Thickness`), or the reservoir (`reservoirThickness`). Here, the # character shown for shales and aquifers should be replaced with the index (e.g., `shale5Thickness` for shale 5). The unit thicknesses in each row of the file represent the stratigraphy at the corresponding *x* and *y* values. Unit thicknesses can vary over space, but they cannot change with time in a simulation.

The Lookup Table Stratigraphy component produces the output using spatial interpolation within the domain defined by the coordinates in the data file used. An error will occur if the component is asked to produce output for a location that lies outside of the domain covered by the file. If this error occurs, the user can expand the domain covered by the data file by adding more rows with unit thicknesses at  $x$  and  $y$  values outside the current ranges.

Descriptions of the component's parameters are provided below.

- **numberOfShaleLayers** [-] (3 to 30) - number of shale layers in the system (default: 3). The shale units must be separated by an aquifer.
- **datumPressure** [ $Pa$ ] (80,000 to 300,000) - pressure at the top of the system (default: 101,325).

The observations from the Lookup Table Stratigraphy component are:

- **shale#Thickness** [ $m$ ] - thickness of shale unit #, where # is an index ranging from one to *numberOfShaleLayers*. If data for a particular shale layer are not included in the file used, that layer will be assigned a default thickness is 250  $m$ .
- **aquifer#Thickness** [ $m$ ] - thickness of aquifer unit #, where # is an index ranging from one to (*numberOfShaleLayers* - 1). If data for a particular aquifer layer are not included in the file used, that layer will be assigned a default thickness of 100  $m$ .
- **reservoirThickness** [ $m$ ] - thickness of the storage reservoir. If data for the reservoir are not included in the file used, the default reservoir thickness is 50  $m$ .
- **shale#Depth** [ $m$ ] - depth to the bottom of shale unit #, where # is an index ranging from one to *numberOfShaleLayers*.
- **aquifer#Depth** [ $m$ ] - depth to the bottom of aquifer unit #, where # is an index ranging from one to (*numberOfShaleLayers* - 1).
- **reservoirDepth** [ $m$ ] - depth to the bottom of the storage reservoir.
- **shale#MidDepth** [ $m$ ] - depth to the middle of shale unit #, where # is an index ranging from one to *numberOfShaleLayers*.
- **aquifer#MidDepth** [ $m$ ] - depth to the middle of aquifer unit #, where # is an index ranging from one to (*numberOfShaleLayers* - 1).
- **reservoirMidDepth** [ $m$ ] - depth to the middle of the storage reservoir.
- **shale#TopDepth** [ $m$ ] - depth to the top of shale unit #, where # is an index ranging from one to *numberOfShaleLayers*.
- **aquifer#TopDepth** [ $m$ ] - depth to the top of aquifer unit #, where # is an index ranging from one to (*numberOfShaleLayers* - 1).
- **reservoirTopDepth** [ $m$ ] - depth to the top of the storage reservoir. The same value can also be produced with the observation name **depth**, however. The use of the **depth** output name is included to conform with the Stratigraphy component.

If the component produces a unit thickness for a shale, aquifer, or reservoir that falls outside the range of 1  $m$  to 1600  $m$ , then that thickness will be set to the lower or upper limit (whichever is closer). For example, if an interpolated thickness is 0.1  $m$ , based on the values included in the data file, then that thickness observation will be set to 1  $m$  instead.

Currently, this component cannot be used to produce unit thicknesses that vary stochastically in a simulation. The unit thicknesses and depths produced by a `LookupTableStratigraphy` component can vary over space within a simulation, but the thicknesses and depths produced for one location cannot vary across different realizations of a simulation. If one wanted to assess how altering unit thicknesses would impact results in a simulation using Latin Hypercube Sampling (LHS), for example, one should run multiple LHS simulations with different files used for the `LookupTableStratigraphy` components (with the files containing different stratigraphy inputs).

For control file examples using the `LookupTableStratigraphy` component, see *ControlFile\_ex32b.yaml* to *ControlFile\_ex32c.yaml*, *ControlFile\_ex38a.yaml* to *ControlFile\_ex38c.yaml*, *ControlFile\_ex39c.yaml*, *ControlFile\_ex55b.yaml*, and *ControlFile\_ex55d.yaml*. For script example examples, see *iam\_sys\_lutstrata.py*, *iam\_sys\_lutstrata\_gridded.py*, *iam\_sys\_lutstrata\_reservoir\_openwell.py*, *iam\_sys\_lutstrata\_reservoir\_mswell.py*, and *iam\_sys\_lutstrata\_reservoir\_cmwell.py*.

## 3.4 Analytical Reservoir Component

The Analytical Reservoir component model is a semi-analytical model for the reservoir. It is focused on flow across relatively large distances and does not take into account discrete features of the flow paths such as fractures, cracks, etc. The model is based on work of Nordbotten et al., [5]. Further reading can be found in [22], [3].

In the NRAP-Open-IAM control file, the type name for the analytical reservoir component is `AnalyticalReservoir`. The description of the component's parameters is provided below:

- **logResPerm** [ $\log_{10} m^2$ ] (-15.3 to -12) - logarithm of reservoir permeability (default: -13.69897)
- **reservoirPorosity** [-] (0.1 to 0.3) - porosity of reservoir (default: 0.15)
- **reservoirRadius** [ $m$ ] (500 to 100,000) - distance between injection well and outer reservoir boundary (default: 100,000). The reservoirRadius should not be smaller than the distance between the injection well and any leakage pathway components connected to the `AnalyticalReservoir` (e.g., wellbores or faults). If the radius is too small, the simulation will print a warning message.
- **brineDensity** [ $kg/m^3$ ] (965 to 1195) - density of brine phase (default: 1045)
- **CO2Density** [ $kg/m^3$ ] (450 to 976) - density of CO<sub>2</sub> phase (default: 479)
- **brineViscosity** [ $Pa \cdot s$ ] (2.3e-4 to 15.9e-4) - viscosity of brine phase (default: 2.535e-4)
- **CO2Viscosity** [ $Pa \cdot s$ ] (0.455e-6 to 1.043e-4) - viscosity of CO<sub>2</sub> phase (default: 3.95e-5)
- **brineResSaturation** [-] (0 to 0.25) - residual saturation of brine phase (default: 0)
- **brineCompressibility** [ $Pa^{-1}$ ] (3.63e-12 to 2.31e-11) - brine compressibility (default: 4.5e-12 = 3.1e-8 1/psi)
- **injRate** [ $m^3/s$ ] (0.0024 to 3.776) - CO<sub>2</sub> injection rate (default: 0.01)
- **numberOfShaleLayers** [-] (3 to 30) - number of shale layers in the system (default: 3); *linked to Stratigraphy*. The shale units must be separated by an aquifer.
- **shaleThickness** [ $m$ ] (1 to 1600) - thickness of shale layers (default: 250); *linked to Stratigraphy*. Thickness of shale layer 1, for example, can be defined by **shale1Thickness**; otherwise, shale layers for which the thickness is not defined will be assigned a default thickness.
- **aquiferThickness** [ $m$ ] (1 to 1600) - thickness of aquifers (default: 100); *linked to Stratigraphy*. Thickness of aquifer 1, for example, can be defined by **aquifer1Thickness**; otherwise, aquifers for which the thickness is not defined will be assigned a default thickness.
- **reservoirThickness** [ $m$ ] (15 to 500) - thickness of reservoir (default: 50); *linked to Stratigraphy*
- **datumPressure** [ $Pa$ ] (80,000 to 300,000) - pressure at the top of the system (default: 101,325); *linked to Stratigraphy*.

Possible observations from the Analytical Reservoir component are:

- **pressure** [ $Pa$ ] - pressure at top of the reservoir at the user defined location(s)
- **pressureAve** [ $Pa$ ] - pressure vertically averaged at the user defined location(s)
- **CO2saturation** [-] - CO<sub>2</sub> saturation vertically averaged at the user defined location(s)
- **mass\_CO2\_reservoir** [ $kg$ ] - (injected total) mass of the CO<sub>2</sub> in the reservoir.

For control file examples using the Analytical Reservoir component, see *ControlFile\_ex1a* to *ControlFile\_ex5*, *ControlFile\_ex8d*, *ControlFile\_ex31a* to *ControlFile\_ex31f*, *ControlFile\_ex39a* to *ControlFile\_ex39c*, and *ControlFile\_ex56a* to *ControlFile\_ex56g*. For script examples, see *iam\_sys\_reservoir\_mswell\_2aquifers.py*, *iam\_sys\_strata\_reservoir\_openwell\_genericaquifer.py*, *iam\_sys\_lutstrata\_reservoir\_mswell.py*, and *iam\_sys\_reservoir\_mswell\_generic\_lhs.py*.

## 3.5 Lookup Table Reservoir Component

The Lookup Table Reservoir component model is a reduced order model based on interpolation of data from a set of lookup tables. The lookup tables are based on the full-physics scale simulations. Each lookup table is determined by a particular set of M input model parameters which define a signature of the given set of lookup tables.

In the NRAP-Open-IAM control file, the type name for the Lookup Table Reservoir component is `LookupTableReservoir`. The component's parameters depend on the M input model parameters used to create lookup tables data. The minimum and maximum values of lookup table parameters determine boundaries of component parameters. Moreover, the component parameters values as a set can only be one of the combination of values that went into one of the lookup table linked to the component.

In the NRAP-Open-IAM control file a `FileDirectory` keyword must be specified. It indicates the directory where files with the simulation data for the lookup tables are located. A `TimeFile` keyword is a name of the .csv file that stores the time points (in years) at which the results in the tables are provided. If `TimeFile` is not specified then, by default, the name of the file with time data is assumed to be *time\_points.csv*. The time file must be located in the directory specified by `FileDirectory`.

A `ParametersFilename` keyword can also be specified. It defines the names and values of lookup table parameters that were used to create the given set of lookup tables. Additionally, it lists the names of the .csv files containing simulation data for each of the lookup tables in the set (e.g., results from different parameterizations of a reservoir simulation). By default, the file with parameters data is assumed to be named *parameters\_and\_filenames.csv*. The parameters file should be in a comma separated values format. The first M entries in the first row of the file are the names of the lookup table parameters which were varied for different realizations; the (M+1)st entry is a word *filename*. Each subsequent row of the parameters file contains the M values of the lookup table parameters followed by the name of file (lookup table) with the corresponding realization data. The provided filename must match with one of the files in the `FileDirectory`.

The user should make sure that the information provided in `ParametersFilename` file on parameters and simulation data files is accurate and complete. In general, a given parameter of the Lookup Table Reservoir component can have any possible name. At the same time the (possibly random) names specified by user in the `ParametersFilename` file should be the same names that the user would use in the control file for the description of the Lookup Table Reservoir component parameters. Due to the way the lookup tables are produced, each parameter of the reservoir component can only take on certain deterministic values. The possible values of a given parameter should be listed after the `values:` keyword followed by the list in square brackets ('['). The weights for each parameter can be specified with the `weights:` keyword followed by the list of weights for each value in square brackets. The weights should sum to 1, otherwise, they will be normalized. If no weights are provided all values are assumed to be equally likely to occur.

There exists an option to sample the data from the lookup tables without direct reference to any of the parameters used for creating the tables. User can use an auxiliary parameter `index` added to the Lookup Table Reservoir component to sample data from a particular lookup table file based on its index in the file *parameters\_and\_filenames.csv*. This option allows to use lookup tables in the scenarios where the total number of lookup table data files is less than the number of all possible combinations of the lookup table parameters.

Simulation data files (listed in `ParametersFilename` file) in a comma separated values format contain the reservoir simulation data, e.g., pressure and CO<sub>2</sub> saturation, varying over time. The data is used to build the Lookup Table Reservoir component output. Each realization file begins with a header line that is ignored by the code. Each subsequent row of the file represents a particular spatial location. The first and the second columns are the x- and y-coordinates of the location, respectively. The subsequent columns contain reservoir simulation data at the location defined in the first



two columns. The names of the columns should represent the data in them and have the form *base.obs.nm\_#* where *base.obs.nm* is the name of observation as used in the system model and *#* is an index of the time point at which the given observation is provided. The indexing of the reservoir simulation data should always start with 1 not with 0. For example, the pressure data at the first time point (even if this time point is 0) should always be indexed as *pressure\_1*. Further, if the column contains pressure data at the second time point, its name should be *pressure\_2*, and so on. If the column contains saturation data at the 12th time point, its name should be *CO2saturation\_12*. The order of the columns in the lookup table except the first two x and y columns is arbitrary. If some reservoir simulation data does not vary in time then the column name should indicate it: in any case its name should not contain underscore symbol *\_* followed by number (time index). For example, column with name *#temperature#* would indicate that the provided temperature data is constant in time.

The Lookup Table Reservoir component produces the output using interpolation in space and time within the spatio-temporal domain defined by the lookup tables simulation model setup. Observations from the Lookup Table Reservoir component are:

- **pressure** [*Pa*] - pressure at top of the reservoir at the wellbore location(s)
- **CO2saturation** [-] - CO<sub>2</sub> saturation at the top of the reservoir at the wellbore location(s).

Observations *pressure* and *CO2saturation* are mandatory for the Lookup Table Reservoir component which means that the linked lookup tables should contain the necessary data to produce them. In addition, the component can return any other type of observations provided in the lookup tables.

For control file examples using the Lookup Table Reservoir component, see *ControlFile\_ex6*, *ControlFile\_ex9c*, *ControlFile\_ex10*, *ControlFile\_ex14*, *ControlFile\_ex24*, *ControlFile\_ex32a* to *ControlFile\_ex32c*, *ControlFile\_ex37*, *ControlFile\_ex40*, and *ControlFile\_ex55a* to *ControlFile\_ex55d*. For script examples, see *iam\_sys\_lutreservoir\_5locs.py*, *iam\_sys\_lutreservoir\_mswell.py*, and *iam\_sys\_lutreservoir\_openwell\_futuregen\_aor\_3d.py*, and *iam\_sys\_lutreservoir\_mswell\_rand\_allocated\_wells\_lhs.py*.

## 3.6 Theis Reservoir Component

The Theis Reservoir component model is an analytical model for pressure in the reservoir. This component can simulate the use of multiple injection and/or extraction wells, with the pressure values produced reflecting the interaction of these wells. Each well can have injection rates that vary over time. Positive rates represent injection, while negative rates represent extraction. This component only produces the **pressure** output; it does not produce usable **CO2saturation** values (if requested, any **CO2saturation** values produced will be zero). One might therefore consider the wells used to be injecting or extracting brine, rather than CO<sub>2</sub>.

In the NRAP-Open-IAM control file, the type name for the Theis Reservoir component is *TheisReservoir*. Descriptions of the component's parameters are provided below:

- **initialPressure** [*Pa*] (8.0e+4 to 1.0e+7) - initial pressure at the top of the reservoir (default: 1.0e+6);
- **reservoirThickness** [*m*] (1 to 1600) - thickness of reservoir (default: 50); *linked to Stratigraphy*
- **logResPerm** [ $\log_{10} m^2$ ] (-14 to -9) - base 10 logarithm of reservoir permeability (default: -12)
- **reservoirPorosity** [-] (0.01 to 1) - porosity of reservoir (default: 0.3)
- **compressibility** [ $Pa^{-1}$ ] (1.0e-11 to 1.0e-6) - compressibility of porous media (default: 1.0e-10)
- **CO2Density** [ $kg/m^3$ ] (100 to 1500) - density of CO<sub>2</sub> phase (default: 479)
- **brineDensity** [ $kg/m^3$ ] (900 to 1500) - density of brine phase (default: 1000)
- **brineViscosity** [*Pa·s*] (1.0e-4 to 5.0e-3) - viscosity of brine phase (default: 2.535e-3).

Possible observations from the Theis Reservoir component are:

- **pressure** [*Pa*] - pressure at top of the reservoir at the user defined location(s)



- **CO2saturation** [-] - CO<sub>2</sub> saturation at the top of the reservoir at the user defined location(s)

The **CO2saturation** output is only included for compatibility purposes. Specifically, wellbore components require **CO2saturation** inputs, so an error could occur if the Theis Reservoir did not provide this output. All **CO2saturation** values produced by the Theis Reservoir component have fixed values of 0 at each time point for any simulation.

For control file examples using the Theis Reservoir component, see *ControlFile\_ex44a* to *ControlFile\_ex47*. For script examples, see *iam\_sys\_theis.py*, *iam\_sys\_theis\_4inj\_wells.py*, and *iam\_sys\_theis\_4inj\_grid.py*.

## 3.7 Multisegmented Wellbore Component

The Multisegmented Wellbore component estimates the leakage rates of brine and CO<sub>2</sub> along wells in the presence of overlying aquifers or thief zones. The model is based on work of Nordbotten et al., [23]. Further reading can be found in [22].

The model is focused on flow across relatively large distances and does not take into account discrete features of the flow paths such as fractures, cracks, etc. It assumes that leakage is occurring in the annulus between the outside of the casing and borehole. This area is assigned an “effective” permeability of the flow path. The permeability is applied over a length along the well that corresponds to the thickness of a shale formation. Each well is characterized by an effective permeability assigned to each segment of the well that crosses an individual formation. For example, if a well crosses N permeable formations, then it is characterized by N different permeability values. The model utilizes the one-dimensional multiphase version of Darcy’s law to represent flow along a leaky well.

In the NRAP-Open-IAM control file, the type name for the Multisegmented Wellbore component is **MultisegmentedWellbore**. The description of the component’s parameters are provided below. Names of the component parameters coincide with those used by model method of the **MultisegmentedWellbore** class.

- **logWellPerm** [ $\log_{10} m^2$ ] (-101 to -9) - logarithm of well permeability along shale layer (default: -13). Logarithm of well permeability along shale 3, for example, can be defined by **logWell3Perm**. Permeability of the well along the shale layers not defined by user will be assigned a default value.
- **logAquPerm** [ $\log_{10} m^2$ ] (-17 to -9) - logarithm of aquifer permeability (default: -12). Logarithm of aquifer 1 permeability, for example, can be defined by **logAqu1Perm**. Aquifer permeability not defined by user will be assigned a default value.
- **brineDensity** [ $kg/m^3$ ] (900 to 1500) - density of brine phase (default: 1000)
- **CO2Density** [ $kg/m^3$ ] (100 to 1000) - density of CO<sub>2</sub> phase (default: 479)
- **brineViscosity** [ $Pa \cdot s$ ] (1.0e-4 to 5.0e-3) - viscosity of brine phase (default: 2.535e-3)
- **CO2Viscosity** [ $Pa \cdot s$ ] (1.0e-6 to 1.0e-4) - viscosity of CO<sub>2</sub> phase (default: 3.95e-5)
- **aquBrineResSaturation** [-] (0 to 0.99) - residual saturation of brine phase in each aquifer (default: 0.0). For example, the residual brine saturation of aquifer2 can be defined by **aqu2BrineResSaturation**; otherwise, aquifer layers for which the residual brine saturation is not defined will be assigned a default value.
- **compressibility** [ $Pa^{-1}$ ] (1.0e-13 to 1.0e-8) - compressibility of brine and CO<sub>2</sub> phases (assumed to be the same for both phases) (default: 5.1e-11)
- **wellRadius** [ $m$ ] (0.01 to 0.5) - radius of leaking well (default: 0.05)
- **numberOfShaleLayers** [-] (3 to 30) - number of shale layers in the system (default: 3); *linked to Stratigraphy*. The shale units must be separated by an aquifer.
- **shaleThickness** [ $m$ ] (1 to 3000) - thickness of shale layers (default: 250); *linked to Stratigraphy*. Thickness of shale layer 1, for example, can be defined by **shale1Thickness**; otherwise, shale layers for which the thickness is not defined will be assigned a default thickness.

- **aquiferThickness** [*m*] (1 to 1600) - thickness of aquifers (default: 100); *linked to Stratigraphy*. Thickness of aquifer 1, for example, can be defined by **aquifer1Thickness**; otherwise, aquifers for which the thickness is not defined will be assigned a default thickness.
- **reservoirThickness** [*m*] (1 to 1600) - thickness of reservoir (default: 30); *linked to Stratigraphy*.
- **datumPressure** [*Pa*] (80,000 to 300,000) - pressure at the top of the system (default: 101,325); *linked to Stratigraphy*

The possible outputs from the Multisegmented Wellbore component are leakage rates of CO<sub>2</sub> and brine to each of the aquifers in the system and atmosphere. The names of the observations are of the form:

- **CO2\_aquifer1, CO2\_aquifer2, ..., CO2\_atm** [*kg/s*] - CO<sub>2</sub> leakage rates
- **brine\_aquifer1, brine\_aquifer2, ..., brine\_atm** [*kg/s*] - brine leakage rates
- **mass\_CO2\_aquifer1, mass\_CO2\_aquifer2, ..., mass\_CO2\_aquiferN** [*kg*] - mass of the CO<sub>2</sub> leaked into the aquifer.

For control file examples using the Multisegmented Wellbore, see *ControlFile\_ex2*, *ControlFile\_ex3*, *ControlFile\_ex8a*, *ControlFile\_ex24*, *ControlFile\_ex31f*, and *ControlFile\_ex39a*. For script examples, see *iam\_sys\_reservoir\_mswell\_2aquifers.py*, *iam\_sys\_lutstrata\_reservoir\_mswell.py*, and *iam\_sys\_reservoir\_mswell\_futuregen\_tfdplot\_dipping\_strata.py*.

## 3.8 Cemented Wellbore Component

The Cemented Wellbore component model is based on a multiphase well leakage model implemented in the NRAP-IAM-CS, [10]. The model is built off detailed full-physics Finite Element Heat and Mass (FEHM) simulations, [38]. The FEHM simulations are three-dimensional (3-D), multiphase solutions of heat and mass transfer of water and supercritical, liquid, and gas CO<sub>2</sub>. After the simulations are completed, the surrogate model is built based on the key input parameters and corresponding output parameters. The approximate (surrogate) model is represented by polynomials in terms of input parameters that then can be sampled to estimate leakage rate for wells. Early development work can be found in [15].

When using the control file interface with more than 3 shale layers, the ThiefZone keyword can be used to specify the thief zone aquifer and the LeakTo keyword can be specified to name the upper aquifer. These values will default to *aquifer1* and *aquifer2*, respectively, if are not provided by user. In the FEHM simulations used to create the surrogate model some of the stratigraphy layers were setup with a fixed thickness. In particular, shale above aquifer had thickness 11.2 m; aquifer and thief zone to which leakage was simulated were set to have thicknesses 19.2 m and 22.4 m, respectively; and reservoir had thickness of 51.2 m.

Component model input definitions:

- **logWellPerm** [ $\log_{10} m^2$ ] (-13.95 to -10.1) - logarithm of wellbore permeability (default: -13)
- **logThiefPerm** [ $\log_{10} m^2$ ] (-13.9991 to -12.00035) - logarithm of thief zone permeability (default: -12.2)
- **wellRadius** [*m*] (0.025 to 0.25) - radius of the wellbore (default: 0.05)
- **initPressure** [*Pa*] (1.0e+5 to 5.0e+7) - initial pressure at the base of the wellbore (default: 2.0e+7 *Pa*, or 20 *MPa*); *from linked component*
- **wellDepth** [*m*] (960 to 3196.8) - depth in meters from ground surface to top of reservoir (default: 1500); *linked to Stratigraphy*
- **depthRatio** [-] (0.30044 to 0.69985) - fraction of well depth to the center of the thief zone from the top of the reservoir (default: 0.5); *linked to Stratigraphy*.

Temporal inputs of the Cemented Wellbore component are not provided directly to the component model method but rather are calculated from the current and several past values of pressure and CO<sub>2</sub> saturation. The calculated temporal

inputs are then checked against the boundary assumptions of the underlying reduced order model. The Cemented Wellbore component model temporal inputs are:

- **deltaP** [ $Pa$ ] (105891.5 to 9326181.69) - difference between the current and initial pressure at the wellbore
- **pressurePrime** [ $Pa/s$ ] (-6675.03 to 2986.7) - first pressure derivative
- **pressureDPrime** [ $Pa/s^2$ ] (-111.265 to 10.806) - second pressure derivative
- **saturation** [-] (0.001 to 1.0) - CO<sub>2</sub> saturation at the wellbore
- **saturationPrime** [ $1/s$ ] (-4.290e-7 to 1.117e-3) - first CO<sub>2</sub> saturation derivative
- **saturationDPrime** [ $1/s^2$ ] (-6.923e-6 to 1.176e-6) - second CO<sub>2</sub> saturation derivative.

The possible outputs from the Cemented Wellbore component are leakage rates of CO<sub>2</sub> and brine to aquifer, thief zone and atmosphere. The names of the observations are of the form:

- **CO2\_aquifer1, CO2\_aquifer2, CO2\_atm** [ $kg/s$ ] - CO<sub>2</sub> leakage rates
- **brine\_aquifer1, brine\_aquifer2, brine\_atm** [ $kg/s$ ] - brine leakage rates
- **mass\_CO2\_aquifer1, mass\_CO2\_aquifer2** [ $kg$ ] - mass of CO<sub>2</sub> leaked into aquifers.

For control file examples using the Cemented Wellbore component, see *ControlFile\_ex1a* to *ControlFile\_ex1b*, *ControlFile\_ex5*, *\*ControlFile\_ex10\**, *ControlFile\_ex31e*, and *ControlFile\_ex56g*. For script examples, see *iam\_sys\_reservoir\_cmwell.py* and *iam\_sys\_lutstrata\_reservoir\_cmwell.py*.

## 3.9 Open Wellbore Component

The Open Wellbore model is a lookup table reduced order model based on the drift-flux approach, see [25]. This model treats the leakage of CO<sub>2</sub> up an open wellbore or up an open (i.e., uncemented) casing/tubing. The lookup table is populated using T2Well/ECO2N Ver. 1.0 [24], which treats the non-isothermal flow of CO<sub>2</sub> and brine up an open wellbore, allows for the phase transition of CO<sub>2</sub> from supercritical to gaseous, with Joule-Thompson cooling, and considers exsolution of CO<sub>2</sub> from the brine phase.

By default, when used within the control file interface the Open Wellbore is connected to the upper aquifer (e.g., aquifer 2 if there are 2 aquifers in the system). For user-defined scenarios the *LeakTo* keyword can be used to specify either the name of the aquifer (e.g., *aquifer1*) CO<sub>2</sub> leaks to or *atmosphere* for leakage to the atmosphere. The default value is *aquifer#* where # is an index of the uppermost aquifer.

The Open Wellbore component can be used to calculate leakage rates following positive change in reservoir pressure or only from changes in reservoir pressure above a critical pressure. To use the latter approach, the argument *crit\_pressure\_approach* should be set to *True* for the setup of the component. Here is an example of this setup in a script application:

```
OpenWellbore(name='ow', parent=sm, crit_pressure_approach=True)
```

To set *crit\_pressure\_approach* to *True* in the control file interface, the *Controls* section in the .yaml entry for the Open Wellbore should be included with an additional entry *critPressureApproach: True* indented beneath *Controls*. For an example setup, see control file example 31a.

If *crit\_pressure\_approach* is set to *True*, the default approach is for critical pressure ( $P_{crit}$ ,  $Pa$ ) to be calculated in the manner shown in section [Equations](#).

Instead of having critical pressure calculated by the component, one can enforce a particular critical pressure (the *critPressure* parameter) by setting the argument *enforce\_crit\_pressure* to *True* for the setup of the component. Here is an example in a script setup:

```
OpenWellbore(name='ow', parent=sm, crit_pressure_approach=True, enforce_crit_pressure=True)
```

To set `enforce_crit_pressure` to `True` in the control file interface, file interface, the `Controls` section in the `.yaml` entry for the Open Wellbore should be included with additional entry `enforceCritPressure: True` indented beneath `Controls`. If `enforce_crit_pressure` is not set to `True`, then the **critPressure** parameter will not be used.

When a critical pressure is used, flow through an Open Wellbore can still occur at pressures beneath the critical pressure if a CO<sub>2</sub> plume is present in the reservoir at the base of the well (i.e., buoyancy effects from the CO<sub>2</sub>).

Component model input definitions:

- **logReservoirTransmissivity** [ $\log_{10} m^3$ ] (-11.27 to -8.40) - reservoir transmissivity (default: -9.83)
- **logAquiferTransmissivity** [ $\log_{10} m^3$ ] (-11.27 to -8.40) - reservoir transmissivity (default: -9.83)
- **brineSalinity** [-] (0 to 0.2) - brine salinity (mass fraction) (default: 0.1)
- **brineDensity** [ $kg/m^3$ ] (900 to 1200) - brine density (default: 1045)
- **wellRadius** [ $m$ ] (0.025 to 0.25) - radius of the wellbore (default: 0.05)
- **wellTop** [ $m$ ] (0 to 500) - depth of well top (default: 500); *linked to Stratigraphy*. Note that this parameter represents how far leakage can extend from the reservoir along the open wellbore. For example, the cement used to plug the well may be of poor quality or damaged, but the damage may not allow leakage to reach the surface at 0 m. When using control files, `wellTop` can be set to the bottom depth of an aquifer by entering 'aquifer#Depth,' where # is the aquifer number. If the aquifer is too deep, however, the limits for this parameter would still be enforced. In control files, if the 'LeakTo' entry is provided as the name of the aquifer receiving leakage from the open wellbore (e.g., 'LeakTo: aquifer2') then the `wellTop` parameter will automatically be set to the bottom depth of the corresponding aquifer. If 'LeakTo' is set to 'atmosphere', `wellTop` will automatically be set to 0.
- **critPressure** [ $Pa$ ] (1.0e+5 to 9.0e+7) - pressure above which the model initiates leakage rates calculations. Default value of this parameter is not defined: either the user provides it through component setup or the value is calculated based on the value of **brineDensity** parameter.
- **reservoirDepth** [ $m$ ] (1000 to 4000) - depth of reservoir (well base) (default: 2000); *linked to Stratigraphy*. Note that if 'shale1Depth' is entered for this parameter in a control file, the parameter will be set to the bottom depth of shale 1 (which is also the top of the reservoir).

The possible outputs from the Open Wellbore component are leakage rates of CO<sub>2</sub> and brine to aquifer and atmosphere. The names of the observations are of the form:

- **CO2\_aquifer** and **CO2\_atm** [ $kg/s$ ] - CO<sub>2</sub> leakage rates
- **brine\_aquifer** and **brine\_atm** [ $kg/s$ ] - brine leakage rates.

For control file examples using the Open Wellbore component, see *ControlFile\_ex4a* to *ControlFile\_ex4b*, *ControlFile\_ex9a* to *ControlFile\_ex9c*, *ControlFile\_ex32a* to *ControlFile\_ex32c*, and *ControlFile\_ex55a* to *ControlFile\_ex55d*. For script examples, see *iam\_sys\_strata\_reservoir\_openwell\_genericaquifer.py*, *iam\_sys\_lutstrata\_reservoir\_openwell.py*, *iam\_sys\_reservoir\_openwell\_futuregen\_aor\_plot.py*, and *iam\_sys\_strata\_reservoir\_openwell\_genericaquifer\_5locs.py*.

## 3.10 Generalized Flow Rate Component

The Generalized Flow Rate component model is a model representing wide range of carbon dioxide (CO<sub>2</sub>) and brine leakage flow rates and created based on the results of multiple wellbore simulations. The generalized models facilitate the implementation of flow rates in an uncertainty quantification (UQ) framework since the relevant leakage rate and time parameters can be generated randomly. The basic shape for these models were constructed from the results of numerical wellbore simulations based on pressure and saturation profiles derived from the Kimberlina reservoir model [32] coupled with wellbore permeability to yield CO<sub>2</sub> and complimentary brine leakage functions. More details covering derivation and application of the model can be found in [21].

In the IAM control file, the type name for the Generalized Flow Rate component is `GeneralizedFlowRate`. The description of the possible component's parameters are provided below.

- **numberOfShaleLayers** [-] (3 to 30) - number of shale layers in the system (default: 3); *linked to Stratigraphy*. The shale units must be separated by an aquifer.
- **logPeakCO2Rate** [ $\log_{10}$  kg/s] (-inf to 5) - logarithm of the largest CO<sub>2</sub> flow rate (default: -5)
- **timePeakCO2Rate** [years] (0 to 1000) - time to reach the largest CO<sub>2</sub> flow rate from initial time (default: 5)
- **durationPeakCO2Rate** [years] (0 to 1000) - length of time period during which CO<sub>2</sub> flow rate was the largest (default: 10)
- **durationPeakZeroCO2Rate** [years] (0 to 1000) - length of time period during which CO<sub>2</sub> flow rate decreased from the largest rate to zero (default: 100)
- **logInitBrineRate** [ $\log_{10}$  kg/s] (-inf to 5) - logarithm of the initial brine flow rate (default: -10)
- **logFinalBrineRate** [ $\log_{10}$  kg/s] (-inf to 5) - logarithm of the final brine flow rate (default: -11.5). Ratio of initial brine rate over final brine rate is recommended to be between 0.2 and 0.3
- **durationInitBrineRate** [years] (0 to 1000) - length of initial brine flow rate time period (default: 2)
- **durationInitFinalBrineRate** [years] (0 to 1000) - length of time period during which brine flow rate decreased from initial to final rate (default: 10)
- **mitigationTime** [years] (0 to inf) - time at which the leakage was remediated (default: 10000)

The possible outputs from the Generalized Flow Rate component are leakage rates of CO<sub>2</sub> and brine to the aquifer specified by user. The names of the observations are of the form:

- **CO2\_aquifer#** [kg/s] - CO<sub>2</sub> leakage rates where # is an aquifer index
- **brine\_aquifer#** [kg/s] - brine leakage rates
- **mass\_CO2\_aquifer#** [kg] - mass of CO<sub>2</sub> leaked into the specified aquifer.

## 3.11 Hydrocarbon Leakage Component

The HydrocarbonLeakage component model is a reduced order model predicting liquid and gas leakage to the shallow aquifer between 100 to 410 years post-injection of CO<sub>2</sub> to depleted hydrocarbon field. The model output begins 100 years and extends to 410 years after injection stops. The component is based on a machine learning regression model fitted to the results of compositional multiphase flow transport simulations using a neural network. Total 192,000 data from ~1,000 numerical simulations were used to develop the model. The model predicts CO<sub>2</sub> and methane leakage in liquid and gas phases to a shallow aquifer. The model also predicts the total liquid (oil) and gas leakage to the shallow aquifer, where these total masses include all hydrocarbons (light, intermediate, and heavy) as well as CO<sub>2</sub>. The depth to the bottom of the shallow aquifer is assumed to be 60 ft (18.288 m) below the surface. The top of the shallow aquifer extends to the surface. Input parameters were sampled using Latin Hypercube Sampling across wide ranges.

Values of input parameters FCO2, FC1, FC4, and FC7Plus must sum to one. To allow some leniency (e.g., for issues related to rounding errors), the sum of these values must be within 0.0001 (0.01%) of one (0.9999 to 1.0001). This option was created for cases when the sum of the values is different from 1 by a small value (e.g., 1.0e-6). If the sum of the provided values is not sufficiently close to one, then a warning message is printed.

Since the temporal bounds for the HydrocarbonLeakage component are years 100 to 410 after injection stops, the component produces zero results for any times outside of this range. Any results outside the applicable time range should not be considered valid, however.

The description of the component's parameters is presented below:

- **reservoirDepth** [m] (914.4 to 2743.2) - depth to the top of the reservoir (default: 2000); *linked to Stratigraphy*

- **NTG** [-] (0.4 to 1.0) - net-to-gross ratio representing the fraction of reservoir contributing to the flow (default: 0.6)
- **logResPerm** [log10 m<sup>2</sup>] (-14.0057 to -13.0057) - logarithm of reservoir permeability (default: -13.5)
- **reservoirPressureMult** [-] (1.0 to 1.2) - factor used to represent a state of the reservoir pressurization post-injection (relative to the reservoir pressure calculated lithostatically) (default: 1.1)
- **logWellPerm** [log10 m<sup>2</sup>] (-17.0057 to -12.0057) - logarithm of wellbore permeability (default: -13.0)
- **avgWaterSaturation** [-] (0.471 to 0.792) - average water saturation in the reservoir (default: 0.5)
- **FCO2** [-] (0.432 to 0.693) - mole fraction of CO<sub>2</sub> in the reservoir post-injection (default: 0.55)
- **FC1** [-] (0.010 to 0.113) - mole fraction of methane in the reservoir post-injection (default: 0.05)
- **FC4** [-] (0.010 to 0.111) - mole fraction of intermediate hydrocarbons in the reservoir post-injection (default: 0.05)
- **FC7Plus** [-] (0.123 to 0.500) - mole fraction of heavy hydrocarbons in the reservoir post-injection (default: 0.35)

Component model outputs:

- **mass\_oil\_aquifer** [kg] - cumulative mass of oil leaked to the aquifer. This output includes all hydrocarbons (light, intermediate, and heavy hydrocarbons) and CO<sub>2</sub> in the liquid phase.
- **mass\_gas\_aquifer** [kg] - cumulative mass of gas leaked to the aquifer. This output includes all hydrocarbons (light, intermediate, and heavy hydrocarbons) and CO<sub>2</sub> in the gas phase.
- **mass\_methane\_gas\_aquifer** [kg] - cumulative mass of methane gas leaked to the aquifer
- **mass\_methane\_oil\_aquifer** [kg] - cumulative mass of the methane in oil phase leaked to the aquifer
- **mass\_CO2\_aquifer** [kg] - cumulative mass of liquid CO<sub>2</sub> leakage to aquifer
- **mass\_CO2\_gas\_aquifer** [kg] - cumulative mass of CO<sub>2</sub> gas leaked to the aquifer

For control file examples using the Hydrocarbon Leakage component, see *ControlFile\_ex52a* and *ControlFile\_ex52b*. For a script examples, see *iam\_sys\_hydrocarbon\_forward\_lhs.py*.

## 3.12 Seal Horizon Component

The Seal Horizon component model simulates the flow of CO<sub>2</sub> through a low permeability but fractured rock horizon (a “seal” formation) overlying the storage reservoir into which CO<sub>2</sub> is injected.

The rock horizon is represented by a number of “cells” arranged (conceptually) in an arbitrary shape grid. A two-phase, relative permeability approach is used with Darcy’s law for one-dimensional (1D) flow computations of CO<sub>2</sub> through the horizon in the vertical direction. The code also allows the simulation of time-dependent processes that can influence such flow.

The model is based on an earlier code, NSealR, created with GoldSim, and described in [19]. A stand-alone version of this code in Python is also available on the NETL EDX system, described as Seal ROM.

In the NRAP-Open-IAM control file, the type name for the component is SealHorizon. The following is a list of the component parameters, including the parameter names, units, accepted value range and the default value.

Reference parameters for each cell:

- **area** [m<sup>2</sup>] (1 to 2.6e+5) - area of the cell (default: 10000.0)
- **thickness** [m] (5 to 1000) - thickness of the cell (vertically) (default: 100)
- **baseDepth** [m] (800 to 9500) - depth to the base of seal (default: 1100)



- **permeability** [ $m^2$ ] (1.0e-22 to 1.0e-15) - cell equivalent initial permeability (default: 1.0e-18)
- **entryPressure** [ $Pa$ ] (100 to 2.0e+6) - entry threshold pressure that controls flow into rock (default: 5000)

Distribution parameters for thickness of the seal layer

- **aveThickness** [ $m$ ] (10 to 1000) - mean of the truncated normal distribution for thickness (default: 100)
- **stdDevThickness** [ $m$ ] (0 to 500) - standard deviation of the thickness distribution (default: 0)
- **minThickness** [ $m$ ] (5 to 1000) - minimum thickness; this value truncates the distribution and limits lower values (default: 75)
- **maxThickness** [ $m$ ] (10 to 1000) - maximum thickness; this value truncates the distribution and limits higher values (default: 125)

Note: The setup of the four distribution parameters above is not yet implemented in the control file interface or GUI of NRAP-Open-IAM and available only in the script interface.

Distribution parameters for permeability of the seal layer:

- **avePermeability** [ $m^2$ ] (1.0e-22 to 1.0e-16) - mean total vertical permeability of a lognormal distribution; equivalent value for fractured rock (default: 2.5e-16)
- **stdDevPermeability** [ $m^2$ ] (0 to 1.0e-17) - standard deviation of the total vertical permeability distribution (default: 0.0)
- **minPermeability** [ $m^2$ ] (1.0e-24 to 1.0e-17) - minimum total vertical permeability; this value truncates (censors) the vertical random distribution and limits lower values (default: 1.0e-18)
- **maxPermeability** [ $m^2$ ] (1.0e-21 to 1.0e-12) - maximum total vertical permeability; this value truncates (censors) the random distribution and limits higher values (default: 1.0e-15)

Note: The setup of the four distribution parameters above is not yet implemented in the control file interface or GUI of NRAP-Open-IAM and available only in the script interface.

- **heterFactor** [-] (1.0e-2 to 100) - increase factor of the permeability of cells selected for heterogeneity, if the heterogeneity approach is used (default: 0.5).

Reference parameters for all cells:

- **aveBaseDepth** [ $m$ ] (800 to 9500) - average depth to base of cell/reservoir top; interpolation depth (default: 1100)
- **aveBasePressure** [ $Pa$ ] (1.0e+6 to 6.0e+7) - average pressure at seal base during injection (default: 3.3e+7)
- **aveTemperature** [ $^{\circ}C$ ] (31 to 180) - average temperature of seal (default: 50)
- **salinity** [ $ppm$ ] (0 to 80000) - average salinity of seal (default: 1.5e+4)
- **staticDepth** [ $m$ ] (800 to 9500) - reference depth for computing static pressure at top of seal (default: 1000)
- **staticPressure** [ $Pa$ ] (1.0e+6 to 6.0e+7) - pressure at static reference depth for computing pressure at the cell top (default: 1.0e+7).

Fluid (conditions) parameters:

- **brineDensity** [ $kg/m^3$ ] (880 to 1080) - density of brine phase (default: 1004)
- **CO2Density** [ $kg/m^3$ ] (93 to 1050) - density of CO<sub>2</sub> phase (default: 597.8)
- **brineViscosity** [ $Pa \cdot s$ ] (1.5e-4 to 1.6e-3) - viscosity of brine phase (default: 5.634e-4)
- **CO2Viscosity** [ $Pa \cdot s$ ] (1.8e-5 to 1.4e-4) - viscosity of CO<sub>2</sub> phase (default: 4.452e-5)
- **CO2Solubility** [ $mol/kg$ ] (0 to 2) - solubility of CO<sub>2</sub> phase in brine (default: 0.035).

Two-phase model parameters for LET model:

- **wetting1** [-] (0.5 to 5) - wetting phase parameter  $L$  (default: 1)
- **wetting2** [-] (0.1 to 30) - wetting phase parameter  $E$  (default: 10)
- **wetting3** [-] (0 to 3) - wetting phase parameter  $T$  (default: 1.25)
- **nonwet1** [-] (0.5 to 5) - nonwetting phase parameter  $L$  (default: 1.05)
- **nonwet2** [-] (0.1 to 30) - nonwetting phase parameter  $E$  (default: 10)
- **nonwet3** [-] (0 to 3) - nonwetting phase parameter  $T$  (default: 1.25)
- **capillary1** [-] (0.01 to 5) - LET-model parameter  $L$  for capillary pressure (default: 0.2)
- **capillary2** [-] (0.01 to 30) - LET-model parameter  $E$  for capillary pressure (default: 2.8)
- **capillary3** [-] (0.01 to 3) - LET-model parameter  $T$  for capillary pressure (default: 0.43)
- **maxCapillary** [ $Pa$ ] (100 to 2.0e+8) - maximum capillary pressure for model (default: 1.0e+7)

Note: Parameters **wetting1**, **wetting2**, **wetting3**, **nonwet1**, **nonwet2**, **nonwet3**, **capillary1**, **capillary2**, **capillary3**, and **maxCapillary** are used only if parameter **relativeModel** is set to *LET*.

Parameters for BC model:

- **lambda** [-] (0 to 5) - lambda parameter in Brooks-Corey model (default: 2.5)

Note: Parameter **lambda** is used only if parameter/keyword argument **relativeModel** is set to *BC*.

Additional parameters for two-phase flow:

- **brineResSaturation** [-] (0.01 to 0.35) - residual brine saturation (default: 0.15)
- **CO2ResSaturation** [-] (0 to 0.35) - residual CO<sub>2</sub> saturation (default: 0)
- **relativeModel** [-] (LET or BC) - relative permeability model (default: LET)
- **permRatio** [-] (0 to 1.5) - ratio of nonwetting to wetting permeability (default: 0.6).

Time-model and rock type parameters:

- **influenceModel** [-] (integer: 0, 1, 2) - time-dependent permeability model (default: 0); deterministic parameter, i.e. cannot be set to be random. Model type used to compute the influence factor of the fluid flow on permeability for time-dependent response:
  - 0: No influence factor used.
  - 1: Use a time-dependent model based on exposure time to CO<sub>2</sub>. Parameters **rateEffect** and **totalEffect** control the initial time delay and the maximum extent of effect.
  - 2: Use a multivariant model that considers **reactivity**, **clayType**, **clayContent** and **carbonateContent** values together with **rateEffect** and **totalEffect** parameters to establish the magnitude of the influence factor.
- **influence** [-] (0 to 1) - initial permeability influence factor (default: 1)
- **rateEffect** [-] (0.01 to 0.65) - time variance parameter; this parameter controls the initial time delay in the permeability effect of the model (default: 0.1)
- **totalEffect** [-] (0.01 to 200) - time variance parameter; this parameter defines the total change in permeability of the model (as a factor) (default: 0.1)
- **reactivity** [-] (0 to 10) - reactivity of time model; factor controls the magnitude of permeability change (default: 8)
- **clayType** [-] (smectite, illite, or chlorite) - predominate clay mineral content in the seal horizon, defined as one of following categories:



- smectite (high swelling material)
- illite (moderate swelling material)
- chlorite (low swelling material) (default: smectite)
- **carbonateContent** [%] (0 to 100) - carbonate content in seal layer rock (default: 8)
- **clayContent** [%] (0 to 100) - clay mineral content in seal layer rock (default: 60).

Note: Parameters **rateEffect** and **totalEffect** are used only when parameter **influenceModel** is set to 1 or 2. These parameters control the initial time delay and the maximum extent of effect.

Note: Parameters **reactivity**, **clayType**, **carbonateContent**, and **clayContent** are used only when parameter **influenceModel** is set to 2.

The possible outputs from the Seal Horizon component are leakage rates of CO<sub>2</sub> and brine to aquifer through seal layer. The names of the observations are of the form:

- **CO2\_aquifer, brine\_aquifer** [*kg/s*] - CO<sub>2</sub> and brine leakage rates to aquifer through seal layer (individual cells) into overlying aquifer
- **mass\_CO2\_aquifer, mass\_brine\_aquifer** [*kg*] - mass of the CO<sub>2</sub> and brine leaked through seal layer (individual cells) into overlying aquifer
- **CO2\_aquifer\_total, brine\_aquifer\_total** [*kg/s*] - cumulative (for all cells) CO<sub>2</sub> and brine leakage rates to aquifer through seal layer into overlying aquifer
- **mass\_CO2\_aquifer\_total, mass\_brine\_aquifer\_total** [*kg*] - cumulative (for all cells) mass of the CO<sub>2</sub> and brine leaked through seal layer into overlying aquifer.

For control file examples using the Seal Horizon component, see *ControlFile\_ex19* and *ControlFile\_ex23*. For script examples, see *iam\_sys\_lutreservoir\_seal\_horizon.py* and *iam\_sys\_lutreservoir\_seal\_horizon\_samplers.py*.

### 3.13 Fault Flow Component

The Fault Flow component model simulates the flow of carbon dioxide along a low permeability fault from an injection horizon (into which carbon dioxide is injected) up to a freshwater aquifer. The theoretical base is predicated on one-dimension (1D), steady-state, two-phase flow of CO<sub>2</sub> through a saturated discontinuity (parallel plates) under CO<sub>2</sub> supercritical conditions. The flow in the current implementation uses the near-surface CO<sub>2</sub> supercritical point to be the upper point of flow. The surrounding rock matrix is considered relatively impermeable.

In the NRAP-Open-IAM control file, the type name for the Fault Flow component is **FaultFlow**. The description of the component's parameters is provided below:

Fault core setup parameters:

- **strike** [°] (0 to 360) - direction of fault: trend of fault strike taken clockwise from north (default: 30)
- **dip** [°] (10 to 90) - inclination of fault plane from strike, using right-hand rule from strike (default: 70)
- **length** [*m*] (0 to 10,000) - length of fault trace at surface from start point (default: 100)
- **xStart** [*m*] (-5.0e+07 to 5.0e+07) - x-coordinate of the fault start point taken as the left point on fault trace (default: 500)
- **yStart** [*m*] (-5.0e+07 to 5.0e+07) - y-coordinate of the fault start point taken as the left point on fault trace (default: 500)
- **nSegments** [-] (1 to 100) - number of separate fault divisions of the fault (default: 4)
- **faultProbability** [%] (0 to 100) - probability of fault existence (default: 100)

Fault aperture setup parameters:

- **aperture** [*m*] (0 to 0.05) - effective aperture of fault (default: 2.5e-6)
- **SGR** [-] (0 to 100) - shale gouge ratio for fault (default: 0)
- **stateVariable** [-] (0 to 1) - correction factor for near-surface flow (default: 1)

For variability of fault properties and orientation setup one can use the following eight distribution parameters.

Note: The setup of the eight distribution parameters below is not yet implemented in the control file interface or GUI of NRAP-Open-IAM and available only in the script interface.

Strike distribution parameters:

- **aveStrike** [°] (0 to 360) - average direction of fault: average trend of fault strike taken clockwise from north (default: 90); also default value for no variation in strike
- **spreadStrike** [°] (0 to 180) - spread in strike orientation (range of 2-sigma around average) (default: 0)

Dip distribution parameters:

- **aveDip** [°] (10 to 90) - average inclination of fault plane from strike, using right-hand rule from strike (default: 90)
- **stdDevDip** [°] (0 to 90) - standard deviation of angle of dip (default: 0)

Aperture distribution parameters:

- **aveAperture** [*m*] (0 to 1.01e-1) - average effective aperture of fault (default: 1.0e-2)
- **stdDevAperture** [*m*] (0 to 2.0e-2) - standard deviation of effective aperture (default: 0.0)
- **minAperture** [*m*] (0 to 1.0e-3) - minimum aperture (default: 1.0e-7)
- **maxAperture** [*m*] (0 to 5.0e-2) - maximum aperture (default: 2.0e-2)

Field parameters:

- **aquiferDepth** [*m*] (200 to 2,000) - depth to base of deepest aquifer along/above fault (default: 240)
- **aquiferTemperature** [°C] (15 to 180) - temperature of brine of deepest aquifer at base (default: 22)
- **aquiferPressure** [*Pa*] (1.0e+6 to 6.0e+8) - pressure at base of aquifer (default: 1.42E+07)
- **injectDepth** [*m*] (860 to 20,000) - reference depth positive below grade to top of injection horizon (default: 1880)
- **injectTemperature** [°C] (31 to 180) - average temperature of brine at injection depth in reservoir (default: 95)
- **fieldPressure** [*Pa*] (1.0e+5 to 6.0e+7) - initial pressure at injection depth before injection starts (default: 1.9140e+07)
- **injectPressure** [*Pa*] (7.0e+6 to 6.0e+8) - average pressure at base during injection period for interpolation of viscosity and density (default: 2.9290E+07)
- **finalPressure** [*Pa*] (1.0e+5 to 6.0e+7) - final average pressure at injection depth for interpolation of viscosity and density (default: 1.9140e+07)
- **injectX** [*m*] (-5.0e+07 to 5.0e+07) - x-coordinate of the location of injection well (default: 0)
- **injectY** [*m*] (-5.0e+07 to 5.0e+07) - y-coordinate of the location of injection well (default: 0)
- **injectEndTime** [years] (0 to 10000) - time when injection stops (default: 50)

Reservoir conditions parameters:

- **salinity** [ppm] (0 to 80000) - salinity of the brine (default: 0). The value is used to compute density and viscosity of the brine

- **CO2Density** [ $kg/m^3$ ] (93 to 1050) - average density of CO<sub>2</sub> phase for fault (default: 673.84). The value is used if interpolation is not conducted by code
- **CO2Viscosity** [ $Pa \cdot s$ ] (1.8e-05 to 1.4e-04) - viscosity of CO<sub>2</sub> phase for fault (default: 5.5173e-05). The value is used if interpolation is not conducted by code
- **brineDensity** [ $kg/m^3$ ] (880 to 1080) - density of brine phase for fault (default: 974.895). The value is used if interpolation is not conducted by code
- **brineViscosity** [ $Pa \cdot s$ ] (1.5e-04 to 1.6e-03) - viscosity of brine phase for fault (default: 3.0491e-04). The value is used if interpolation is not conducted by code
- **CO2Solubility** [ $mol/kg$ ] (0 to 2) - solubility of CO<sub>2</sub> phase in brine for fault (default: 0.035). The value is used if interpolation is not conducted by code

Aquifer conditions parameters:

- **aquiferCO2Density** [ $kg/m^3$ ] (93 to 1050) - density of CO<sub>2</sub> phase in the aquifer (default: 886.44)
- **aquiferCO2Viscosity** [ $Pa \cdot s$ ] (1.1e-05 to 1.4e-04) - viscosity of CO<sub>2</sub> phase in the aquifer (default: 8.8010e-05)
- **aquiferBrineDensity** [ $kg/m^3$ ] (880 to 1080) - density of brine phase in the aquifer (default: 1004.10)
- **aquiferBrineViscosity** [ $Pa \cdot s$ ] (1.5e-04 to 1.6e-03) - viscosity of brine phase in the aquifer (default: 3.0221e-04)

Relative flow parameters:

- **brineResSaturation** [-] (0.01 to 0.35) - residual wetting brine saturation used in two-phase model (default: 0.15)
- **CO2ResSaturation** [-] (0 to 0.35) - residual nonwetting CO<sub>2</sub> saturation used in two-phase model (default: 0.0)
- **relativeModel** [-] (LET or BC) - relative permeability model (default: LET)
- **permRatio** [-] (0 to 1.5) - ratio of maximum nonwetting permeability to the maximum wetting permeability (default: 0.6)
- **entryPressure** [ $Pa$ ] (100 to 2.0e+6) - entry/threshold/bubbling pressure that controls flow into rock (default: 5000)

Two-phase model parameters for LET model:

- **wetting1** [-] (0.5 to 5) - wetting phase parameter  $L$  (default: 1)
- **wetting2** [-] (1 to 30) - wetting phase parameter  $E$  (default: 10)
- **wetting3** [-] (0 to 3) - wetting phase parameter  $T$  (default: 1.25)
- **nonwet1** [-] (0.5 to 5) - nonwetting phase parameter  $L$  (default: 1.05)
- **nonwet2** [-] (1 to 30) - nonwetting phase parameter  $E$  (default: 10)
- **nonwet3** [-] (0 to 3) - nonwetting phase parameter  $T$  (default: 1.25)
- **capillary1** [-] (0.01 to 5) - LET-model parameter  $L$  for capillary pressure (default: 0.2)
- **capillary2** [-] (0.01 to 30) - LET-model parameter  $E$  for capillary pressure (default: 2.8)
- **capillary3** [-] (0.01 to 3) - LET-model parameter  $T$  for capillary pressure (default: 0.43)
- **maxCapillary** [ $Pa$ ] (100 to 2.0e+8) - maximum capillary pressure for model (default: 1.0e+7)

Note: Parameters **wetting1**, **wetting2**, **wetting3**, **nonwet1**, **nonwet2**, **nonwet3**, **capillary1**, **capillary2**, **capillary3**, and **maxCapillary** are used only if parameter **relativeModel** is set to *LET*.

BC model parameters:

- **lambda** [-] (0 to 5) - lambda term in Brooks-Corey model (default: 2.5)

Note: Parameter **lambda** is used only if parameter **relativeModel** is set to *BC*.

Stress parameters:

- **maxHorizontal** [*Pa*] (0 to  $5.0\text{e}+7$ ) - secondary maximum horizontal principal stress at top of injection horizon (default:  $3.0\text{e}+7$ )
- **minHorizontal** [*Pa*] (0 to  $5.0\text{e}+7$ ) - secondary minimum horizontal principal stress at top of injection interval (default:  $2.0\text{e}+7$ )
- **maxTrend** [ $^{\circ}$ ] (0 to 180) - strike of secondary maximum horizontal stress clockwise from north (default: 55)

The possible outputs from the Fault Flow component are leakage rates of CO<sub>2</sub> and brine to aquifer through fault. The names of the observations are of the form:

- **CO2\_aquifer, brine\_aquifer** [*kg/s*] - CO<sub>2</sub> and brine leakage rates to aquifer through fault (individual segments) into overlying aquifer
- **mass\_CO2\_aquifer, mass\_brine\_aquifer** [*kg*] - mass of the CO<sub>2</sub> and brine through fault (individual segments) to overlying aquifer
- **CO2\_aquifer\_total, brine\_aquifer\_total** [*kg/s*] - cumulative CO<sub>2</sub> and brine leakage rates to aquifer through fault into overlying aquifer
- **mass\_CO2\_aquifer\_total, mass\_brine\_aquifer\_total** [*kg*] - cumulative mass of the CO<sub>2</sub> and brine through fault (individual cells) to overlying aquifer.

Observations with names **CO2\_aquifer**, **brine\_aquifer**, **mass\_CO2\_aquifer** and **mass\_brine\_aquifer** are provided as arrays of values of length equal to the number of fault segments. To output observations corresponding to a particular fault segment (e.g., segment 1) one can add observations with names **CO2\_aquifer\_seg#** where # is an index of a segment of interest (e.g., **CO2\_aquifer\_seg1**) to the output of the Fault Flow component.

For control file examples using the Fault Flow component, see *ControlFile\_ex17*, *ControlFile\_ex18*, and *ControlFile\_ex48* to *ControlFile\_ex50*. For script examples, see the examples at the end of *fault\_flow\_component.py*.

## 3.14 Fault Leakage Component

The Fault Leakage Model component uses deep neural networks to estimate the flow of brine carbon dioxide along a fault. The dynamics of multiphase flow in the storage reservoir and shallow aquifer into which the fault leaks are both taken into consideration in the estimation of leakage rates. The CO<sub>2</sub> is treated as supercritical throughout the leakage process, including within the shallow aquifer. No phase change occurs. The fault is modeled as a continuous, homogeneous, isotropic porous medium with Darcy flow. This component assumes the following:

- The storage reservoir is 4000 m long (distance away from the fault), 2400 m wide, and 200 m thick.
- The shallow aquifer has similar dimensions to the storage reservoir, but is located on the opposite side of the fault.
- The fault has a thickness of 3 m and a width of 2400 m, contacting the entire width of both the storage reservoir and the shallow aquifer.
- The fault dip angle varies.
- There is a caprock on both sides of the fault with a thickness of 100 m perpendicular to the fault. The caprock has a permeability of  $1.0\text{e}-18$  (impermeable) and a porosity of 0.1.
- The storage reservoir has no-flow boundaries.
- The shallow aquifer has a constant pressure boundary at its side boundary. All other boundaries are no-flow.
- The domain is 1750 m in depth from the top of the shallow aquifer to the bottom of the storage reservoir.

- The pressure and temperature at the top of the shallow aquifer are 8.15 MPa and 50 °C, respectively.
- The CO<sub>2</sub> injection temperature is 32 °C.
- The thermal conductivity of rock is 3 W/(m\*K).
- The specific heat capacity of rock is 920 J/(kg\*K).
- Well is located about 178-190 m above the bottom of the storage reservoir.

The description of the component's parameters is provided below:

- **damage\_zone\_perm** [ $\log_{10} m^2$ ] (-15 to -12) - the permeability of the fault, including both the fault core and damage zone (default: -13.5)
- **damage\_zone\_por** [-] (0.001 to 0.1) - the porosity of the fault, including both the fault core and damage zone (default: 0.01)
- **shallow\_aquifer\_perm** [ $\log_{10} m^2$ ] (-14 to -12) - the permeability of the shallow aquifer (default: -13.0)
- **deep\_aquifer\_perm** [ $\log_{10} m^2$ ] (-14 to -12) - the permeability of the storage aquifer (e.g., reservoir) (default: -13.0)
- **shallow\_aquifer\_por** [-] (0.05 to 0.5) - the porosity of the shallow aquifer (default: 0.25)
- **deep\_aquifer\_por** [-] (0.05 to 0.35) - the porosity of the deep aquifer (default: 0.2)
- **well\_index** [-] (integer: 0, 1, 2) - a proxy for the horizontal distance of the well from the fault; value of 0 means that the well is about 200 m from the fault, value of 1 means that the well is about 400 m from the fault; value of 2 means that the well is about 600 m from the fault (default: 0)
- **well\_rate** [ $kg/s$ ] (0.5 to 25) - injection rate of the well for the aquifer (default: 15.8)
- **dip\_angle** [°] (integer: 40, 60, 80, 100, 120, 140) - dip angle of the fault, measured from the horizontal plane (default: 60)
- **injection\_time** [*years*] (10 to 50) - duration of injection (default: 30)
- **geothermal\_gradient** [°C/km] (8 to 44) - the geothermal gradient in the formation (default: 30)

The possible outputs from the Fault Leakage component are the leakage rates and cumulative leakage amounts of brine and CO<sub>2</sub> to the shallow aquifer through the fault. The names of the observations are:

- **brine\_aquifer**, **CO2\_aquifer** [ $kg/s$ ] - brine and CO<sub>2</sub> leakage rates to the shallow aquifer, respectively.
- **mass\_brine\_aquifer**, **mass\_CO2\_aquifer** [ $kg$ ] - cumulative brine and CO<sub>2</sub> leakage into the shallow aquifer, respectively.

Notes:

- Due to the use of the trapezoidal rule in integrating instantaneous leakage rates, the cumulative leakage values might not conserve mass for CO<sub>2</sub> depending on the time step size used.
- Leakage estimates are available only up to 100 years.
- After extensive sensitivity analysis the key parameters for the model have been found to be: damage zone permeability, deep aquifer permeability, injection rate of the well, injection duration, and dip angle. The time at which the leakage rate or amount is to be estimated (i.e., simulation time) is also a key parameter. Uncertainties in these variables are, therefore, most important.

## 3.15 Carbonate Aquifer Component

The Carbonate Aquifer component model is a reduced-order model that can be used to predict the impact that carbon dioxide (CO<sub>2</sub>) and brine leaks from a CO<sub>2</sub> storage reservoir might have on overlying aquifers. The model predicts the size of “impact plumes” according to nine water quality metrics, see [2], [6], [17].

Although the Carbonate Aquifer model was developed using site-specific data from the Edwards aquifer, the model accepts aquifer characteristics as variable inputs and, therefore, may have more broad applicability. Careful consideration should be given to the hydrogeochemical character of the aquifer before using this model at a new site. Guidelines and examples are presented in [16].

The size of “impact plumes” are calculated using two alternative definitions of “impact” which should be selected by user: 1) changes that cause an exceedance of a drinking water standard or maximum contaminant level (MCL); and 2) changes that are above and beyond “natural background variability” in the aquifer, [18].

Component model input definitions:

- **ithresh** [-] (1 or 2) - threshold, either 1: MCL or 2: No-impact (default: 2)
- **rmin** [*m*] (0 to 100) - maximum distance between leaks for them to be considered one leak (default: 15)
- **perm\_var** [ $\log_{10} m^4$ ] (0.017 to 1.89) - logarithm of permeability variance (default: 0.9535)
- **corr\_len** [*m*] (1 to 3.95) - correlation length (default: 2.475)
- **aniso** [-] (1.1 to 49.1) - anisotropy factor: ratio of horizontal to vertical permeability (default: 25.1)
- **mean\_perm** [ $\log_{10} m^2$ ] (-13.8 to -10.3) - logarithm of mean permeability (default: -12.05)
- **hyd\_grad** [-] (2.88e-4 to 1.89e-2) - horizontal hydraulic gradient (default: 9.59e-03)
- **calcite\_ssa** [ $m^2/g$ ] (0 to 1.0e-2) - calcite surface area (default: 5.5e-03)
- **organic\_carbon** [-] (0 to 1.0e-2) - organic carbon volume fraction (default: 5.5e-03)
- **benzene\_kd** [ $\log_{10} K_{oc}$ ] (1.49 to 1.73) - benzene distribution coefficient (default: 1.61)
- **benzene\_decay** [ $\log_{10}$  day] (0.15 to 2.84) - benzene decay constant (default: 0.595)
- **nap\_kd** [ $\log_{10} K_{oc}$ ] (2.78 to 3.18) - naphthalene distribution coefficient (default: 2.98)
- **nap\_decay** [ $\log_{10}$  day] (-0.85 to 2.04) - naphthalene decay constant (default: 0.595)
- **phenol\_kd** [ $\log_{10} K_{oc}$ ] (1.21 to 1.48) - phenol distribution coefficient (default: 1.35)
- **phenol\_decay** [ $\log_{10}$  day] (-1.22 to 2.06) - phenol decay constant (default: 0.42)
- **cl** [ $\log_{10}$  molality] (0.1 to 6.025) - brine salinity (default: 0.776)
- **logf** [-] (0 or 1) - type of transform of output plume volume; 0: linear, 1: log (default: 0)
- **aqu\_thick** [*m*] (100 to 500) - aquifer thickness (default: 300); *linked to Stratigraphy*

Component model dynamic inputs:

- **brine\_rate** [*kg/s*] (0 to 0.075) - brine rate
- **brine\_mass** [*kg*] (0 to 2.0e+8) - cumulative brine mass
- **co2\_rate** [*kg/s*] (0 to 0.5) - CO<sub>2</sub> rate
- **co2\_mass** [*kg*] (0 to 2.0e+9) - cumulative CO<sub>2</sub> mass.

Possible observations from the Carbonate Aquifer component are:

- **pH\_volume** [*m*<sup>3</sup>] - volume of aquifer below pH threshold
- **Flux** [*kg/s*] - CO<sub>2</sub> leakage rate to atmosphere

- **dx** [*m*] - length of impacted aquifer volume in x-direction
- **dy** [*m*] - width of impacted aquifer volume in y-direction
- **TDS\_volume** [*m*<sup>3</sup>] - volume of aquifer above TDS threshold in *mg/L*
- **As\_volume** [*m*<sup>3</sup>] - volume of aquifer above arsenic threshold in *μg/L*
- **Pb\_volume** [*m*<sup>3</sup>] - volume of aquifer above lead threshold in *μg/L*
- **Cd\_volume** [*m*<sup>3</sup>] - volume of aquifer above cadmium threshold in *μg/L*
- **Ba\_volume** [*m*<sup>3</sup>] - volume of aquifer above barium threshold in *μg/L*
- **Benzene\_volume** [*m*<sup>3</sup>] - volume of aquifer above benzene threshold
- **Naphthalene\_volume** [*m*<sup>3</sup>] - volume of aquifer above naphthalene threshold
- **Phenol\_volume** [*m*<sup>3</sup>] - volume of aquifer above phenol threshold.

For control file examples using the Carbonate Aquifer component, see *ControlFile\_ex3*, *ControlFile\_ex4a*, *ControlFile\_ex13*, and *ControlFile\_ex42*. For script examples, see *iam\_sys\_reservoir\_mswell\_2aquifers.py*, *iam\_sys\_reservoir\_mswell\_4aquifers\_timeseries.py*, and *iam\_sys\_reservoir\_mswell\_aquifer\_lhs.py*.

## 3.16 Deep Alluvium Aquifer Component

The Deep Alluvium Aquifer component model is a reduced order model which can be used to predict the changes in diluted groundwater chemistry if CO<sub>2</sub> and brine were to leak into a deep alluvium aquifer similar to the one located below the Kimberlina site, in the Southern San Joaquin Valley, California. The protocol allows uncertainty and variability in aquifer heterogeneity, fluid transport, and potential CO<sub>2</sub> and brine leakage rates from abandoned or damaged oil and gas wells to be collectively evaluated to assess potential changes in groundwater pH, total dissolved solids (TDS), and changes in the aquifer pressure resulting from leakage.

Although the Deep Alluvium Aquifer model was developed using site-specific data from the LLNL's Kimberlina Model (version 1.2), the model accepts aquifer characteristics as variable inputs and, therefore, may have broader applicability. Careful consideration should be given to the hydrogeochemical character of the aquifer before using this model at a new site.

Model was created using the *py-earth* Python package [28]. Simulation data used to build this model was created by Mansoor et al. [20]. In the NRAP-Open-IAM control file, the type name for the Deep Alluvium Aquifer component is *DeepAlluviumAquifer*.

Component model input definitions:

- **logK\_sand1** [ $\log_{10} m^2$ ] (-12.92 to -10.92) - permeability of layer 1 at depth between 10 and 546 *m* (default: -11.92)
- **logK\_sand2** [ $\log_{10} m^2$ ] (-12.72 to -10.72) - permeability of layer 2 at depth between 546 and 1225 *m* (default: -11.72)
- **logK\_sand3** [ $\log_{10} m^2$ ] (-12.7 to -10.7) - permeability of layer 3 at depth between 1225 and 1411 *m* (default: -11.70)
- **logK\_caprock** [ $\log_{10} m^2$ ] (-16.699 to -14.699) - permeability of caprock at depth between 0 and 10 *m* (default: -15.70)
- **correlationLengthX** [*m*] (200 to 2000) - correlation length in x-direction (default: 1098.99)
- **correlationLengthZ** [*m*] (10 to 150) - correlation length in z-direction (default: 79.81)
- **sandFraction** [-] (0.7 to 0.9) - sand volume fraction (default: 0.8)



- **groundwater\_gradient** [-] (0.001000 to 0.001667) - regional groundwater gradient ( $dh/dx$ =change in hydraulic head/distance) (default: 0.001333)
- **leak\_depth** [m] (424.36 to 1341.48) - depth of leakage interval (default: 885.51).

Component model dynamic inputs:

- **brine\_rate** [kg/s] (0 to 0.017) - brine rate (default: 0.0003)
- **brine\_mass** [kg] (238.14419 to 8689604.29) - cumulative brine mass (default: 84722.74=10\*\*4.928)
- **co2\_rate** [kg/s] (0 to 0.385) - CO<sub>2</sub> rate (default: 0.045)
- **co2\_mass** [kg] (1.002 to 1.621e+9) - cumulative CO<sub>2</sub> mass (default: 1.636e+7=10\*\*7.214).

Observations from the Deep Alluvium Aquifer component are:

- **TDS\_volume** [m<sup>3</sup>] - volume of plume above baseline TDS change in mg/L (change in TDS > 100 mg/L)
- **TDS\_dx** [m] - length of plume above baseline TDS change in mg/L (change in TDS > 100 mg/L)
- **TDS\_dy** [m] - width of plume above baseline TDS change in mg/L (change in TDS > 100 mg/L)
- **TDS\_dz** [m] - height of plume above baseline TDS change in mg/L (change in TDS > 100 mg/L)
- **Pressure\_volume** [m<sup>3</sup>] - volume of plume above baseline pressure change in Pa (change in pressure > 500 Pa)
- **Pressure\_dx** [m] - length of plume above baseline pressure change in Pa (change in pressure > 500 Pa)
- **Pressure\_dy** [m] - width of plume above baseline pressure change in Pa (change in pressure > 500 Pa)
- **Pressure\_dz** [m] - height of plume above baseline pressure change in Pa (change in pressure > 500 Pa)
- **pH\_volume** [m<sup>3</sup>] - volume of plume below pH threshold (pH < 6.75)
- **pH\_dx** [m] - length of plume below pH threshold (pH < 6.75)
- **pH\_dy** [m] - width of plume below pH threshold (pH < 6.75)
- **pH\_dz** [m] - height of plume below pH threshold (pH < 6.75).

For control file examples using the Deep Alluvium Aquifer, see *ControlFile\_ex10* and *ControlFile\_ex43*. For a script example, see *iam\_sys\_deepalluvium.py*.

## 3.17 FutureGen2 Aquifer Component

The FutureGen 2.0 Aquifer component model is a reduced order model that can be used to predict the impact that carbon dioxide (CO<sub>2</sub>) and brine leakage from the CO<sub>2</sub> storage reservoir at the FutureGen 2.0 site might have on overlying aquifers or monitoring units. The model predicts the size of “impact plumes” according to four metrics: pH, total dissolved solids (TDS), pressure and dissolved CO<sub>2</sub>.

The FutureGen 2.0 Aquifer model is a regression model fitted to the results of STOMP-CO2E-R multiphase flow and reactive transport simulations of CO<sub>2</sub> and brine leakage using the *py-earth* Python package ([28]). The *py-earth* package is a Python implementation of the Multivariate Adaptive Regression Splines algorithm ([8]), in the style of *scikit-learn* ([26]), a library of machine-learning methods.

The aquifer simulations used to train the FutureGen 2.0 Aquifer component model were based on modeling done for monitoring program design at the FutureGen 2.0 site ([31]), as well as porosity and permeability values from the ELAN logs and core samples taken from the characterization well. Isothermal simulations were performed for training the aquifer component model.

In the NRAP-Open-IAM control file, the type name for the FutureGen 2.0 Aquifer component is *FutureGen2Aquifer*. The description of the possible component's parameters are provided below:

- **aqu\_thick** [*m*] (30 to 90) - thickness of unit (default: 33.2); *linked to Stratigraphy*
- **depth** [*m*] (100 to 700) - depth to bottom of unit (default: 590.1); *linked to Stratigraphy*
- **por** [-] (0.02 to 0.2) - porosity of unit (default: 0.118)
- **log\_permh** [ $\log_{10} m^2$ ] (-14 to -11) - horizontal permeability (default: -13.39)
- **log\_aniso** [ $\log_{10}$ ] (0 to 3) - anisotropy ratio (default: 0.3)
- **rel\_vol\_frac\_calcite** [-] (0 to 1) - relative volume fraction of calcite in solid phase (default: 0.01).

Component model dynamic inputs:

- **brine\_rate** [*kg/s*] (0 to 31.622) - brine rate
- **brine\_mass** [*kg*] (0 to 6.985e+10) - cumulative brine mass
- **co2\_rate** [*kg/s*] (0 to 31.622) - CO<sub>2</sub> rate
- **co2\_mass** [*kg*] (0 to 6.985e+10) - cumulative CO<sub>2</sub> mass.

Observations from the FutureGen 2.0 Aquifer component are:

- **Pressure\_volume** [*m*<sup>3</sup>] - volume of plume where relative change in pressure > 0.065%
- **Pressure\_dx** [*m*] - length of plume where relative change in pressure > 0.065%
- **Pressure\_dy** [*m*] - width of plume where relative change in pressure > 0.065%
- **Pressure\_dz** [*m*] - height of plume where relative change in pressure > 0.065%
- **pH\_volume** [*m*<sup>3</sup>] - volume of plume where absolute change in pH > 0.2
- **pH\_dx** [*m*] - length of plume where absolute change in pH > 0.2
- **pH\_dy** [*m*] - width of plume where absolute change in pH > 0.2
- **pH\_dz** [*m*] - height of plume where absolute change in pH > 0.2
- **TDS\_volume** [*m*<sup>3</sup>] - volume of plume where relative change in TDS > 10%
- **TDS\_dx** [*m*] - length of plume where relative change in TDS > 10%
- **TDS\_dy** [*m*] - width of plume where relative change in TDS > 10%
- **TDS\_dz** [*m*] - height of plume where relative change in TDS > 10%
- **Dissolved\_CO2\_volume** [*m*<sup>3</sup>] - volume of plume where relative change in dissolved CO<sub>2</sub> concentration > 20%
- **Dissolved\_CO2\_dx** [*m*] - length of plume where relative change in dissolved CO<sub>2</sub> concentration > 20%
- **Dissolved\_CO2\_dy** [*m*] - width of plume where relative change in dissolved CO<sub>2</sub> concentration > 20%
- **Dissolved\_CO2\_dz** [*m*] - height of plume where relative change in dissolved CO<sub>2</sub> concentration > 20%

For control file examples using the FutureGen 2.0 Aquifer component, see *ControlFile\_ex8c*, *ControlFile\_ex14*, *ControlFile\_ex26*, *ControlFile\_ex31a*, *ControlFile\_ex32a*, and *ControlFile\_ex55a*. For script examples, see *iam\_sys\_reservoir\_openwell\_futuregen\_aor\_plot.py* and *iam\_sys\_lutreservoir\_mswell\_futuregen\_dream.py*.

## 3.18 FutureGen2 AZMI Component

The FutureGen 2.0 Above Zone Monitoring Interval (AZMI) component model is a reduced order model that can be used to predict the impact that carbon dioxide (CO<sub>2</sub>) and brine leakage from the CO<sub>2</sub> storage reservoir at the FutureGen 2.0 site might have on overlying aquifers or monitoring units. The model predicts the size of “impact plumes” according to five metrics: pH, total dissolved solids (TDS), pressure, dissolved CO<sub>2</sub> and temperature.

The FutureGen 2.0 AZMI model is a regression model fitted to the results of STOMP-CO2E-R multiphase flow and reactive transport simulations of CO<sub>2</sub> and brine leakage using the `py-earth` Python package ([28]). The `py-earth` package is a Python implementation of the Multivariate Adaptive Regression Splines algorithm ([8]), in the style of `scikit-learn` ([26]), a library of machine-learning methods.

The aquifer simulations used to train the FutureGen 2.0 AZMI component model were based on modeling done for monitoring program design at the FutureGen 2.0 site ([31]), as well as porosity and permeability values from the ELAN logs and core samples taken from the characterization well. Nonisothermal simulations were performed for training the AZMI component model.

In the NRAP-Open-IAM control file, the type name for the FutureGen 2.0 AZMI component is `FutureGen2AZMI`. The description of the possible component's parameters are provided below:

- **aqu\_thick** [*m*] (30 to 90) - thickness of unit (default: 33.2); *linked to Stratigraphy*
- **depth** [*m*] (700 to 1600) - depth to bottom of unit (default: 1043.9); *linked to Stratigraphy*
- **por** [-] (0.02 to 0.2) - porosity of unit (default: 0.118)
- **log\_permh** [ $\log_{10} m^2$ ] (-14 to -11) - horizontal permeability (default: -13.39)
- **log\_aniso** [ $\log_{10}$ ] (0 to 3) - anisotropy ratio (default: 0.3)
- **rel\_vol\_frac\_calcite** [-] (0 to 1) - relative volume fraction of calcite in solid phase (default: 0.01).

Component model dynamic inputs:

- **brine\_rate** [*kg/s*] (0 to 31.622) - brine rate
- **brine\_mass** [*kg*] (0 to 6.985e+10) - cumulative brine mass
- **co2\_rate** [*kg/s*] (0 to 31.622) - CO<sub>2</sub> rate
- **co2\_mass** [*kg*] (0 to 6.985e+10) - cumulative CO<sub>2</sub> mass.

Observations from the FutureGen 2.0 AZMI component are:

- **Pressure\_volume** [*m*<sup>3</sup>] - volume of plume where relative change in pressure > 0.065%
- **Pressure\_dx** [*m*] - length of plume where relative change in pressure > 0.065%
- **Pressure\_dy** [*m*] - width of plume where relative change in pressure > 0.065%
- **Pressure\_dz** [*m*] - height of plume where relative change in pressure > 0.065%
- **pH\_volume** [*m*<sup>3</sup>] - volume of plume where absolute change in pH > 0.2
- **pH\_dx** [*m*] - length of plume where absolute change in pH > 0.2
- **pH\_dy** [*m*] - width of plume where absolute change in pH > 0.2
- **pH\_dz** [*m*] - height of plume where absolute change in pH > 0.2
- **TDS\_volume** [*m*<sup>3</sup>] - volume of plume where relative change in TDS > 10%
- **TDS\_dx** [*m*] - length of plume where relative change in TDS > 10%
- **TDS\_dy** [*m*] - width of plume where relative change in TDS > 10%
- **TDS\_dz** [*m*] - height of plume where relative change in TDS > 10%

- **Dissolved\_CO2\_volume** [ $m^3$ ] - volume of plume where relative change in dissolved CO<sub>2</sub> concentration > 20%
- **Dissolved\_CO2\_dx** [ $m$ ] - length of plume where relative change in dissolved CO<sub>2</sub> concentration > 20%
- **Dissolved\_CO2\_dy** [ $m$ ] - width of plume where relative change in dissolved CO<sub>2</sub> concentration > 20%
- **Dissolved\_CO2\_dz** [ $m$ ] - height of plume where relative change in dissolved CO<sub>2</sub> concentration > 20%
- **Temperature\_volume** [ $m^3$ ] - volume of plume where relative change in temperature > 0.03%
- **Temperature\_dx** [ $m$ ] - length of plume where relative change in temperature > 0.03%
- **Temperature\_dy** [ $m$ ] - width of plume where relative change in temperature > 0.03%
- **Temperature\_dz** [ $m$ ] - height of plume where relative change in temperature > 0.03%

For control file examples using the FutureGen 2.0 AZMI component, see *ControlFile\_ex15*, *ControlFile\_ex39a*, and *ControlFile\_ex39b*. For script examples, see *iam\_sys\_reservoir\_openwell\_futuregen\_aor\_plot.py* and *iam\_sys\_lutreservoir\_mswell\_futuregen\_dream.py*.

## 3.19 Generic Aquifer Component

The Generic Aquifer component model is a surrogate model that can be used to predict the leakage of carbon dioxide (CO<sub>2</sub>) and brine from a CO<sub>2</sub> storage reservoir. The model predicts the mass fraction of CO<sub>2</sub> and salt on a 100x10 radial grid surrounding the leaky well and outputs these as gridded observations. The model also predicts the volume and dimensions of aquifer where pore water concentrations exceed specified threshold values of dissolved CO<sub>2</sub> and salt.

The Generic Aquifer model is a machine learning regression model fitted to the results of STOMP-CO2E-R multiphase flow and reactive transport simulations of CO<sub>2</sub> and brine leakage using Tensorflow 2.4. 50,000 nonisothermal multiphase flow simulations were used to train the Generic Aquifer component model. Input parameters were varied using Latin Hypercube Sampling across wide ranges.

In the NRAP-Open-IAM control file, the type name for the Generic Aquifer component is **GenericAquifer**. The gridded output of the Generic Aquifer component can be displayed using the **GriddedRadialMetric** plot type; this plot type can also save .csv files showing results across the domain.

Descriptions of the component's parameters are provided below:

- **aqu\_thick** [ $m$ ] (25 to 250) - thickness of unit (default: 33.2); *linked to Stratigraphy*
- **top\_depth** [ $m$ ] (100 to 4100) - depth to the top of the aquifer (default: 590.1); *linked to Stratigraphy*
- **por** [-] (0.02 to 0.25) - porosity of unit (default: 0.118)
- **log\_permh** [ $\log_{10} m^2$ ] (-14 to -10) - horizontal permeability (default: -13.39)
- **log\_aniso** [-] (0 to 3) - anisotropy ratio Kh/Kv (default: 0.3)
- **aquifer\_salinity** [-] (0.0 to 0.015) - salt mass fraction in the aquifer's water (default: 0.005). This value is a dimensionless mass fraction.
- **reservoir\_salinity** [-] (0.015 to 0.05) - salt mass fraction in the water leaked from the reservoir to the aquifer (default: 0.03). This value is a dimensionless mass fraction.
- **dissolved\_salt\_threshold** [-] (0.0 to 1.0) - threshold for salt mass fraction (default: 0.02). This threshold is a dimensionless mass fraction. Any regions of the aquifer with dissolved salt mass fractions exceeding this value will be included in the dissolved salt plumes calculated by this component.
- **dissolved\_co2\_threshold** [-] (0.0 to 1.0) - threshold for CO<sub>2</sub> mass fraction (default: 0.01). This threshold is a dimensionless mass fraction. Any regions of the aquifer with dissolved CO<sub>2</sub> concentrations exceeding this mass fraction threshold will be included in the dissolved CO<sub>2</sub> plumes calculated by this component.

The **aquifer\_salinity**, **reservoir\_salinity**, and **dissolved\_salt\_threshold** parameters of this component are dimensionless mass fractions representing the aquifer's initial salinity, the reservoir's salinity, and the threshold for dissolved salt plume formation in the aquifer, respectively. Total dissolved solid (TDS) concentrations in **[mg/L]** can be converted to mass fractions by converting to **[mg/m<sup>3</sup>]** (multiply by 1000 **[L/m<sup>3</sup>]**), then converting to **[g/m<sup>3</sup>]** (multiply by 0.001 **[mg/g]**), then converting to **kg/m<sup>3</sup>** (multiply by 0.001 **[g/kg]**), and finally dividing by a water density in **kg/m<sup>3</sup>**. When considering the **dissolved\_salt\_threshold** parameter, the bulk density of the mixture can change once brine leaks into the aquifer (likely increasing slightly, if the brine has a higher density). For the purposes of this conversion, however, you can use a water density of 1000 **kg/m<sup>3</sup>**. Using this approach, the default **dissolved\_salt\_threshold** value of 0.02 would correspond with a TDS threshold of 20,000 **[mg/L]**. Any regions of the aquifer with dissolved salt concentrations exceeding this threshold will be included in the dissolved salt plumes determined by this component. Additionally, with this approach the default **aquifer\_salinity** and **reservoir\_salinity** values of 0.005 and 0.03 would correspond with TDS concentration of 5000 **[mg/L]** and 30,000 **[mg/L]**, respectively. If the brine in the reservoir has a density higher than 1000 **kg/m<sup>3</sup>**, however, using that density instead of 1000 **kg/m<sup>3</sup>** would change the conversion. For example, using a brine density of 1100 **kg/m<sup>3</sup>** would cause the default **reservoir\_salinity** value of 0.03 to correspond with a TDS concentration of 33,000 **[mg/L]**.

Component model dynamic inputs:

- **brine\_mass** [*kg*] (0 to 6.985e+10) - cumulative brine mass
- **co2\_mass** [*kg*] (0 to 6.985e+10) - cumulative CO<sub>2</sub> mass.

Observations from the Generic Aquifer component are:

- **Dissolved\_salt\_volume** [*m<sup>3</sup>*] - volume of plume where relative change in salt mass fraction > dissolved\_salt\_threshold
- **Dissolved\_salt\_dr** [*m*] - radius of plume where relative change in salt mass fraction > dissolved\_salt\_threshold
- **Dissolved\_salt\_dz** [*m*] - height of plume where relative change in salt mass fraction > dissolved\_salt\_threshold
- **Dissolved\_CO2\_volume** [*m<sup>3</sup>*] - volume of plume where dissolved CO<sub>2</sub> mass fraction > dissolved\_co2\_threshold
- **Dissolved\_CO2\_dr** [*m*] - radius of plume where dissolved CO<sub>2</sub> mass fraction > dissolved\_co2\_threshold
- **Dissolved\_CO2\_dz** [*m*] - height of plume where dissolved CO<sub>2</sub> mass fraction > dissolved\_co2\_threshold

Gridded observations from the Generic Aquifer component are:

- **Dissolved\_CO2\_mass\_fraction** [-] - mass fraction of CO<sub>2</sub> in aquifer pore water on a 100x10 radial grid surrounding the leaky well
- **Dissolved\_salt\_mass\_fraction** [-] - mass fraction of salt in aquifer pore water on a 100x10 radial grid surrounding the leaky well
- **r\_coordinate** [*m*] - radial coordinates of the points in the 100x10 radial grid surrounding the leaky well. The 100 radii range from 1.62 *m* to about 77.5 **[km]**.
- **z\_coordinate** [*m*] - depth coordinates of the points in the 100x10 radial grid surrounding the leaky well. The 10 depths used are within the aquifer modeled by the GenericAquifer. The minimum depth is 5% of the aquifer's thickness above the base of the aquifer, while the maximum depth is 95% of the aquifer's thickness above the base of the aquifer. The increment used between depth values is 10% of the aquifer's thickness.

For control file examples using the Generic Aquifer component, see to *ControlFile\_ex24*, *ControlFile\_ex31b*, *ControlFile\_ex34*, *ControlFile\_ex41*, and *ControlFile\_ex54a* to *ControlFile\_ex54d*. For script examples, see *iam\_sys\_strata\_reservoir\_openwell\_genericaquifer.py*, *iam\_sys\_analytical\_mswell\_generic.py*, *iam\_sys\_analytical\_mswell\_generic\_lhs.py*, and *iam\_sys\_strata\_reservoir\_openwell\_genericaquifer\_5locs.py*.

## 3.20 Atmospheric Model Component

The Atmospheric model is meant to be used for performing scoping studies for CO<sub>2</sub> dispersion after leakage out of the ground. The employed method is an extension of the nomograph approach of Britter and McQuaid (1988) [4] developed for estimating dense gas plume length from a single or multiple leakage sources. The method is very fast and, therefore, amenable to general system-level geologic carbon sequestration (GCS) risk assessment. The method is conservative: it assumes the wind could be from any direction and handles multiple sources by a simple superposition approach [37]. A user's manual for the standalone model is available at [36].

The model is intended to be used for large CO<sub>2</sub> leakage rates (e.g., leakage from an open wellbore). It may not be suitable for very small leakage rate, as, in general, small release rates (e.g., less than 1.0e-5 kg/s) do not form a dense gas release due to ambient mixing. The inputs to the model are leakage rate(s) from leaky well(s), location(s) of leaky well(s), ambient conditions (wind speed), and receptor locations (home or business locations where people are present). The outputs from the model are flags at receptors indicating whether the CO<sub>2</sub> concentration at the location exceeds a pre-defined critical value, and the critical downwind distance from the sources.

Within the control file interface, receptor locations can be specified with the `receptors` keyword argument assigned a full path (including a name) to a csv file containing x- and y-coordinates of the receptors. Alternatively, the `x_receptor` and `y_receptor` keywords can be assigned a list of x- and y-coordinates of the receptors, respectively. In the NRAP-Open-IAM control file, the type name for the Atmospheric model component is `AtmosphericROM`.

Component model input definitions:

- **T\_amb** [°C] (5 to 40) - ambient temperature (default: 15)
- **P\_amb** [atmosphere] (0.7 to 1.08) - ambient pressure (default: 1)
- **V\_wind** [m/s] (1.e-10 to 20) - wind velocity (default: 5)
- **C0\_critical** [-] (0.002 to 0.1) - critical concentration (default: 0.01)
- **T\_source** [°C] (5 to 50) - released CO<sub>2</sub> temperature (default: 15)
- **x\_receptor** [m] - x-coordinate of receptor
- **y\_receptor** [m] - y-coordinate of receptor

Possible observations from the Atmospheric model component are:

- **outflag\_r###** [-] - count of critical distances receptor is within from original leak points; here, ### is a receptor number starting at 000
- **num\_sources** [-] - number of sources. The possible maximum is a number of leakage points; could be less as leakage sources can potentially coalesce.
- **x\_new\_s###** [m] - x-coordinate of leakage source; here ### is a source number starting at 000
- **y\_new\_s###** [m] - y-coordinate of leakage source
- **critical\_distance\_s###** [m] - critical downwind distance from each source.

For control file examples using the Atmospheric model, see *ControlFile\_ex9a* to *ControlFile\_ex9c*. For script examples, see *iam\_sys\_reservoir\_openwell\_atmosphere.py* and *iam\_sys\_reservoir\_openwell\_atmosphere\_5locs\_lhs.py*.

## 3.21 Plume Stability Component

The Plume Stability component model produces quantitative metrics of the area, change in area over time, mobility and spreading [11]. Plume mobility is the effective centroid velocity including the speed and direction of movement. Plume spreading is the effective longitudinal dispersion of the plume along its direction of maximum elongation. This direction is returned by the model as well. The mobility and spreading metrics are comprehensive in that they can effectively handle and account for complex continuous and discontinuous plumes and intra-plume migration. The metrics are calculated using 2D-scalar attribute field values as inputs. The model can read in field values formatted in the NRAP-Open-IAM dataset format. In order to process 3D scalar field data for the model, it is recommended to collapse the 3D data to 2D using a maximization approach as described in [11].

In the NRAP-Open-IAM control file, the type name for the Plume Stability component is `PlumeStability`. The data files providing input for the Plume Stability component need to satisfy the same requirements imposed on the data files used as input for the Lookup Table Reservoir component. In particular, for control file setup of the Plume Stability component the following three keywords have the same meaning:

- `FileDirectory` is a directory where files with the simulation data for the component are located, and which contains files described below;
- `TimeFile` keyword specifies a name of the .csv file that stores the time points (in years) at which the results in the data files are provided;
- `ParametersFilename` keyword contains a name of the .csv file containing the names and values of the parameters used to create the given set of data files; in addition, it lists the names of the .csv files in the folder `FileDirectory` containing simulation data for each of the data file in the set.

The additional keywords of the component's control file interface are:

- `Variables` is a list of observations names provided in the data files and for which some (or all) metrics will be calculated;
- `Thresholds` is a dictionary of pairs (observation name, value) providing threshold value above which the change in the observation value should be taken into account for the calculation of the plume stability metrics.

The only component model input parameter is `index` which indicates the index of the data file from the list in the last column of `ParametersFilename` to be used to produce plume stability metrics. The minimum and maximum value of the parameter is defined by the indices of data files provided in the list.

Possible observations from the Plume Stability component are

- `{obs}_areas` - area of the plume above the predefined threshold
- `{obs}_areas_dt` - change in the area of the plume above the predefined threshold
- `{obs}_mobility` - velocity of centroid of the plume above the predefined threshold
- `{obs}_mobility_angles` - angles/direction at which the centroid of the plume above the predefined threshold is changing
- `{obs}_spreading` - longitudinal dispersion of the plume above the predefined threshold along its direction of maximum elongation
- `{obs}_spreading_angles` - angles/direction at which the dispersion of the plume occurs.

Above, `{obs}` determines the name of observation for which the plume stability metrics are to be calculated and for which the data is provided in the data files used as input for the component, e.g. **pressure**, **CO2saturation**, etc.

For a control file example using the Plume Stability component, see *ControlFile\_ex16*. For script examples, see *iam\_plume\_stability\_FEHM.py*, *iam\_plume\_stability\_Kimb.py*, and *iam\_plume\_stability\_Kimb\_3d.py*.



## 3.22 Chemical Well Sealing Component

The Chemical Well Sealing component is based on the model described in [13]. It predicts whether a fracture at the cement caprock interface, upon exposure to CO<sub>2</sub>, would self-seal or not due to calcite precipitation. The model couples two-phase flow of supercritical CO<sub>2</sub> and brine through fractures, advective and diffusive transport along the fracture, diffusive transport within the cement, and chemical reactions between cement and carbonated brine. If the fracture is predicted to self-seal, the time required for sealing is also computed. The original model is described in [33], [35], and [14] and was calibrated using experimental data presented in [33], [35], and [34].

Component model input definitions:

- **fractureAperture** [*m*] (1.0e-5 to 2.0e-3) - aperture of the fractured leakage path (default: 2.0e-5). Any input aperture that is lower than 10 micron (lower bound) is set to 10 micron.
- **fractureLength** [*m*] (10.0 to 400.0) - length of the fractured leakage path (default: 20)
- **maxOverpressure** [*Pa*] (1.0e+6 to 1.5e+7) - maximum overpressure the base of the fracture is expected to experience (default: 5.0e+6).

The output from the Chemical Well Sealing component informs about the sealing ability of the fractured leakage pathway. In case the fracture seals, the component would also report sealing time.

- **seal\_flag** [-] - flag informing whether a fracture would seal (1) or not (0) due to calcite precipitation
- **seal\_time** [*s*] - predicted time for sealing a fracture by calcite precipitation. If fracture doesn't seal this variable is set to 0.0.

For a control file example using the Chemical Well Sealing component, see *ControlFile\_ex22*. For a script example, see *iam\_sys\_reservoir\_chem\_well\_seal.py*.



## COMPONENT COMPARISON

This section of the documentation provides the comparison tables for the wellbore component models available in NRAP-Open-IAM.

The following table provides comparison of the wellbore components model parameters.

Table 4.1: Comparison of wellbore components in NRAP-Open-IAM

Input variable	Cemented wellbore ROM	Multisegmented wellbore ROM	Open wellbore ROM
logWellPerm (well Permeability)	-13.95 to -10.1	-17 to -9	Open tubing or casing
logAquPerm or logThiefPerm (aquifer or thief zone permeability)	-13.995 to -12	-14 to -9	log of aquifer and reservoir transmissivity: -11.27 to -8.39
wellRadius (radius of the leaky well)	0.025 to 0.25	0.01 to 0.5	0.025 to 0.25
Brine and CO <sub>2</sub> properties	Included in the ROM simulations	Calculated as a function of depth. Pressure gradient: 9792 Pa/m. Temperature gradient: 25 C/km.	Mass fraction of salt: 0 to 0.2. Temperature gradient: 25 C/km. Temperature at surface: 15 degrees Celcius.
Number of shale layers	3	3 to 30	1
reservoirDepth and wellDepth (depth of the reservoir, well)	960 to 3200	No limitation	1000 to 4000
shaleThickness, aquiferThickness, reservoirThickness (thickness of shale, aquifer or reservoir)	reservoir: 51.2 m, thief zone: 22.4 m, aquifer: 29.2 m, upper caprock layer: 11.2 m	1 to 1600 m	Well Top: 0 to 500 m
brineResSaturation (residual brine saturation)	0.001	Used as tuning parameter to allow CO <sub>2</sub> to accumulate in intermediate aquifers	Not needed
deltaP (increase in pressure due to injection)	0.1 to 9.3 MPa	No limitations	0 to 20 MPa
Energy transport consideration	Nonisothermal	Isothermal	Nonisothermal
Governing wellbore equation employed	Two phase Darcy equation	Two phase Darcy equation	Drift flux
Type of ROM	numerical	analytical	numerical

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Table 4.1 – continued from previous page

Input variable	Cemented wellbore ROM	Multisegmented wellbore ROM	Open wellbore ROM
Reference	D. R. Harp, R. Pawar, J. W. Carey, C. W. Gable, Reduced order models of transient CO <sub>2</sub> and brine leakage along abandoned wellbores from geologic carbon sequestration reservoirs, Int. J. Greenhouse Gas Control, 45 (2016), pp. 150-162	S. Baek, D. H. Bacon, N. J. Huerta, NRAP-Open-IAM Multisegmented Wellbore Reduced-Order Model, 2021, PNNL-32364	L. H. Pan, S. W. Webb, C. M. Oldenburg, Analytical solution for two-phase flow in a wellbore using the drift-flux model. Adv. Water Resour. 34 (2011), pp. 1656-1665

Limitations of each model are summarized in the next table.

Table 4.2: Limitations of the wellbore components in NRAP-Open-IAM

Cemented wellbore ROM	Multisegmented wellbore ROM	Open wellbore ROM
Limited flexibility in specifying layer thicknesses	Deviations from full physics simulations due to analytical nature of the model	The ROM was derived from short-time simulations. Simulations were terminated after 1,000-time steps or 100 hours, whichever came first. It should not be used for long term predictions if leakage rate is high and would result in significant mass loss from reservoir.
Limited flexibility in specifying the number of thief zones (only 1 thief zone supported)	Large shale thicknesses, "Large wellbore segment length for shale (or cap rock) layers may lead to errors due to the nature of the analytical approach employed. Less than 100 m for each segment is recommended with the current model. Although a particular combination of model parameters can relax the degree of the error or enlarge the applicable length of the segment, the quality of the performance is not guaranteed with large segment lengths."	

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Table 4.2 – continued from previous page

Cemented wellbore ROM	Multisegmented wellbore ROM	Open wellbore ROM
Limitations on total reservoir pressure change. Reservoir pressure changes greater than 9.3 MPa are outside the bounds of the model. Pressure changes in saline aquifers are typically below this range. However, pressure changes can be larger than this range in other formations that are more limited in extent (e.g., depleted gas fields)	No leakage of CO <sub>2</sub> in intermediate aquifers, “The analytical model does not consider the lateral leakage into intermediate aquifers, resulting in zero accumulation of CO <sub>2</sub> in intermediate aquifers. All the CO <sub>2</sub> is transported to the topmost aquifer, yielding an overestimation of the amount of CO <sub>2</sub> leaking into the topmost aquifer. Residual brine saturation can be used as a proxy for lateral leakage into intermediate aquifers. Users can calibrate this parameter to yield a desirable CO <sub>2</sub> leakage rate/mass into the thief layers (intermediate aquifers). The parameter would have to be recalibrated if any of the other input parameters, like reservoir pressure, well permeability, etc., are changed.”	
Limitations of the first and second derivative of pressure and saturation changes		



## USE CASES

In the folder *examples/Control\_Files* there are a number of example control files distributed with the NRAP-Open-IAM tool. The description of the files is provided in the following table.

Table 5.1: Control file examples distributed with NRAP-Open-IAM

File name	Included components	Comments
ControlFile_ex1a.yaml	Analytical reservoir, cemented wellbore	Forward simulation. The saturation and pressure output produced by analytical reservoir model is used as input for cemented wellbore model. The example produces three TimeSeries type of plots. The first two plots is of the CO <sub>2</sub> leakage rates into the intermediate aquifer (aquifer 1) and the shallow aquifer (aquifer 2). The third plot is of the pressure in the reservoir at the wellbore locations.
ControlFile_ex1b.yaml	Analytical reservoir, cemented wellbore	Forward simulation. The saturation and pressure output produced by analytical reservoir model is used as input for cemented wellbore model. The example produces several TimeSeries type of plots. In addition, this example demonstrates the use of the UseMarkers, UseLines, and VaryLineStyle entries for TimeSeries plots.
ControlFile_ex2.yaml	Analytical reservoir, multi-segmented wellbore	Latin hypercube sampling, 30 realizations. A fixed seed can be specified for the sampling setup: symbol # before the seed has commented it out. This example shows a way to specify wellbore locations inside the wellbore component specification with the Locations keyword.
ControlFile_ex3.yaml	Analytical reservoir, multi-segmented wellbore, carbonate aquifer	Latin hypercube sampling, 30 realizations. Example illustrates use of known and random wellbore locations.
ControlFile_ex4a.yaml	Analytical reservoir, open wellbore, carbonate aquifer	Latin hypercube sampling, 30 realizations. Example demonstrates several plotting options: TimeSeries, TimeSeriesStats, and TimeSeriesAndStats. For analysis of the large number of realizations a user might find it more helpful to plot the time series statistics rather than the data for each realization using TimeSeriesStats plotting option. TimeSeriesAndStats plots the simulated values and the related statistics. The subplot keyword can be used to present some of the results in subplots.

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Table 5.1 – continued from previous page

File name	Included components	Comments
ControlFile_ex4b.yaml	Analytical reservoir, open wellbore	Latin hypercube sampling, 30 realizations. Example demonstrates several plotting options: <code>TimeSeries</code> , <code>TimeSeriesStats</code> , and <code>TimeSeriesAndStats</code> . For analysis of the large number of realizations a user might find it more helpful to plot the time series statistics rather than the data for each realization using <code>TimeSeriesStats</code> plotting option. <code>TimeSeriesAndStats</code> plots the simulated values and the related statistics. The <code>subplot</code> keyword can be used to present some of the results in subplots. Example also illustrates an option to setup open wellbore with an option to calculate the critical pressure.
ControlFile_ex5.yaml	Analytical reservoir, cemented wellbore	Example is used to demonstrate parameter study analysis.
ControlFile_ex6.yaml	Lookup table based reservoir	Latin hypercube sampling, 10 realizations. To run this example, the additional <i>Kimberlina</i> data set have to be downloaded and unzipped into the <i>source/components/reservoir/lookuptables/Kimb_54_sims</i> folder. The data files can be downloaded from the same EDX directory where the NRAP-Open-IAM was downloaded or from <a href="https://gitlab.com/NRAP/Kimberlina_data">https://gitlab.com/NRAP/Kimberlina_data</a> . The data set comes from simulation work done by Wainwright et. al. [32].
ControlFile_ex7a.yaml	Multisegmented wellbore	Latin hypercube sampling, 30 realizations. Example illustrates the use of dynamic input to drive a wellbore model without attaching a reservoir model. This functionality can be used to quickly evaluate behavior of a single component model with a fixed input or interactions between several component models without constructing a full systems model. In the example the dynamic input is provided as a list of pressure and CO <sub>2</sub> saturation data at the simulation time points.
ControlFile_ex7b.yaml	Multisegmented wellbore	Forward simulation. Example illustrates the use of dynamic input to drive a wellbore model without attaching a reservoir model. In the example the dynamic input is provided as a path to the file containing pressure and CO <sub>2</sub> saturation data at the simulation time points.
ControlFile_ex8a.yaml	Analytical reservoir, multi-segmented wellbore	Latin hypercube sampling, 300 realizations. Example demonstrates the use of the <code>Analysis</code> section to compute correlation and sensitivity coefficients and create visualizations of the analysis results. Each subsection of the <code>Analysis</code> section shows some of the available options for the user to control.
ControlFile_ex8b.yaml	Analytical reservoir, multi-segmented wellbore	Latin hypercube sampling, 300 realizations. Example demonstrates the use of the <code>Analysis</code> section to compute correlation and sensitivity coefficients and create visualizations of the analysis results. Each subsection of the <code>Analysis</code> section shows some of the available options for the user to control.

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Table 5.1 – continued from previous page

File name	Included components	Comments
ControlFile_ex8c.yaml	Analytical reservoir, open wellbore, FutureGen2 aquifer	Latin hypercube sampling, 100 realizations. Example demonstrates the use of the <b>Analysis</b> section to compute correlation and sensitivity coefficients and create visualizations of the analysis results. Each subsection of the <b>Analysis</b> section shows some of the available options for the user to control.
ControlFile_ex9a.yaml	Analytical reservoir, open wellbore, atmospheric ROM	Latin hypercube sampling, 30 realizations. The <b>AtmosphericROM</b> model simulates CO <sub>2</sub> dispersion in the atmosphere as a dense gas leak. Example illustrates two plotting options available only for the <b>AtmosphericROM</b> model. <b>AtmPlumeSingle</b> keyword can be used to plot the critical distance from leakage points for a single realization. For multiple Monte-Carlo simulations probability of being within a critical distance can be estimated and plotted using <b>AtmPlumeEnsemble</b> keyword. Both types of plots produce a map-view of the release area for each time step.
ControlFile_ex9b.yaml	Analytical reservoir, open wellbore, atmospheric ROM	Forward simulation. The <b>AtmosphericROM</b> model simulates CO <sub>2</sub> dispersion in the atmosphere as a dense gas leak. Example illustrates <b>AtmPlumeSingle</b> plotting option.
ControlFile_ex9c.yaml	Lookup table based reservoir, open wellbore, atmospheric ROM	Latin hypercube sampling, 30 realizations. The <b>AtmosphericROM</b> model simulates CO <sub>2</sub> dispersion in the atmosphere as a dense gas leak. Example illustrates two plotting options available only for the <b>AtmosphericROM</b> model. <b>AtmPlumeSingle</b> keyword can be used to plot the critical distance from leakage points for a single realization. For multiple Monte-Carlo simulations probability of being within a critical distance can be estimated and plotted using <b>AtmPlumeEnsemble</b> keyword. Both types of plots produce a map-view of the release area for each time step.
ControlFile_ex10.yaml	Lookup table based reservoir, cemented wellbore, deep alluvium aquifer	Latin hypercube sampling, 30 realizations. Example illustrates a setup connecting lookup table based reservoir, wellbore and aquifer impact components.
ControlFile_ex11.yaml	Analytical reservoir, multisegmented wellbore, carbonate aquifer	Forward simulation. The saturation/pressure output produced by analytical reservoir model is used to drive leakage from five multisegmented wellbore models separated into two groups according to their properties. Two carbonate aquifer components are linked to both groups of wellbores and estimate the impact from the leakage of CO <sub>2</sub> and brine into the aquifers 1 and 2.
ControlFile_ex12.yaml	Generalized flow rate	Forward simulation. Example simulates brine and CO <sub>2</sub> leakage rates to the deepest aquifer (aquifer 1) illustrating the use of the generalized flow rate component. The component does not require linking to other components.
ControlFile_ex13.yaml	Generalized flow rate, carbonate aquifer	Forward simulation. Example computes the leakage rates of brine and CO <sub>2</sub> to aquifers 1 and 2 utilizing the generalized flow rate component, and calculates the associated impact with the help of carbonate aquifer component.

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Table 5.1 – continued from previous page

File name	Included components	Comments
ControlFile_ex14.yaml	Lookup table based reservoir, multisegmented wellbore, FutureGen2 aquifer	Latin hypercube sampling, 50 realizations. Example illustrates a setup connecting lookup table based reservoir, multisegmented wellbore and aquifer impact components.
ControlFile_ex15.yaml	Lookup table based reservoir, multisegmented wellbore, FutureGen2 AZMI	Latin hypercube sampling, 100 realizations. Example illustrates a setup connecting lookup table based reservoir, multisegmented wellbore and aquifer impact components.
ControlFile_ex16.yaml	Plume stability	Example is used to demonstrate parameter study analysis for plume stability component.
ControlFile_ex17.yaml	Fault flow	Example demonstrates forward simulation for fault flow component. Dynamic inputs are provided as arrays.
ControlFile_ex18.yaml	Lookup table based reservoir, fault flow	Latin hypercube sampling, 10 realizations. To run this example, the additional <i>Kimberlina</i> data set have to be downloaded and unzipped into the <i>source/components/reservoir/lookuptables/Kimb_54_sims</i> folder. The data files can be downloaded from the same EDX directory where the NRAP-Open-IAM was downloaded or from <a href="https://gitlab.com/NRAP/Kimberlina_data">https://gitlab.com/NRAP/Kimberlina_data</a> . The data set comes from simulation work done by Wainwright et. al. [32].
ControlFile_ex19.yaml	Lookup table based reservoir, seal horizon	Latin hypercube sampling, 5 realizations. To run this example, the additional <i>Kimberlina</i> data set have to be downloaded and unzipped into the <i>source/components/reservoir/lookuptables/Kimb_54_sims</i> folder. The data files can be downloaded from the same EDX directory where the NRAP-Open-IAM was downloaded or from <a href="https://gitlab.com/NRAP/Kimberlina_data">https://gitlab.com/NRAP/Kimberlina_data</a> . The data set comes from simulation work done by Wainwright et. al. [32].
ControlFile_ex20.yaml	Analytical reservoir	Latin hypercube sampling, 10 realizations. Example illustrates use of analytical reservoir component.
ControlFile_ex21.yaml	Multisegmented wellbore, deep alluvium aquifer (ML)	Forward simulation. Example illustrates use of deep alluvium aquifer component (ML). Dynamic input provided through input files is used to drive wellbore component.
ControlFile_ex22.yaml	Chemical well sealing component	Forward simulation. Example illustrates setup of the chemical well sealing component.
ControlFile_ex23.yaml	Lookup table based reservoir, seal horizon	Forward simulation. To run this example, the additional <i>Kimberlina</i> data set have to be downloaded and unzipped into the <i>source/components/reservoir/lookuptables/Kimb_54_sims</i> folder. The data files can be downloaded from the same EDX directory where the NRAP-Open-IAM was downloaded or from <a href="https://gitlab.com/NRAP/Kimberlina_data">https://gitlab.com/NRAP/Kimberlina_data</a> . The data set comes from simulation work done by Wainwright et. al. [32].

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Table 5.1 – continued from previous page

File name	Included components	Comments
ControlFile_ex24.yaml	Lookup table based reservoir, multisegmented wellbore, generic aquifer	Forward simulation. Example illustrates a setup connecting lookup table based reservoir, multisegmented wellbore and aquifer impact components. This example requires the additional <i>FutureGen 2.0</i> data set that can be downloaded from the following source: <a href="https://edx.netl.doe.gov/dataset/phase-iii-nrap-open-iam/resource/71aeb591-1609-430b-8392-4d75ee84750c">https://edx.netl.doe.gov/dataset/phase-iii-nrap-open-iam/resource/71aeb591-1609-430b-8392-4d75ee84750c</a> . The data set has to be placed into the <i>source/components/reservoir/lookuptables/FutureGen2/1008_sims</i> folder.
ControlFile_ex25.yaml	Multisegmented wellbore, generic aquifer	Latin hypercube sampling, 10 realizations. Example illustrates the use of dynamic input to drive a wellbore component providing input for aquifer impact component.
ControlFile_ex26.yaml	Lookup table based reservoir, multisegmented wellbore, FutureGen2 aquifer	Forward simulation. Example illustrates a setup connecting lookup table based reservoir linked to 3d data to multisegmented wellbore and aquifer impact components. This example requires the additional <i>FutureGen 2.0</i> data set that can be downloaded from the following source: <a href="https://edx.netl.doe.gov/dataset/phase-iii-nrap-open-iam/resource/71aeb591-1609-430b-8392-4d75ee84750c">https://edx.netl.doe.gov/dataset/phase-iii-nrap-open-iam/resource/71aeb591-1609-430b-8392-4d75ee84750c</a> . The data set has to be placed into the <i>source/components/reservoir/lookuptables/FutureGen2/1008_sims</i> folder.
ControlFile_ex27.yaml	Lookup table based reservoir, multisegmented wellbore	Forward simulation. Example illustrates a setup connecting lookup table based reservoir and multisegmented wellbore components. Reservoir component is linked to lookup tables with additional temperature metric.
ControlFile_ex28.yaml	Lookup table based reservoir, multisegmented wellbore	Forward simulation. Example illustrates a setup connecting lookup table based reservoir and multisegmented wellbore components. Reservoir component is linked to lookup tables with additional data metric.
ControlFile_ex29.yaml	Lookup table based reservoir, multisegmented wellbore	Forward simulation. Example illustrates use of known and random wellbore locations in 3d domain. The pressure and CO2 saturation are obtained from a reservoir component linked to 3d lookup tables.
ControlFile_ex30.yaml	Lookup table based reservoir, multisegmented wellbore	Forward simulation. Example illustrates different options of setting locations both for lookup table reservoir and multisegmented wellbore components: through direct entry, through file, as equally spaced array points, and as a grid. This example requires the additional <i>FutureGen 2.0</i> data set that can be downloaded from the following source: <a href="https://edx.netl.doe.gov/dataset/phase-iii-nrap-open-iam/resource/71aeb591-1609-430b-8392-4d75ee84750c">https://edx.netl.doe.gov/dataset/phase-iii-nrap-open-iam/resource/71aeb591-1609-430b-8392-4d75ee84750c</a> . The data set has to be placed into the <i>source/components/reservoir/lookuptables/FutureGen2/1008_sims</i> folder.
ControlFile_ex31a.yaml	Analytical reservoir, open wellbore, FutureGen2 aquifer	Forward simulation. Example demonstrates the setup of the AoR plot as well as an option for user to control which and in what form the model outputs are saved.

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Table 5.1 – continued from previous page

File name	Included components	Comments
ControlFile_ex31b.yaml	Analytical reservoir, open wellbore, generic aquifer	Forward simulation. Example demonstrates the setup of the AoR plot as well as an option for user to control which and in what form the model outputs are saved.
ControlFile_ex31c.yaml	Analytical reservoir, open wellbore, FutureGen2 aquifer	Latin hypercube sampling, 70 realizations. Example demonstrates the setup of the AoR plot.
ControlFile_ex31d.yaml	Analytical reservoir, open wellbore, FutureGen2 aquifer	Latin hypercube sampling, 70 realizations. Example demonstrates the setup of the AoR plot with time list option.
ControlFile_ex31e.yaml	Analytical reservoir, cemented wellbore, FutureGen2 aquifer	Forward simulation. Example demonstrates the setup of the AoR plot.
ControlFile_ex31f.yaml	Analytical reservoir, multi-segmented wellbore, FutureGen2 aquifer	Forward simulation. Example demonstrates the setup of the AoR plot.
ControlFile_ex32a.yaml	Lookup table based reservoir, open wellbore, FutureGen2 aquifer	Forward simulation. Locations for wellbore components are provided through an input file. Example demonstrates the use of the AoR plot as well as an option for user to control which and in what form the model outputs are saved. This example requires the additional <i>FutureGen 2.0</i> data set that can be downloaded from the following source: <a href="https://edx.netl.doe.gov/dataset/phase-iii-nrap-open-iam/resource/71aeb591-1609-430b-8392-4d75ee84750c">https://edx.netl.doe.gov/dataset/phase-iii-nrap-open-iam/resource/71aeb591-1609-430b-8392-4d75ee84750c</a> . The data set has to be placed into the <i>source/components/reservoir/lookuptables/FutureGen2/1008_sims</i> folder.
ControlFile_ex32b.yaml	Lookup table based reservoir, open wellbore, FutureGen2 aquifer	Forward simulation. Locations for wellbore components are provided through an input file. Stratigraphy is setup as spatially varying. Example demonstrates the use of the AoR and stratigraphy plot types as well as an option for user to control which and in what form the model outputs are saved. This example requires the additional <i>FutureGen 2.0</i> data set that can be downloaded from the following source: <a href="https://edx.netl.doe.gov/dataset/phase-iii-nrap-open-iam/resource/71aeb591-1609-430b-8392-4d75ee84750c">https://edx.netl.doe.gov/dataset/phase-iii-nrap-open-iam/resource/71aeb591-1609-430b-8392-4d75ee84750c</a> . The data set has to be placed into the <i>source/components/reservoir/lookuptables/FutureGen2/1008_sims</i> folder.
ControlFile_ex32c.yaml	Lookup table based reservoir, open wellbore, FutureGen2 aquifer	Forward simulation. Locations for wellbore components are provided through an input file. Stratigraphy is setup as spatially varying. Example demonstrates the use of the AoR and stratigraphy plot types. This example requires the additional <i>FutureGen 2.0</i> data set that can be downloaded from the following source: <a href="https://edx.netl.doe.gov/dataset/phase-iii-nrap-open-iam/resource/71aeb591-1609-430b-8392-4d75ee84750c">https://edx.netl.doe.gov/dataset/phase-iii-nrap-open-iam/resource/71aeb591-1609-430b-8392-4d75ee84750c</a> . The data set has to be placed into the <i>source/components/reservoir/lookuptables/FutureGen2/1008_sims</i> folder.

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Table 5.1 – continued from previous page

File name	Included components	Comments
ControlFile_ex33a.yaml	Analytical reservoir, open wellbore	Forward simulation. Stratigraphy is setup as spatially varying. Parameters of wellbore component are defined as linked to the stratigraphy explicitly. Setup of stratigraphy and stratigraphic column plots is illustrated.
ControlFile_ex33b.yaml	Analytical reservoir, open wellbore	Forward simulation. Stratigraphy is setup as spatially varying. Parameters of wellbore component are defined as linked to the stratigraphy explicitly. Setup of stratigraphy and stratigraphic column plots is illustrated.
ControlFile_ex34.yaml	Analytical reservoir, open wellbore, generic aquifer	Forward simulation. Stratigraphy is setup as spatially varying. Parameters of wellbore component are defined as linked to the stratigraphy explicitly. Setup of default stratigraphy plot is illustrated.
ControlFile_ex35.yaml	Analytical reservoir, open wellbore	Forward simulation. Stratigraphy is setup as spatially varying. Parameters of wellbore component are defined as linked to the stratigraphy explicitly. Setup of stratigraphy plot is illustrated.
ControlFile_ex36.yaml	Analytical reservoir, open wellbore, generic aquifer	Forward simulation. Stratigraphy is setup as spatially varying. Parameters of wellbore component are defined as linked to the stratigraphy explicitly. Locations are setup with grid option. Setup of stratigraphy plot is illustrated.
ControlFile_ex37.yaml	Lookup table based reservoir, multisegmented wellbore	Forward simulation. Setup of stratigraphy plot is illustrated. This example requires the additional <i>FutureGen 2.0</i> data set that can be downloaded from the following source: <a href="https://edx.netl.doe.gov/dataset/phase-iii-nrap-open-iam/resource/71aeb591-1609-430b-8392-4d75ee84750c">https://edx.netl.doe.gov/dataset/phase-iii-nrap-open-iam/resource/71aeb591-1609-430b-8392-4d75ee84750c</a> . The data set has to be placed into the <i>source/components/reservoir/lookuptables/FutureGen2/1008_sims</i> folder.
ControlFile_ex38.yaml	Simple reservoir, open wellbore	Forward simulation. Stratigraphy is setup as spatially varying. Parameters of wellbore component are defined as linked to the stratigraphy explicitly. Setup of stratigraphy plot is illustrated.
ControlFile_ex39a.yaml	Analytical reservoir, multi-segmented wellbore, FutureGen2 AZMI	Latin hypercube sampling, 30 realizations. Example demonstrates the setup of the TTFD plot.
ControlFile_ex39b.yaml	Analytical reservoir, multi-segmented wellbore, FutureGen2 AZMI	Forward simulation. Example demonstrates the setup of the TTFD and stratigraphy plots.
ControlFile_ex40.yaml	Lookup table based reservoir, multisegmented wellbore, FutureGen2 AZMI	Latin hypercube sampling, 30 realizations. Example demonstrates the setup of the TTFD plot.
ControlFile_ex41.yaml	Analytical reservoir, open wellbore, generic aquifer	Forward simulation. Example demonstrates the setup of the TTFD plot.
ControlFile_ex42.yaml	Analytical reservoir, multisegmented wellbore, carbonate aquifer	Latin hypercube sampling, 30 realizations. Example demonstrates the setup of the TTFD plot.
ControlFile_ex43.yaml	Analytical reservoir, multi-segmented wellbore, deep alluvium aquifer	Latin hypercube sampling, 30 realizations. Example demonstrates the setup of the TTFD plot.

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File name	Included components	Comments
ControlFile_ex44a.yaml	Theis reservoir	Forward simulation. Example demonstrates the setup of simulation with Theis reservoir component. Injection rates and times are defined with the array of values.
ControlFile_ex44b.yaml	Theis reservoir	Latin hypercube sampling, 30 realizations. Example demonstrates the setup of simulation with Theis reservoir component. Injection rates and times are defined with the array of values.
ControlFile_ex45a.yaml	Theis reservoir	Forward simulation. Example demonstrates the setup of simulation with Theis reservoir component. Injection rates and times are defined with the array of values.
ControlFile_ex45b.yaml	Theis reservoir	Forward simulation. Example demonstrates the setup of simulation with Theis reservoir component. Injection rates and times are defined through input files.
ControlFile_ex45c.yaml	Theis reservoir	Forward simulation. Example demonstrates the setup of simulation with Theis reservoir component. Injection rates and times are defined through input files.
ControlFile_ex45d.yaml	Theis reservoir	Forward simulation. Example demonstrates the setup of simulation with Theis reservoir component. Injection rates and times are defined through the same input file.
ControlFile_ex46a.yaml	Theis reservoir	Forward simulation. Example demonstrates the setup of simulation with Theis reservoir component. The reservoir component is setup with multiple injection wells. Injection rates and times are defined with multidimensional arrays.
ControlFile_ex46b.yaml	Theis reservoir	Forward simulation. Example demonstrates the setup of simulation with Theis reservoir component. The reservoir component is setup with multiple injection wells. Injection rates and times are defined through input files.
ControlFile_ex47.yaml	Generic reservoir	Forward simulation. Example illustrates setup of generic reservoir component.
ControlFile_ex48.yaml	Fault flow	Forward simulation. Example illustrates setup of fault flow component. Dynamic inputs are provided as arrays.
ControlFile_ex49.yaml	Fault flow	Forward simulation. Example illustrates setup of fault flow component. Dynamic inputs are provided as arrays.
ControlFile_ex50.yaml	Analytical reservoir, fault flow	Latin hypercube sampling, 30 realizations. Example demonstrates the application of fault flow component. Input to the component is provided from the linked reservoir component.
ControlFile_ex51a.yaml	Analytical reservoir, cemented wellbore (wr)	Forward simulation. Example illustrates setup of system model with analytical reservoir component providing required input for cemented wellbore (wr) component. Example also illustrates use of known and random wellbore locations.
ControlFile_ex51b.yaml	Analytical reservoir, cemented wellbore (wr)	Latin hypercube sampling, 32 realizations. Example illustrates setup of system model with analytical reservoir component providing required input for cemented wellbore (wr) component. Example also illustrates use of only random wellbore locations.

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Table 5.1 – continued from previous page

File name	Included components	Comments
ControlFile_ex51c.yaml	Analytical reservoir, cemented wellbore (wr)	Forward simulation. Example illustrates setup of system model with analytical reservoir component providing required input for cemented wellbore (wr) component. Example illustrates setup of cemented wellbore components simulating leakage to different aquifer layers.
ControlFile_ex52a.yaml	Hydrocarbon leakage	Forward simulation. Example illustrates setup of system model with hydrocarbon leakage component.
ControlFile_ex52b.yaml	Hydrocarbon leakage	Latin hypercube sampling, 30 realizations. Example illustrates setup of system model with hydrocarbon leakage component.
ControlFile_ex53a.yaml	Fault leakage	Forward simulation. Example illustrates setup of system model with fault leakage component.
ControlFile_ex53b.yaml	Fault leakage	Latin hypercube sampling, 25 realizations. Example illustrates setup of system model with fault leakage component.
ControlFile_ex54a.yaml	Analytical reservoir, open wellbore, generic aquifer	Forward simulation. Example illustrates setup of gridded radial metric plot.
ControlFile_ex54b.yaml	Analytical reservoir, open wellbore, generic aquifer	Forward simulation. Example illustrates setup of gridded radial metric plot with default settings.
ControlFile_ex54c.yaml	Lookup table based reservoir, multisegmented wellbore, generic aquifer	Forward simulation. Example illustrates setup of gridded radial metric plot.
ControlFile_ex54d.yaml	Analytical reservoir, open wellbore, generic aquifer	Latin hypercube sampling, 30 realizations. Example illustrates setup of gridded radial metric plot.
ControlFile_ex55a.yaml	Lookup table based reservoir, open wellbore, FutureGen2 aquifer	Forward simulation. Example illustrates setup of AoR workflow section.
ControlFile_ex55b.yaml	Lookup table based reservoir, open wellbore, FutureGen2 aquifer	Forward simulation. Example illustrates setup of spatially variable stratigraphy and AoR workflow sections.
ControlFile_ex55c.yaml	Lookup table based reservoir, open wellbore, generic aquifer	Forward simulation. Example illustrates setup of AoR workflow section with default settings.
ControlFile_ex55d.yaml	Lookup table based reservoir, open wellbore, FutureGen2 aquifer	Forward simulation. Example illustrates setup of spatially variable stratigraphy and AoR workflow sections.
ControlFile_ex56a.yaml	Analytical reservoir, open wellbore, generic aquifer	Forward simulation. Example illustrates simplified setup of AoR workflow sections.
ControlFile_ex56b.yaml	Analytical reservoir, open wellbore, FutureGen2 aquifer	Forward simulation. Example illustrates setup of AoR analysis and workflow sections.
ControlFile_ex56c.yaml	Analytical reservoir, open wellbore, FutureGen2 aquifer	Forward simulation. Example illustrates setup of AoR workflow section.
ControlFile_ex56d.yaml	Analytical reservoir, open wellbore, FutureGen2 aquifer	Latin hypercube sampling, 30 realizations. Example illustrates setup of AoR workflow section.
ControlFile_ex56e.yaml	Analytical reservoir, multisegmented wellbore, FutureGen2 aquifer	Forward simulation. Example illustrates setup of AoR workflow section.

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Table 5.1 – continued from previous page

File name	Included components	Comments
ControlFile_ex56f.yaml	Analytical reservoir, multi-segmented wellbore, FutureGen2 aquifer	Forward simulation. Example illustrates setup of AoR workflow section.
ControlFile_ex56g.yaml	Analytical reservoir, cemented wellbore, FutureGen2 aquifer	Forward simulation. Example illustrates setup of AoR workflow section.
ControlFile_ex57a.yaml	Lookup table based reservoir, multisegmented wellbore, FutureGen2 AZMI	Latin hypercube sampling, 30 realizations. Example illustrates setup of TTFD workflow section.
ControlFile_ex57b.yaml	Lookup table based reservoir, multisegmented wellbore, FutureGen2 AZMI	Forward simulation. Example illustrates setup of TTFD workflow section.
ControlFile_ex57c.yaml	Analytical reservoir, open wellbore, Generic aquifer	Forward simulation. Example illustrates setup of TTFD workflow section.
ControlFile_ex58a.yaml	Lookup table based reservoir, open wellbore	Forward simulation. Example illustrates setup of LeakageAssessment workflow section.
ControlFile_ex58b.yaml	Analytical reservoir, multi-segmented wellbore	Latin hypercube sampling, 30 realizations. Example illustrates setup of LeakageAssessment workflow section.
ControlFile_ex58b.yaml	Analytical reservoir, open wellbore	Latin hypercube sampling, 30 realizations. Example illustrates setup of LeakageAssessment workflow section.

Beyond control files the described scenarios can be implemented with a help of a Python script. Example scripts can be found in the *examples/scripts* folder.

## JUPYTER NOTEBOOKS

NRAP-Open-IAM has multiple examples illustrating functionality and applications of the tool for different types of applicable interfaces. In the folder *examples/Jupyter\_notebooks* there are a number of Jupyter notebooks (<https://docs.jupyter.org/en/latest/>) that provide guidance on the possible use cases and workflows. Some of these materials are included as part of documentation (available online only).



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