

OLI Studio V12.x

think simulation

getting the chemistry right

Introduction to OLI Studio

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Version: OLI Studio V12.x

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Disclaimer

This manual was produced using the OLI Studio 12.0.1 build 1 (12.0.0.6). As time progresses, new data and refinements to existing data sets can result in values that you obtain being slightly different than what is presented in this manual. This is a natural progress and cannot be avoided. When large systematic changes to the software occur, this manual will be updated.

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About This Manual

This manual will provide an introductory guide for new OLI Studio Software users. OLI Studio is a software suite containing several modules including Stream Analyzer, think simulation | getting the chemistry right and Corrosion Analyzer. A client's license determines which modules are enabled within the OLI Studio Software.

This introductory manual consists of one chapter showcasing the OLI Studio User Interface, and four main chapters showcasing the three main modules of OLI Studio: Stream Analyzer, Corrosion Analyzer, EVS Analyzer, and think simulation | getting the chemistry right.

If you have not yet installed the software, please install it following the instructions given in the Installation and Security page on our Wiki page at: https://support.olisystems.com

OLI Studio Components

Stream Analyzer

Stream Analyzer is standalone software, and it is the main interface of the OLI Studio. Stream Analyzer is a comprehensive thermodynamic tool that calculates speciation, phase equilibria, enthalpies, heat capacities and densities in mixed-solvent, multicomponent systems. Capabilities and features of Stream Analyzer are:

Three different thermodynamic frameworks:
Mixed Solvent Electrolyte (MSE) model (Default)
Aqueous (AQ) model
Mixed Solvent Electrolyte and Soave-Redlich-Kwong (MSE-SRK) model

Thermophysical properties: Stream Analyzer has thermophysical models to predict surface tension, interfacial tension, viscosity, electrical conductivity, thermal conductivity, diffusivity, and osmotic pressure.

Molecular and ionic inflows: Stream Analyzer accepts molecular inflows, typical of a process stream, and ion inflows, typical of a sample water analysis.

Corrosion Analyzer

Corrosion Analyzer is a module within the OLI Studio. A separate license enables this module. Corrosion Analyzer is a first-principles corrosion prediction tool. It is used to predict the corrosion rates of general corrosion, propensity of alloys to undergo localized corrosion, depletion profiles of heat-treated alloys, and thermodynamic stability of metals and alloys. It enables users to address the causes of aqueous corrosion by identifying its mechanistic reasons. As a result, users take informed action on how to mitigate or eliminate this risk.

Corrosion Analyzer calculates corrosion by quantifying the bulk chemistry, transport phenomena, and surface reactions through a thermophysical and electrochemical module.

The thermophysical module calculates the aqueous solution speciation and obtains concentrations, activities and transport properties of the reacting species.

The electrochemical module simulates partial oxidation and reduction process on the metal surface.

The tool reproduces the active-to-passive transition and the effects of solution species on passivity.

Effects of temperatures, pressure, pH, concentration, and velocity on corrosion are also included. Capabilities and features of Corrosion Analyzer are:

Generation of Pourbaix (E vs pH) diagrams
Calculation of general corrosion rates
Localized corrosion susceptibility
Heat treatment effect
Generation of polarization curves plots

EVS Analyzer

Extreme value statistics (EVS) is a powerful statistical techniques that is used extensively to extrapolate damage (maximum pit depth) from small samples in the laboratory to larger area samples in the field.

ScaleChem

ScaleChem is a simulation software tool that predicts scaling problems during oil and gas production. ScaleChem simulates fluid production from the reservoir to the sales point, and computes the phase mass balance, scale tendencies, scale mass, and nucleation induction times in production fluid at each location in the production line. Capabilities and features of think simulation | getting the chemistry right are:

Brine, gas and oil analysis

Scaling scenarios

Compatibility testing of brines using the stream mixing function

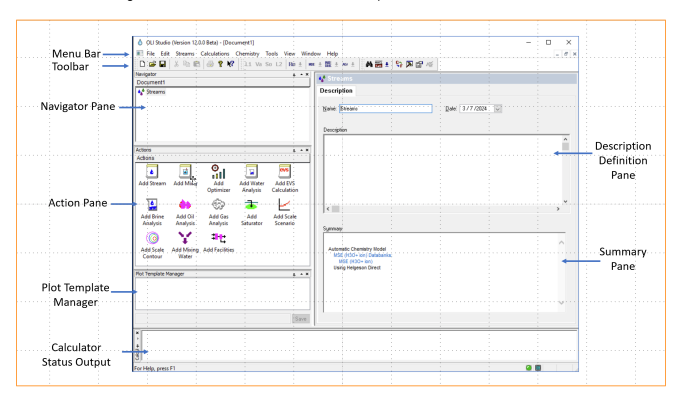
Phase equilibrium calculation for four-phase reservoir saturation, from which as whole fluid reservoir composition is determined.

Contour plots to study produced brine properties across a broad temperature and pressure range

Chapter I – The OLI Studio Software User Interface

General View of the User Interface

The OLI Studio Desktop User Interface provides the environment to create, analyze and interpret the results of your application chemistry. In this manual, an overview of the different windows and tools that you will use, as well as how to navigate the OLI Studio User Interface, will be provided here.



The screenshot shown above is what you will see when you first start OLI Studio. You can customize the desktop to your own needs. The windows can be resized, moved, docked, and detached.

Menu Bar

The Menu Bar gives access to the following options: File, Edit, Streams, Calculations, Chemistry, Tools, View, Window and Help.

File: Gives access to functionality such as New, Open, Close, and Save a file

Edit: Cut, Copy, Paste, Delete and Clear calculation results

Stream: The Streams menu contains actions that can also be performed using the Actions Pane

Calculations: The calculation menu contains all the calculations found in the Actions Pane

Chemistry: Advanced changes to the chemistry can be made here.

Tools: Gives access to tools such as component search, names manager, units manager and other customizations.

View: Gives the option to customize the view of the interface.

Window: Allows to arrange the different OLI Studio. documents in the window.

Help: Here you can have access to Technical Support or any content that you may need related to OLI Software.

Tool Bar

The Tool Bar gives quick access to different functions and sits below the Menu Bar. Tools within this bar can be added, removed or repositioned. When a tool is selected (or turned ON) it is highlighted in light blue. When deselected (or turned OFF) it comes back to gray.



They are laid out in the following order:

| Quick access to File: | |
|------------------------------------|-----------------------------------|
| Quick access to Edit: | X ₽ ® |
| Quick access to Help : | % № ? |
| Quick access to Chemistry : | L1 Va So L2 Re ± MSE ± MSE ± AQ ± |
| Quick access to Tools : | AA B ± C; ∑ B A € |

Icons in the Chemistry section

In the quick access to **Chemistry**, you find: Phases, Redox and Databanks.

Phases: Turns ON/OFF specific phases. Four different phases are available:

L1: Liquid 1 or water-rich phase

Va: Vapor phase

So: Solid phase

L2: Usually organic rich phase and sometimes a critical fluid. It is also referred to as Liquid 2 phase.

Redox: Denoted as **Re**. Turns ON/OFF Reduction/Oxidation (REDOX) reactions.

<u>Databanks</u>: Turn ON/OFF a specific thermodynamic databank. There are three thermodynamic databanks available:

MSE: Mixed Solvent Electrolyte databank (Default)

MSE-SRK: Mixed Solvent Electrolyte and Soave-Redlich-Kwong databank

AQ: Aqueous databank

Icons in the Tools section

A description of each icon in the Tools section is given below:

Component search:

This tool helps you to look for a component using Formula, CAS number, chemical name or using the periodic table.

This tool shows you the name of components in different styles in tables and reports. The style options are: Display name, Formula or OLI Name.

Units manager:

This tool allows you to select or change to preferred units for all calculations.

This tool allows you to remove or add preferred tools to the Tool Bar. For example, you can add or remove Chemistry from the Tool Bar.

General options:

This tool allows you to adjust or change default software settings.

This tools allows you to include or exclude different types of properties into

Calculation options:

Navigator Pane

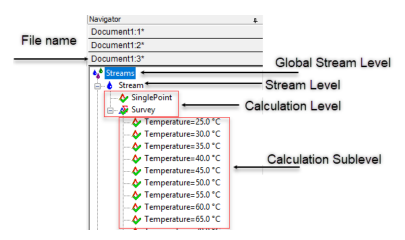
The Navigator Pane (or tree level) contains the list of streams and calculations that are active within a file. This view contains the icons and names of each action in a hierarchical tree. The Description/Definition pane changes depending on the level that is highlighted. A plus sign next to an object in the stream level indicates that that stream or object has sub-streams or branches. There are four levels: Global Stream Level, Stream Level, Calculation Level, and Calculation Sublevel.

Global Stream Level: Provides the broadest view of the navigator objects. At this level the user can define default units, default components name, and general preferred calculation options for the working file.

Stream Level: Chemistry options such as phase selection, REDOX reactions, and the thermodynamic databanks can be selected as this level.

Calculation Level: Calculation types such as Single point calculations, Survey calculations, etc., are subordinate to streams, and appear in this level. A more detailed explanation of calculation types can be found in Chapter II of this manual.

Calculation Sublevel: Some calculations, such as Survey calculations, have their own calculation sublevels. They can be expanded or maximized using the small icon (\blacksquare) next to the calculation type.

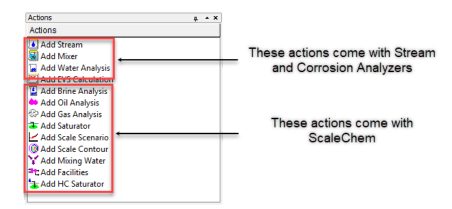


Note: A star (*) next to the file name indicates that the file has not been saved.

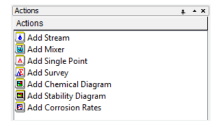
Actions Pane

This view contains selectable action icons. Each icon represents either a new stream input or new calculation. Additional actions will appear depending on what kind of stream we are working with. You can change the view of the actions pane by right clicking in the white area. You can show the icons as a List, Small Icons or Large Icons. In this case the List option was selected.

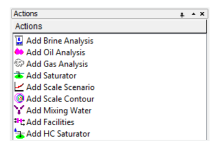
When located at the **Global Stream Level** the following action icons appear on the Actions Pane.



If you select an action item that belongs to Corrosion Analyzer, this action item will be located at the **Stream Level**, and only the following action icons appear on the Actions Pane.

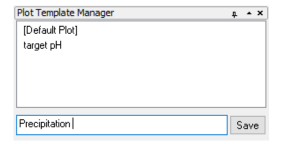


If you select an action item that belongs to think simulation | getting the chemistry right, this action item will be located at the **Stream Level**, and only the following action icons appear on the Actions pane.



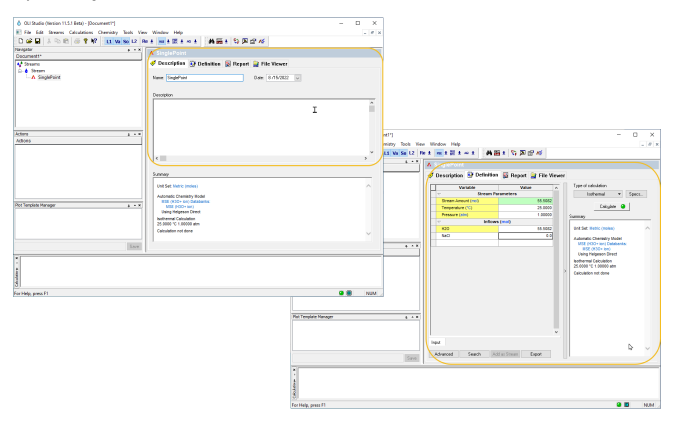
Plot Template Manager

The default location for the Plot Template Manager is in the lower left-hand corner of the main window. The Plot Template Manager is a tool that allows the user to create, save and use plot templates. This tool allows a fast plotting and analysis of the user's results.



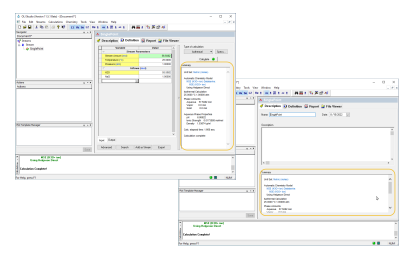
Description / Definition Pane

Users work most of the time on the Description / Definition area, which changes depending on which action object is being used.



Summary Pane

The Summary Pane can be viewed in both the Description and Definition Tabs. The Summary Pane shows inputs, calculation outputs, warnings, and has hyperlinks to the Units and Databank Managers. This window varies with object.



Calculation Status Output

The Calculation Status Output window shows progress, errors, warnings, temporary file locations, and other data.

```
Starting Calculation
Calculating SinglePoint
Unit Set: Metric (moles)
Automatic Chemistry Model
Aqueous (H+ ion)
Using K-fit Polynomials
T-span: 25.0 - 225.0
P-span: 1.0 - 201.0
* = differs from default.

Trace enabled: Level=1 Filename="C:\Users\DIANA"1.MIL\AppData\Local\Temp\OL1872a0\OL12c2a2_SinglePoint.oue"

Calculation Complete!
```

Thermodynamic Frameworks and their Databases

You can access the databanks from the Menu Bar: **Chemistry > Model Options...** or via the quick access from the Tool Bar.

There are three main thermodynamic frameworks in OLI Studio. The user can pick the thermodynamic framework that is more suitable for their chemistry. These are the **AQ** (Aqueous), the **MSE** (Mixed Solvent Electrolyte), and the **MSE-SRK** (Mixed Solvent Electrolyte – Soave Redlich-Kwong) Thermodynamic Frameworks. A more detailed description of each framework is given below:

Aqueous (AQ) Thermodynamic Framework

The OLI Aqueous (AQ) thermodynamic framework is a mature electrolyte activity coefficient model that predicts the properties of solutions up to 30 molal ionic strength. Its accompanying database contains 6,000 species for 80+ elements (metals and non-metals) across multiple oxidation states.

This AQ framework is suitable for applications involving electrolytes, gases and hydrocarbons dissolved in water.

The AQ framework utilizes the Bromley-Zemaitis activity model:

$$\log \gamma_{\pm} = -\frac{A|Z_{+}Z_{-}|\sqrt{I}}{1+\sqrt{I}} + \frac{(0.06+0.6B)|Z_{+}Z_{-}|\sqrt{I}}{\left(1+\frac{1.5}{|Z_{+}Z_{-}|}I\right)^{2}} + BI + CI^{2} + DI^{3}$$

The model can produce valid results in the following ranges:

Temperature -50°C to 300°C Pressure 0 - 1500 bar Ionic Strength 0 - 30 molal

The AQ-framework selects the PUBLIC ¹ database by default. Frequently there is more to the chemistry than what is covered in the PUBLIC database. The following is a partial list of additional databases available in the AQ framework:

Aqueous (H⁺ ion) – Public Database and selected by default Geochemical (AQ) Ceramics (AQ) Corrosion (AQ) Low Temperature (AQ) Alloys (AQ) Ion Exchange (AQ)

Surface Complexation Double Layer Model (AQ)

Mixed Solvent Electrolyte (MSE) Thermodynamic Framework

A system's thermodynamic properties are calculated from two sources; the first is from the Temperature and Pressure dependent standard-state Gibbs energies (i.e., $\overline{G_l}^0(T,P)$) of each species present. The second is from the temperature, pressure and composition dependent excess Gibbs energy (i.e., $\gamma_i(m,T)$) for each species present. In the combined relationship, the partial molal Gibbs energy of the i^{th} species is, $\overline{G_l} = \overline{G_l^0} + RT \ln m_i \gamma_i$, where $\overline{G_l^0}$ is the standard-state partial Gibbs energy and γ_i is the activity coefficient. This activity coefficient γ_i , is computed using the MSE theory. A comprehensive explanation of the MSE theory is given in Wang et al. [7].

The MSE framework utilizes the MSE activity model which contains the extended Debye-Huckel term, that accounts for **long-range** interactions, an UNIQUAC term that accounts for **short-range** interactions, and a **middle-range** that includes the ionic interactions:

$$\log \gamma_i = \log \gamma_i^{SR} + \log \gamma_i^{MR} + \log \gamma_i^{LR}$$

The model can produce results for the following ranges:

Temperature -50°C - to 90% T_{crit}

Pressure 0 - 1500 bar

Ionic Strength no limit

The MSE framework contains the following databanks:

MSE (H₃O⁺ ion) – Selected by default Corrosion (MSE) Geochemical (MSE) Urea (MSE) Surface Complexation Double Layer Model (MSE)

¹ The PUBLIC database is the older OLI database, containing nearly 70 percent of the thermodynamic data available from OLI and 100 percent of the supporting information.

Mixed Solvent Electrolyte with Soave-Redlich-Kwong Equation of State (MSE-SRK) Thermodynamic Framework

The MSE-SRK model is based on the Mixed-Solvent Electrolyte (MSE) framework, which provides a very accurate representation of electrolyte systems in both aqueous and mixed-solvent (e.g., glycol-containing) environments. The MSE-SRK framework combines an equation of state for standard-state properties of individual species, an excess Gibbs energy model to account for solution non-ideality in the aqueous electrolyte phase, and the Soave-Redlich-Kwong equation of state (SRK EOS) to calculate the properties of the gas phase. The MSE-SRK framework, however, provides a different treatment of the non-electrolyte-rich second liquid phase for liquid-liquid equilibria. MSE-SRK assumes the second (usually organic-rich) liquid phase to be non-ionic and reproduces its properties using the SRK EOS. This allows the MSE-SRK framework to reproduce the critical behavior of nonelectrolyte systems more easily.

In the MSE-SRK model, the electrolyte-containing (usually aqueous) liquid phase is represented by a combination of the Helgeson-Kirkham-Flowers (HKF) equation of state for standard-state properties and the MSE activity coefficient model for solution nonideality. Accordingly, the chemical potential of a species i in a liquid (electrolyte) phase is calculated as:

$$\mu_i^L = \mu_i^{L,0,x}(T,P) + RT \ln x_i \gamma_i^{x,*}(T,P,x)$$
 (1)

where $\mu_i^{L,0,x}(T,P)$ is the standard-state chemical potential from the HKF theory [9], [10], x_i is the mole fraction, and $x_i \gamma_i^{x,*}(T,P,x)$ is the activity coefficient from the MSE theory of Wang et al. [7], which accounts for long-range electrostatic, specific ionic, and short-range intermolecular interactions.

The second liquid phase is assumed to be non-ionic and is modeled using the Soave-Redlich-Kwong equation of state (SRK-EOS) [11]. The chemical potential in the non-ionic liquid phase is then calculated as:

$$\mu_i^G = \mu_i^{G,0}(T) + RT \ln \frac{Py_i \varphi_i(T, P, y)}{P^0}$$
 (2)

where $\mu_i^{G,0}(T)$ is the chemical potential of pure component i in the ideal gas state, y_i is the mole fraction, $\varphi_i(T, P, y)$ is the fugacity coefficient from the SRK-EOS, P is the total pressure, and $P^0 = 1$ atm.

The properties of the gas phase are also obtained from the SRK equation according to Equation (2).

The MSE-SRK framework contains the following databanks:

MSE (H₃O⁺ ion) – Selected by default MSE-SRK (H₃O⁺ ion) – Selected by default Corrosion (MSE) Geochemical (MSE) Urea (MSE) Surface Complexation Double Layer Model (MSE)

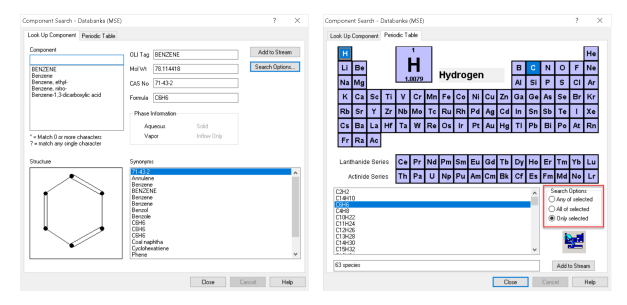
Component Search

You can access the **Component Search** from the Menu Bar: **Tools > Component Search...** or via the quick access from the Tool Bar.

Note: When using the **Component Search** tool, you should be aware that the tool will only show components in the database that you have selected.

The **Component Search** option opens a new window where you can search for the component of interest by typing the component name or look up components using the Periodic Table option. In the periodic table option, you have 3 search options:

Any of selected: Will show elements that you have selected All of selected Only selected



In any of the two options for component search, you can click on the **Add to Stream** button to add the component in your chemistry.

Names Manager

You can access the **Names Manager** from the Menu Bar: **Tools > Names Manager...** or via the quick access from the Tool Bar

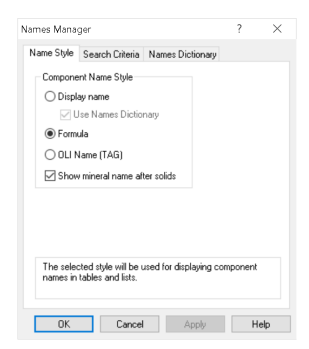
The **Names Manager** option opens a new window where you can select the component name style. There are three different styles:

Display name: This is the name that is commonly displayed when a species is entered. This is the default display. For example, Benzene, Cyclohexane, Sodium Chloride

Formula: This is the chemical formula name. For example, C6H6, C6H12, NaCl

OLI Name (TAG): This is the traditional name for the species stored internally in the OLI software. For example: BENZENE, CYCLOHEXAN, NACL (Note: These names are usually for OLI internal use).

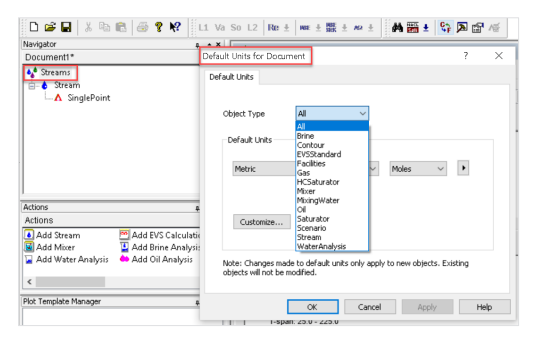
Additionally, you can select if you prefer the mineral name to appear after the solids. For example, NaCl (halite), KCl (sylvite), etc.



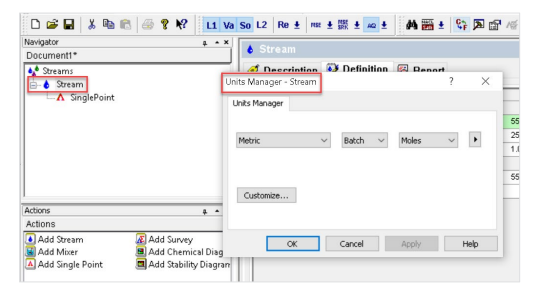
Units Manager

You can access the **Units Manager** from the Menu Bar: **Tools > Units Manager...** or via the quick access from the Tool Bar ...

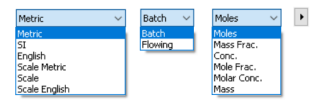
If you are changing units at the Global Stream Level, the **Units Manager** tool will open a new window where you can change the default units for the whole document. You can also select the default units of the different calculation objects.



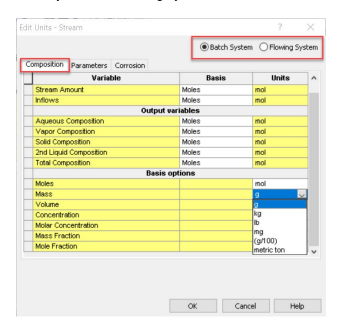
At the Stream Level, the Units Manager tool will open a slightly different window and will change the units only at the Stream Level and Calculation Level.

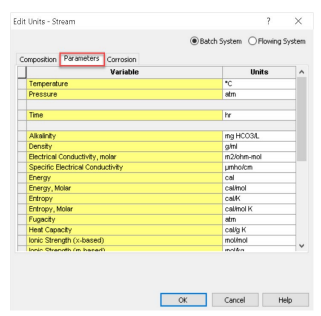


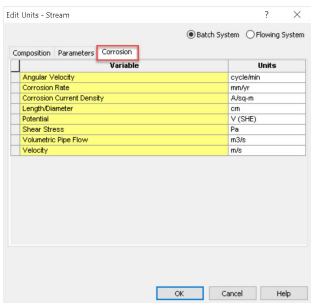
The default units are Metric, Batch and Moles, but you can change to any of the following default options:



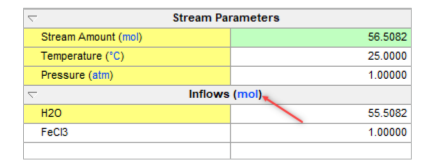
You can customize specific units, by clicking on the **Customize...** button. This will open a new window where you can customize composition units, parameters units, and corrosion units. You can also select units for a Batch system or a flowing system.







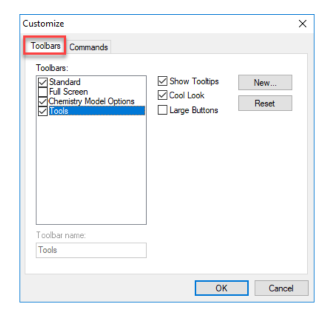
Another option to access the **Units Manager tool** is by clicking on the units highlighted in **blue**. This is a hyperlink to the Units Manager tool, and there you can make changes to the units.

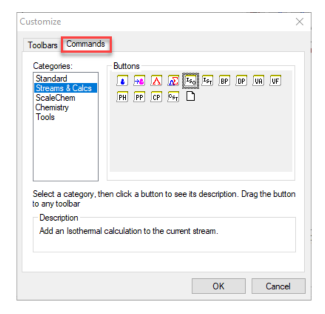


Customize Toolbars

You can access the **Customize Toolbars** from the Menu Bar: **Tools > Customize...** or via the quick access from the Tool Bar. This will open a new window where you can disable or enable toolbars under **Toolbars** tab.

Under the **Commands** tab, you can also add your preferred buttons in the tool bar. If you click on any of them, it will give you a description of the actions the selected button will perform.





Calculation Options

You can access the **Calculation Options** from the Menu Bar: **Tools > Options...** or via the quick access from the Tool Bar . This will open a new window. Under the **Calculation Options** tab you can enable or disable the following options:

General

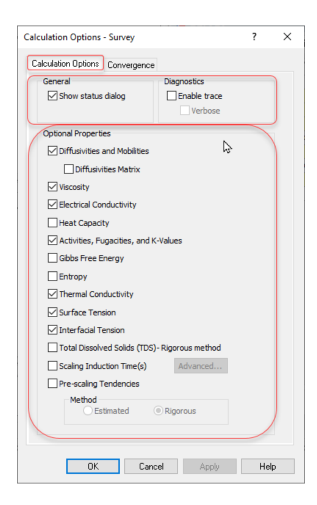
Allows you to disable or enable the status dialog (a popup window that appears after pressing the calculate button)

Diagnostics

Enable trace. This is a file containing the detailed solver output. It is generally used to diagnose why a calculation may have failed.

Optional Properties

Diffusivities, electrical conductivity, heat capacity, activities, fugacities, and K-values, Gibbs free energy, entropy, thermal conductivity, surface tension, interfacial tension, Total Dissolved Solids, Scaling Induction Time, and pre-scaling tendencies.

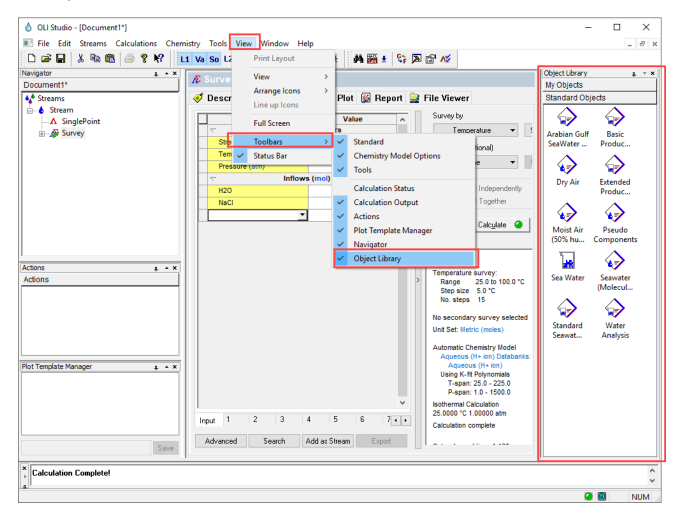


Object Library

You can access the **Object Library** from the Menu Bar: **View > Toolbars > Object Library**.

This option will create a new window (to the right). You can find commonly used objects like standard sea water, dry air, etc.

My Objects – save your own objects, such as a commonly used stream. You can drag an object and save it here for your future use.



Tip: If by accident we lose a tool bar, we can go to View > Toolbars >... and select the toolbar that is missing.

Chapter II – OLI Studio: Stream Analyzer

For all calculations we will create one or more objects, referred to as a **Streams**, which are used to define a particular chemistry, temperature, and pressure.

There are five different types of calculations that can be carried out in OLI Studio: Stream Analyzer: Single point, Survey (multiple point), Water Analysis, Mixer, and Chemical Diagram calculations. A brief definition of each type of calculation is given below.



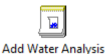
Add Stream is used to add a **New Stream** as a molecular input and define a specific chemistry.



Single Point Calculations are used to find information (pH, volume, speciation, etc.) at one specific equilibrium state.



Survey Calculations are useful for plotting changes in stream parameters against temperature, pressure, or composition.



Water Analysis allows you to enter ionic inflows, i.e., allows you to enter anions and cations.



Mixer Calculations are useful for mixing different streams. You will familiarize yourselves with its four different mixing options, Single Point Mix, Multiplier, Ratio, and Volume.

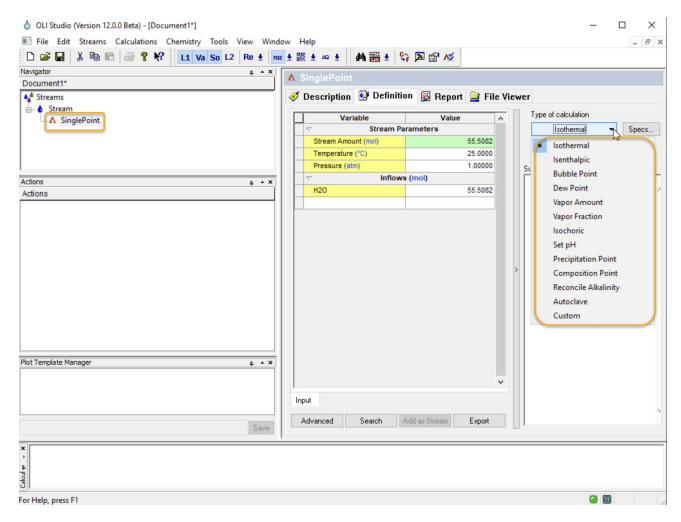


Add Chemical Diagram **Chemical Diagram Calculation** allows you to create a stability map for species based on concentration and other parameters such as pH. A contour map is created showing the user where some solids are stable and where others are not.

In this chapter several examples will be provided to cover all these calculation types.

Section 1. Single Point Calculations

Single point calculations are the simplest set of calculations in the software. There are 13 different single-point calculation types, and a brief explanation of each single point calculation is provided below.



Isothermal: The software computes solution properties based on a known chemical composition at a constant temperature and pressure.

Isenthalpic: A constant enthalpychange (loss/gain) is specified, and temperature or pressure is adjusted to meet the heat requirements.

Bubble Point: The temperature or pressure is adjusted to reach a condition where a small amount of vapor (bubble) begins to appear (a.k.a. boiling point).

Dew Point: The temperature or pressure is adjusted to reach a condition where a small amount of aqueous liquid (dew) begins to appear.

Vapor Amount: The temperature or pressure is adjusted to produce a user-specified amount (in moles) of vapor in the system.

Vapor Fraction: The temperature or pressure is adjusted to produce a user-specified amount of vapor as a fraction of the total quantity of moles in the system.

Isochoric: The temperature or pressure is adjusted to produce the user-specified total volume (constant volume calculation).

Set pH: The software adjusts the flowrate of an acid or base titrant to maintain an aqueous solution at a user-specified pH.

Precipitation Point: The software adjusts the flowrate of a species until a small amount of solid precipitates. This can also be interpreted as the solid's solubility point.

Composition Point: The composition point calculation is used to fix a species value. The software adjusts the flowrate of a species until it reaches the user - specified/fixed species value.

Reconcile Alkalinity: The software calculates or reconciles the alkalinity of a solution. There are several reconciliation types within this option: Reconcile pH, Reconcile Alkalinity and pH, and Reconcile Alkalinity, pH and TIC.

Autoclave: The software simulates a constant volume vessel (autoclave) in which mass, pressure and temperature are allowed to vary to reach a user-specified mole fraction or partial pressure of key gases in the vapor phase.

Custom: With the calculations stated so far, variables are predefined. For instance, we must select either temperature or pressure as a variable in the dew point calculation. With Custom single point calculations, we can manipulate a wider variety of variables; for example, you can set up a custom calculation to determine the solubility of a gas in solution.

In this section, we will learn how to set up each one of these single point calculations, and will also introduce how to use custom units, the names manager, modify the report, and other useful tips to get the most out of your simulation results.

Isothermal

The default and most basic single point calculation is the **Isothermal calculation**. The software computes solution properties based on a known composition, pressure, and temperature.

Example 1: Speciation and its importance for pH calculations

After completing this example, you will learn how to set up an **Isothermal calculation** and will also get a better understanding of the importance of full speciation on the calculation of pH. Let's calculate the pH of a 1 m FeCl₃ solution at 25 °C and 1 atm.

Starting the Simulation

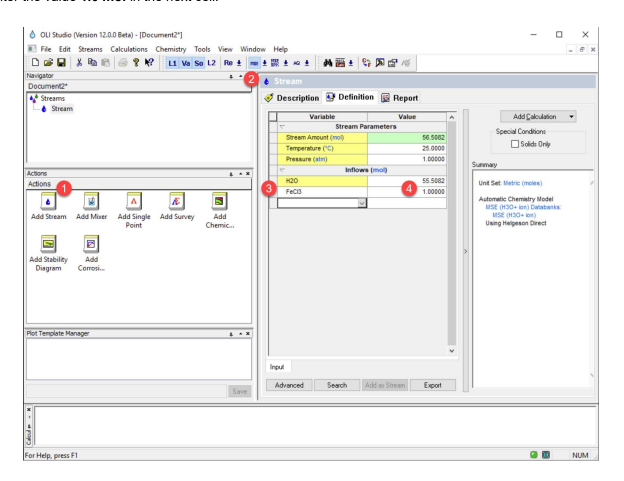
To start the software, double-click the OLI Studio icon on the desktop, which will take you to the OLI Studio interface where you can start creating your calculations.



Let's create a new stream. Click on the **Add Stream** located in the **Action Pane**. When a stream is created only H2O is present in the grid. Its cell is yellow because it cannot be removed, it is a permanent inflow.

For this example, we are going to use the **MSE-Databank** (The default databank)

Type **FeCI3** in the white cell below H2O inflows grid, and press **<Tab>** or **<Enter>** Enter the value **1.0 mol** in the next cell.



Note: By default, the software populates the stream parameters table with 25°C, 1 atm, and 55.5082 moles of water. This amount of water is 1 kg of water. All of OLI's internal aqueous concentrations are based on the molal concentration scale. You will see this value frequently throughout this manual. This effectively makes any component concentration a molal concentration.

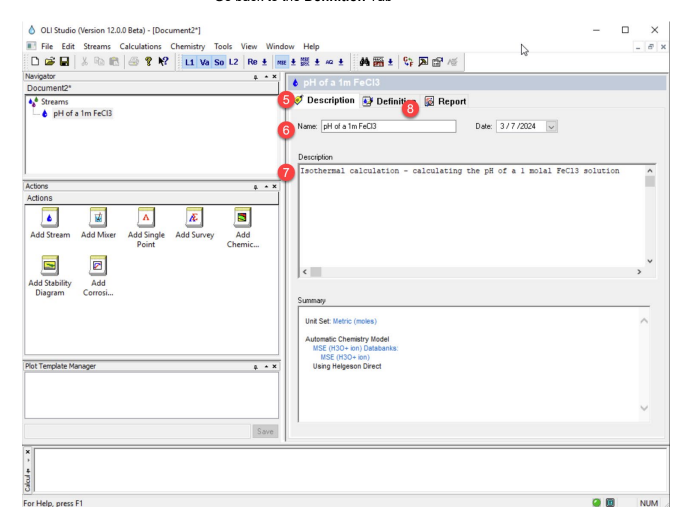
Also, notice that the stream amount will be automatically calculated from the sum of the component inflows. To indicate that the summation has occurred, the grid will highlight the stream amount cell in green.

Click the **Description** tab to change the name of the Stream. You can also change the name using the **<F2>** key or by **right-mouse clicking** on the object and selecting rename.

Change the generic name Stream to pH of 1 m FeCI3

Add the following **Description**: *Isothermal calculation - calculating the pH of FeCl3 1 molal solution*

Go back to the **Definition** Tab

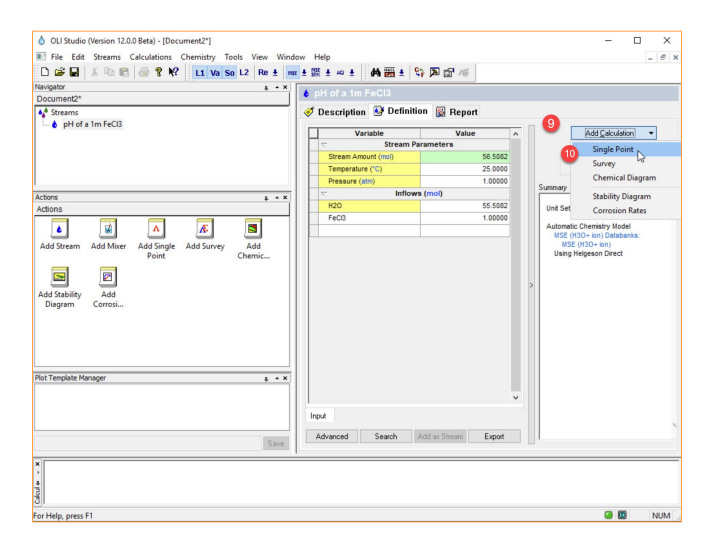


Now, we are ready to perform a calculation

Go to the Add Calculation button

Select Single Point

Note: By default, the software selects the Isothermal type of calculation



All the required variables have been entered, and the Calculate button has turned green. Click on the

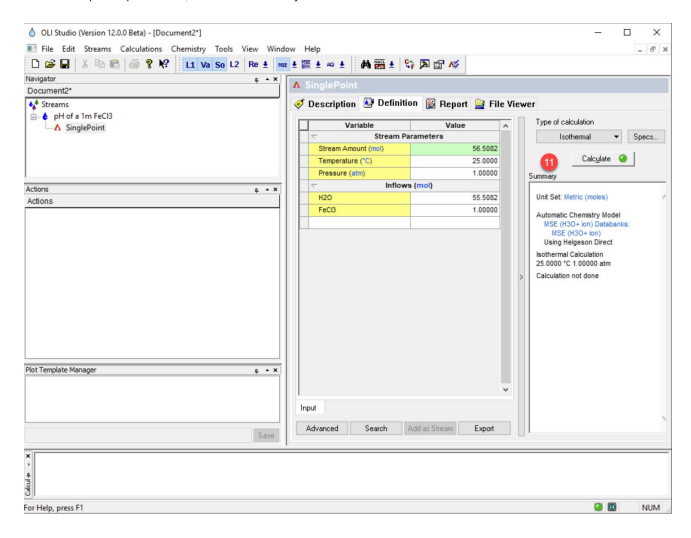
Calculate button. You can also press the **<F9>** key to run the calculation.

Note: The calculation button has three colors depending upon the specifications:

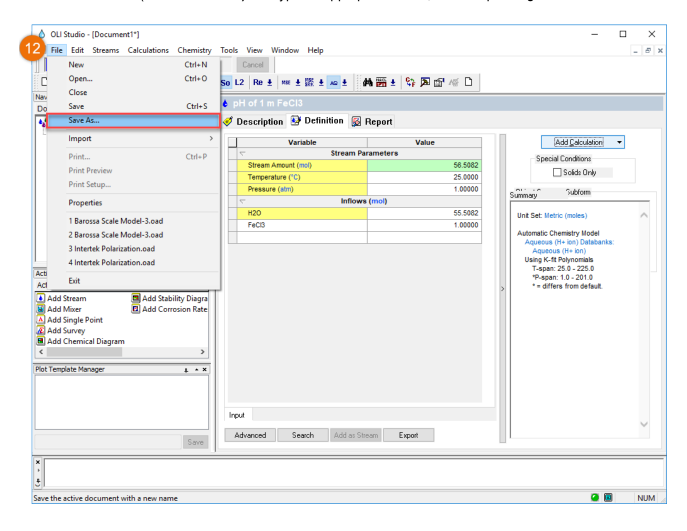
Red - Insufficient specifications for a calculation

Yellow - Incomplete specifications but calculation can continue

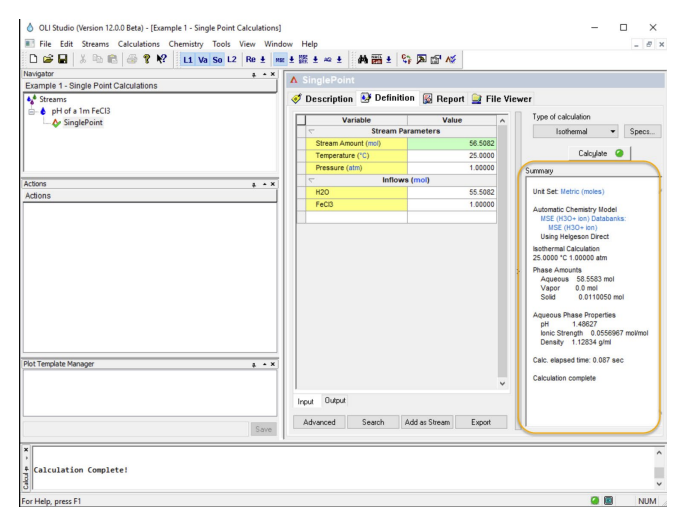
Green - Completed specifications, the calculation is ready



Please save the file (File >Save as...) and type an appropriate name, for example Single Point Calculations.



When the "Calculation is complete" a check mark appears on the calculation object . Now, let us analyze the results of the simulation.



Analyzing the Results

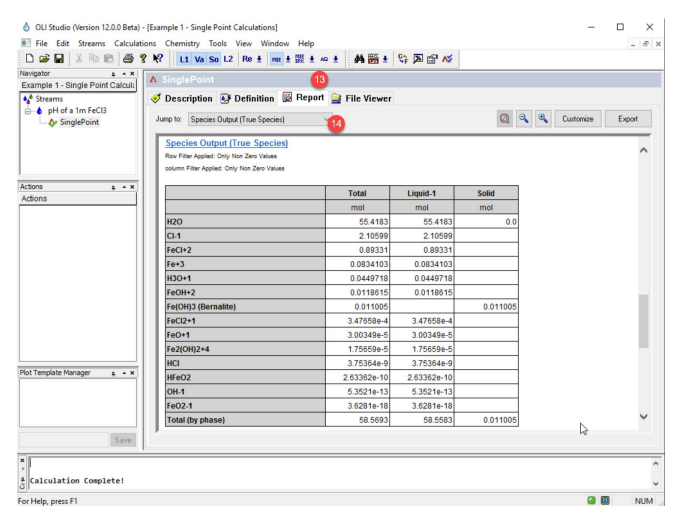
The summary box displays a partial set of results including pH, density, and volume, as well as the total calculation time.

The pH is calculated to be approximately 1.49

Let's analyze the simulation results in more detail

Click on the Report Tab

Scroll down and find **Species Output (True Species).** You will find a list of all the different species present in the aqueous phase.



Why is the pH so low?

The aqueous iron species complexes the hydroxide ions. The water dissociation reaction shifts in the direction that replenishes the hydroxide ions ².

² Le Châtelier's Principle. P.W.Atkins. Physical Chemistry. W.H. Freeman and Company, San Francisco (1982) p 269.

This equilibrium is always present:

$$H_2O = H^+ + OH^-$$
3

Speciation Reactions

First iron (III) chloride dissociates:

$$FeCl_3 \rightarrow Fe^{3+} + 3Cl^{-}$$

Then the Fe³⁺ interacts with water; hydrolysis reaction:

$$Fe^{3+} + H_2O \rightarrow FeOH^{2+} + 2H^+$$

Then another water molecule enters into the reaction:

$$FeOH^{2+} + H_2O \leftrightarrow Fe(OH)_2^+ + H^+$$

These last two reactions take up the **OH**⁻ from solution, and release **H**⁺ into the solution.

More speciation reactions occur, but the reactions above were shown as an illustration. The following is a list of all the species that are formed in the aqueous phase:

| Fe ⁺³ | FeCl ₂ ⁺¹ | Fe(OH) ₃ 0 | H_2O^0 |
|--------------------|---------------------------------|------------------------|--------------------------------------|
| FeCl ⁺² | Fe(OH) ₂ +1 | OH-1 | HCI ⁰ |
| FeOH ⁺² | H+1 | FeCl ₄ -1 | Fe ₂ (OH) ₂ +4 |
| Cl ⁻¹ | FeCl ₃ ⁰ | Fe(OH) ₄ -1 | |

pH Calculation

The pH is calculated using the following formula:

$$pH = -\log(m_H + \gamma_H + \gamma_H)$$

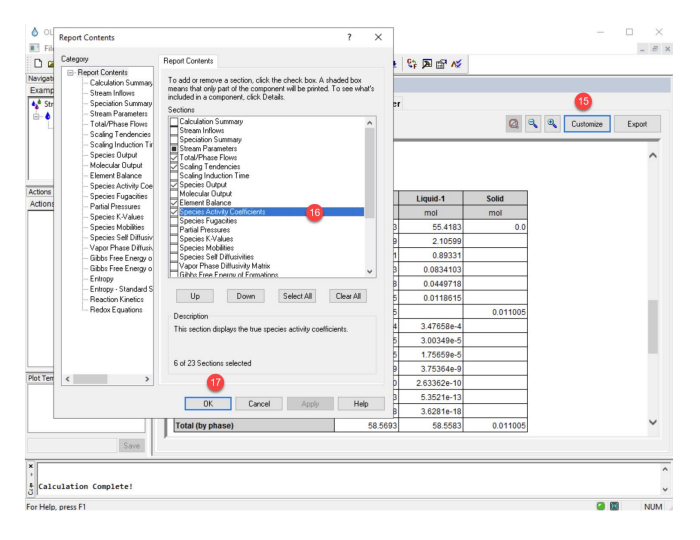
For the pH calculation the molality and the activity coefficient of the H^+ species is needed. To reveal the activity coefficients calculated by software follow the steps below.

³ This hydrolysis reaction is the generic form found in most aquatic textbooks. OLI uses this reaction in the AQ framework. In the MSE framework, a more advanced reaction is used: 2H2O=H3O+ + OH-

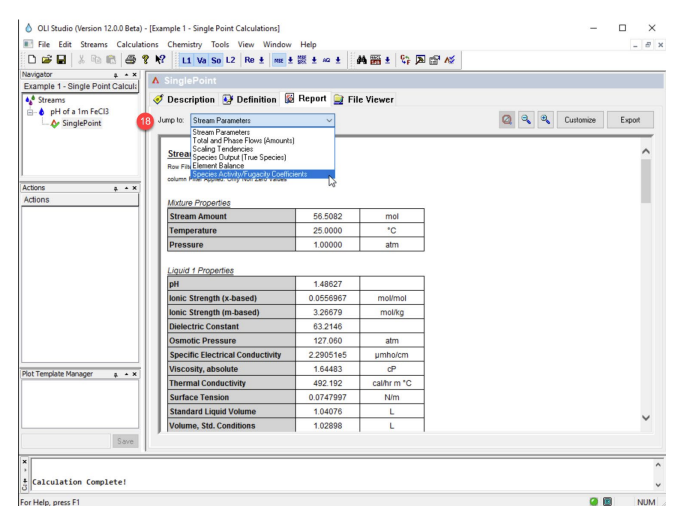
Click on the Customize Button

Select Species Activity Coefficients

Click OK



Go to the Report and scroll down or use the Jump to option and select Species Activities/Fugacity Coefficients.



Thus, the pH is:

$$pH = -\log(m_{H^{+}}\gamma_{H^{+}})$$

$$pH = -\log[(0.049012)(0.660195)]$$

$$pH = -\log[0.032357]$$

$$pH = 1.49$$

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. We save this example initially as *Single Point Calculations*.

Example 2: Calculating the pH of an acetic acid solution

In this example, we will explore an isothermal calculation, and how to set up the right units before you start your calculation. Let us calculate the pH of a 10 wt% acetic acid solution. The temperature and pressure will be 75°C and 1 atm, respectively.

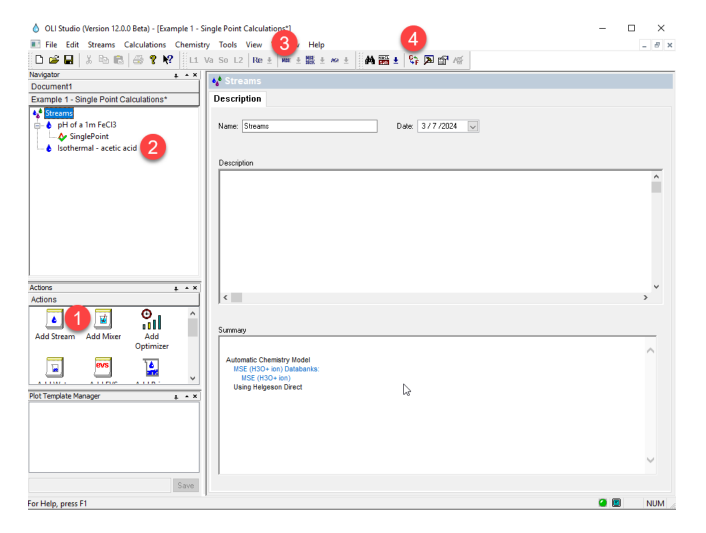
Starting the Simulation

Add a new Stream

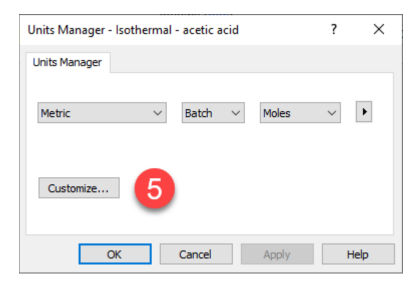
Click on the new Stream and press <F2> to change the name to Isothermal – acetic acid

Select the MSE thermodynamic Framework (selected by default)

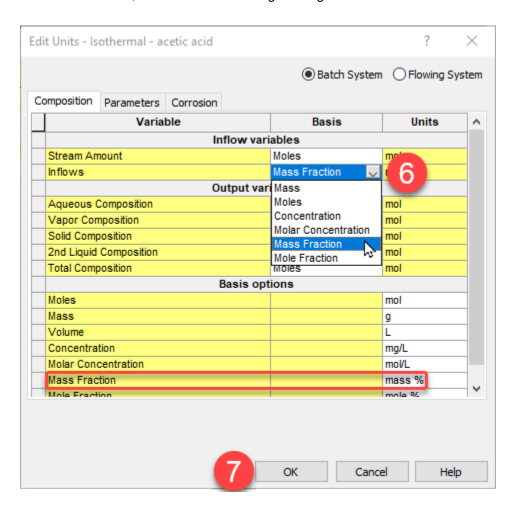
Click on the Units Manager Icon, and the Units Manger window opens



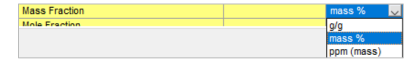
Click on the Customize Button



Let's change only the **Inflows** units. Click on the white box, **select** the drop-down arrow next to the unit, and **select Mass Fraction**. **Click OK**, to exit the Units Manager and go back to the **Definition** tab.



Note: The default unit is mass%, however you can change it to other to ppm (mass) or g/g.



We are now ready to enter the information to set up the single point calculation.

Go to the Add Calculation button and select Single Point. Select the default calculation type – Isothermal Change the name to *pH* using the <F2> key or by right-mouse click on the object and selecting rename Change the temperature to 75°C and pressure to 1 atm

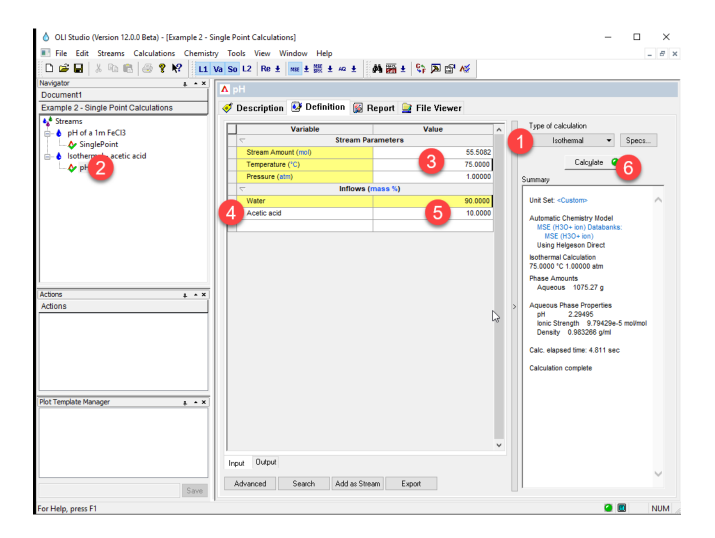
Type Acetic Acid in the white cell below H2O inflows grid, and press <Tab> or <Enter>

Note: If the name *Acetic Acid* changed to the formula type i.e., *CH3COOH*, or the OLI TAG name, i.e., *ACETACID*, you can change the name style to *Display Name* by clicking on the **Names Manager** icon

Enter the value 10 mass% in the next cell.

Note: When using mass-fraction units, it is assumed that the amount of water will be the difference of the components entered. In this case, the value field is highlighted in yellow to inform you that the value will be determined from the values of the other components.

Click on the Calculate button



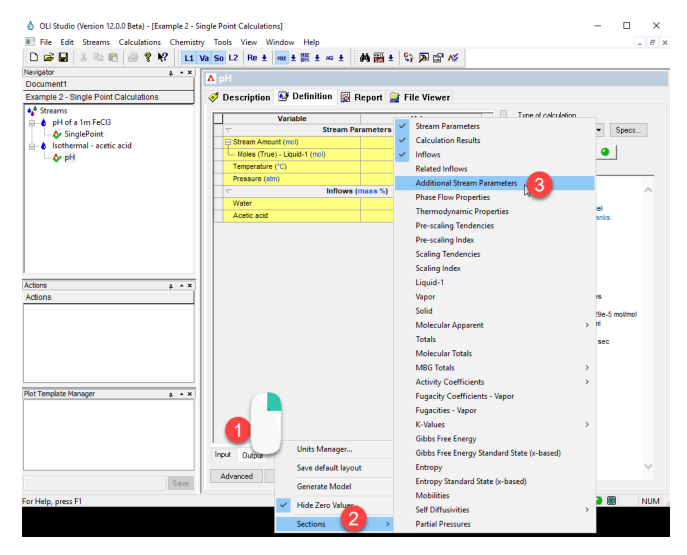
Analyzing the Results

After the calculation is complete, another way of analyzing the results is using the Output mini-tab.

Click on the Output mini-tab at the bottom of the grid

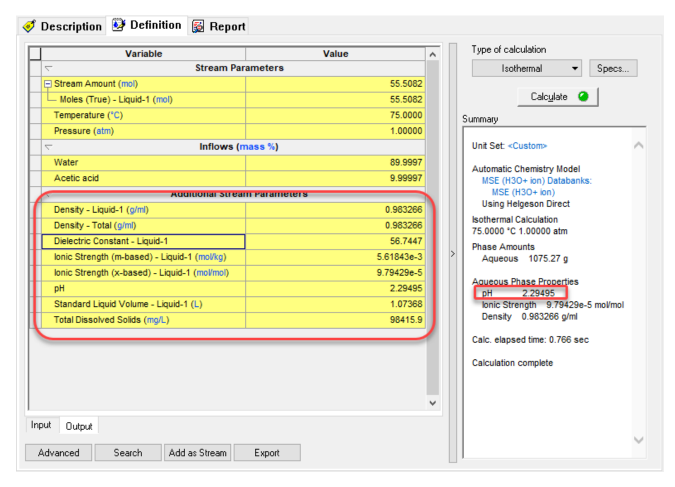
Right-click on the gray area and select Sections

Select Additional Stream Parameters



The **Sections** section offer more results such as Thermodynamic Properties, Scaling Tendencies, Equilibrium constants (K-values), etc., that you can further explore.

So, the calculated pH of this acetic acid solution is approximately 2.3.



It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created in Example 1 named as *Single Point Calculations*.

Bubble Point

Liquid boils when its vapor pressure exceeds its confining pressure. Boiling occurs by adding heat (isobaric boiling) or reducing confining pressure (isothermal boiling). Either way, the effect is the same: a vapor phase forms. Stream Analyzer calculates this phenomenon using a calculation type called **Bubble Point**. Since the OLI software does not assume an air phase, the confining pressure acts like a plunger on a liquid's surface. The pressure the plunger exerts is the pressure specified in the software.

With the software, the temperature or pressure is adjusted such that a very small amount of vapor will form (i.e. 1×10^{-10} of the total stream amount in gmoles). This is another way of saying that the bubble point calculation is a determination of the boiling point. Usually, we determine the temperature at which a solution will boil. If the pressure is set to 1 atm, then we calculate the **Normal Boiling Point**. If we set a constant temperature, then we determine the bubble point pressure.

Example 3: Calculating the Bubble Point of a 1 M Acetone Solution

In this example, we are going to compute the bubble point of a 1 M Acetone solution by changing the temperature (isobaric) and then the pressure (isothermal). Additionally, you will learn how to change the **Inflows** units from moles to moles/L (M).

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Bubble Point Calculation | | | |
|---|--------------------------|---------------|------------|
| Calculation Settings Stream Composition and (| | | |
| Calculation Type | Single Point Calculation | Stream Amount | 1 L |
| Calculation Sub-type | Bubble Point | Temperature | 25 °C |
| Stream Name | Acetone Bubble Point | Pressure | 1 atm |
| Name Style | Display Name | Water | Calculated |
| Unit Set | Custom | Acetone | 1 mol/L |
| Framework | MSE | | |

Calculating the Bubble Point Temperature (Isobaric)

Add a new Stream

Click on the new Stream and press **<F2>** to change the name to *Acetone Bubble Point* Select the **MSE** thermodynamic Framework

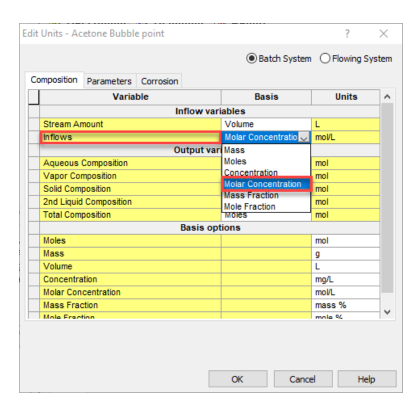
Click on the Names Manager Icon, and select the Display Name option, and click OK



Click on the hyperlink mol next to Stream Amount

| | Variable | | Value |
|---|---------------------|----------|----------|
| П | | tream Pa | rameters |
| | Stream Amount (mol) | | 55.5082 |
| | Temperature (°C) | | 25.0000 |
| Г | Pressure (atm) | | 1.00000 |
| Г | ▽ | Inflows | (mol) |
| Г | Water | | 55.5082 |
| | | | |

This will open the **Units Manager.** Under **Inflows**, select the option of **Molar Concentration**. It will automatically set the Stream amount as **Volume** in **Liters**. Then click **OK**.



Type **Acetone** in the white cell below H2O inflows grid, and press **<Tab>** or hit **<Enter>**, and then enter the value **1.0 mol/L** in the next cell.

Go to the Add Calculation button and select Single Point calculation

Change the **SinglePoint** name to **Bubble Point Temperature** using the **<F2>** key Select **Bubble point** as Type of Calculation.

A new section appears in the grid, called **Calculation Parameters**. By default, the Temperature Bubble Point calculation is selected. For this calculation leave the default calculation: **Temperature**

Variable Value Stream Parameters Stream Amount (L) 1.00000 Density correlation Parent stream condition Stream: Temperature (°C) 25.0000 Stream: Pressure (atm) 1.00000 Temperature (°C) 25.0000 1.00000 Pressure (atm) Calculation Parameters Calculate Temperature Temperature Inflows Pressure Water Acetone 1.00000

Note: In the drop-down arrow you have the option to select Bubble Point Temperature or Bubble Point Pressure.

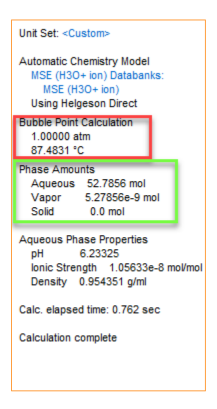
Note: There are also two dots to the left of the Temperature and Pressure rows. The yellow dot represents the "dependent" variable. The temperature value, 25.000 is colored green. This value is now an initial guess since the final value will be computed. The brown dot adjacent to the Pressure variable indicates that this potentially dependent variable is "fixed".

We are ready to perform the calculation. Click on the Calculate button or press the <F9> key.

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Single Point Calculations*.

Analyzing the Results

Review the Summary Box. The temperature at the bubble point is computed to be **87.48°C** at 1 atm. Also notice that the **Vapor** amount is 5.27e-9 moles out of the total **Aqueous** amount of 52.79 moles. This is by design; the software sets the amount of vapor at 1/10th the moles of the stream amount.



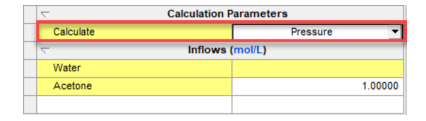
Calculating the Bubble Point Pressure (Isothermal)

This next calculation computes the bubble point pressure for the same stream.

Select the Stream named **Acetone Bubble point**

Go to the **Add Calculation** button and select **Single Point** calculation
Change the **SinglePoint** name to **Bubble Point Pressure** using the **<F2>** key
Select **Bubble point** as Type of Calculation.

Change the Calculate-Temperature row to Calculate-Pressure

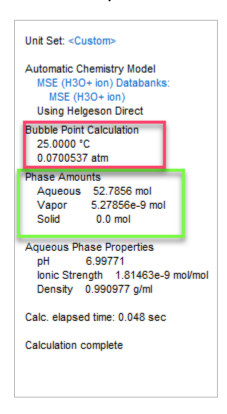


Click on the **Calculate** button or press the **<F9>** key.

It is time to save your file (File >Save as...) or using the save icon in the tool bar.

Analyzing the Results

Review the Summary Box. The bubble point pressure is computed to be **0.0700550** at 25 °C. Also notice that the **Vapor** amount is 5.27e-9 moles out of the total **Aqueous** amount of 52.78 moles.



Dew Point

The **Dew Point** is defined as the temperature at which a condensable component of a gas, for example water vapor in the air, starts to condensate into a liquid. OLI Studio: Stream Analyzer calculates this phenomenon using a calculation type called Dew Point. With the software, the temperature or pressure is adjusted such that a very small amount of liquid will form.

It is frequently useful to determine the temperature or pressure at which a gas will condense. If the pressure is set constant, we calculate the dew point temperature; if the temperature is constant, then we determine the dew point pressure.

Example 4: Calculating the Dew Point of a Simple Sour Natural Gas

In this example, we are going to compute the dew point of a simple sour natural gas, by changing the temperature (isobaric) and then the pressure (isothermal).

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Dew Point Calculation | | | | |
|-----------------------|----------------------------|---------------------------------------|-----------|--|
| Calculation Settings | | Stream Composition and Conditions | | |
| Calculation Type | Single Point Calculation | Stream Amount Default – 55.5082 moles | | |
| Calculation Sub-type | Dew Point | Temperature | 120 °C | |
| Stream Name | Sour Natural Gas Dew Point | Pressure | 100 atm | |
| Name Style | Display Formula | H2O Calculated | | |
| Unit Set | Metric, Mole Fraction | CO2 1 mole % | | |
| Framework MSE | | CH ₄ | 95 mole % | |
| | | H ₂ S | 3 mole % | |

Calculating the Dew Point Temperature (Isobaric)

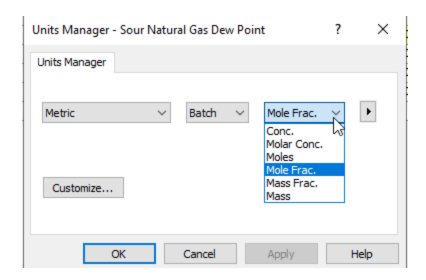
Add a new Stream

Click on the new Stream and press **<F2>** to change the name to *Sour Natural Gas Dew Point* Select the **MSE** thermodynamic Framework

Click on the Names Manager Icon, and select the Formula option, and click OK



Click on the Units Manager Icon, and select Metric, Batch, Mole Frac. This will change all inflows to mole %.



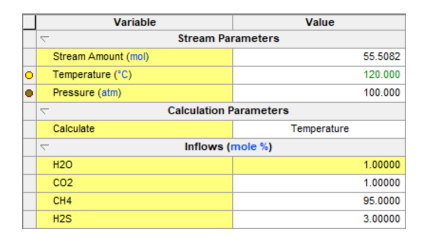
Enter the composition of the gas given in the table above

Go to the Add Calculation button and select Single Point calculation

Change the **SinglePoint** name to **Dew Point Temperature** using the **<F2>** key

Select **Dew point** as Type of Calculation

Keep the Calculate-Temperature option



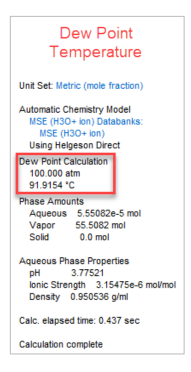
We are ready to perform the calculation. Click on the Calculate button or press the <F9> key

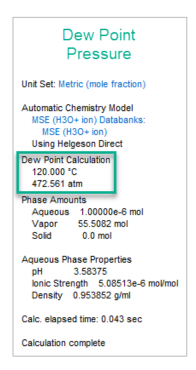
Create a **new SinglePoint** and name it **Dew Point Pressure**. Repeat the steps above, and select the Calculate-**Pressure** option

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Single Point Calculations*.

Analyzing the Results

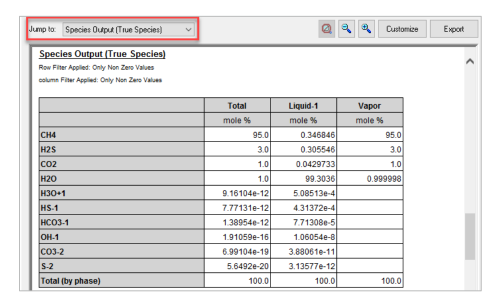
Review the Summary Box. The calculated dew point temperature is 94.9154 °C . The calculated dew point pressure is 472.561





Note: The software sets the amount of liquid to 1/10⁶ of the stream amount for the dew point calculation.

You can study the composition of the acid gases such as CO₂ and H₂S dissolved in the Aqueous phase. After you have calculated the Dew Point Pressure, click on the **Report** Tab. Scroll down or use the **Jump to** option to go to the **Species Output (True Species).**



Note: This image is for the "Dew Point Pressure" calculation

The aqueous phase, also known as Liquid-1 in the MSE framework, is shown in the center column. The dissolved CH_4 , H_2S , and CO_2 are 0.347, 0.305, and 0.043 mole % respectively.

Isochoric (constant volume)

The **Isochoric calculation** fixes the total system **volume** and adjusts the temperature or pressure. This can be understood in a gas-phase from the basis of the Ideal Gas Law:

$$PV = nRT$$

If temperature is the free variable, then the above equation becomes $T = \frac{PV}{nR}$, meaning that for a given volume, pressure, and gas moles, temperature can be obtained.

You will run a calculation, in which the system moles, temperature and volume are defined.

Example 5: Calculating the Total Pressure of a 10 L Vessel Containing Water and Air

In this example, you will add 1 kg water to a 10 L vessel and then fill the void (head space) with Air. The temperature is 25 °C.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Autoclave Calculation | | | |
|-----------------------|--------------------------|-----------------------------------|-------------------|
| Calculation Settings | | Stream Composition and Conditions | |
| Calculation Type | Single Point Calculation | Stream Amount | Default - 55.5082 |
| Calculation Sub-type | Isochoric | Ambient Temperature | 25 °C |
| Stream Name | Isochoric | H2O | 55.5082 |
| Name Style | Display Formula | N2 | 77 moles |
| Unit Set | Metric, Moles | O2 | 21 moles |
| Framework | MSE | CO2 | 1 mol |
| Calculate | Pressure | Ar | 0.04 moles |
| Vessel Volume | 10 L | | |

Add a new Stream

Click on the new Stream and press **<F2>** to change the name to *Isochoric*

Select the MSE (default) thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the gas composition in the table above

Go to the **Add Calculation** button and select **Single Point** calculation

Change the SinglePoint name to 10L Vessel using the <F2> key

Select **Isochoric** as Type of Calculation

In the **Calculation Parameters** section set **Total Volume** to **10 L** and **Calculate Pressure**. (To select Pressure, click on the drop-down arrow).

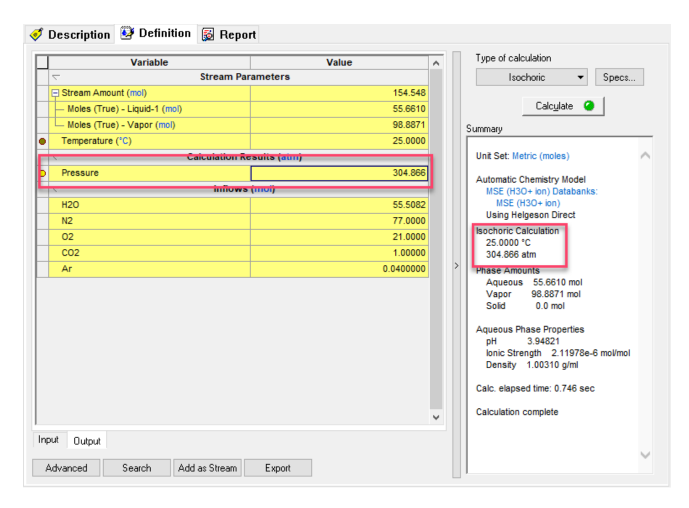
| | Variable | Value |
|---|---------------------|-------------------------------|
| | ▽ | Stream Parameters |
| | Stream Amount (mol) | 154.548 |
| • | Temperature (°C) | 25.0000 |
| 0 | Pressure (atm) | 1.00000 |
| | ▽ | Calculation Parameters |
| | Total Volume (L) | 10.0000 |
| | | |
| | Calculate | Pressure ▼ |
| | Calculate | Pressure Inflows (mol) |
| | | |
| | ▽ | Inflows (mol) |
| | √ H2O | Inflows (mol) 55.5082 |
| | H2O N2 | Inflows (mol) 55.5082 77.0000 |

We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Single Point Calculations*.

Analyzing the Results

Review the **Summary Box** or Click on the **Output-Minitab** to see the Results.



About **304.9 atm** of pressure are needed to compress the 154.55 moles of gas and liquid into a 10-L volume vessel at 25 °C.

Go to the Report tab and view the Total and Phase Flows (Amount) table.

Total and Phase Flows (Amounts)

column Filter Applied: Only Non Zero Values

| | Total | Liquid-1 | Vapor |
|-------------|---------|----------|---------|
| | mol | mol | mol |
| Mole (True) | 154.548 | 55.6610 | 98.8871 |
| Mole (App) | 154.548 | 55.6611 | 98.8871 |
| | g | g | g |
| Mass | 3874.62 | 1005.05 | 2869.56 |
| | L | L | L |
| Volume | 10.0001 | 1.00194 | 8.99811 |

The system is 10 L as defined (within the 1/10⁶ tolerance). The water phase makes up slightly more than 1 L and the balance is vapor. At this pressure, about 0.1527 moles of water evaporates, and contributes to the total moles of vapor.

Vapor Amount and Vapor Fraction

The **vapor amount** and **vapor fraction** calculations are identical in nature to the <u>Bubble Point</u> calculation, except that instead of the software defining the vapor amount as 1×10^{-10} of the total stream amount, the user defines the vapor size in either mole fraction units (vapor fraction) or mole units (vapor amount). To create a specified amount of vapor (or vapor fraction) the software can adjust the temperature (or pressure).

Example 6: Evaporating a Brine

In this example, you will concentrate a brine via evaporation adjusting the temperature, until reaching a vapor fraction amount of 95 mole %.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Vapor Fraction / Vapor Amount Point Calculation | | | |
|---|------------------------------------|-------------------|----------------|
| Calculation Settings Stream Composition and Conditi | | | |
| Calculation Type | Single Point Calculation | Stream Amount | Default - 1 kg |
| Calculation Sub-type | Vapor Fraction | Temperature | 25 °C |
| Stream Name | Brine Evaporation – Vapor Fraction | Pressure | 1 atm |
| Name Style | Display Formula | H2O | Calculated |
| Unit Set | Metric, Mass Fraction | NaCl | 9 mass % |
| Framework | MSE | CaSO ₄ | 1 mass % |

Calculating the Vapor Fraction Temperature (Isobaric)

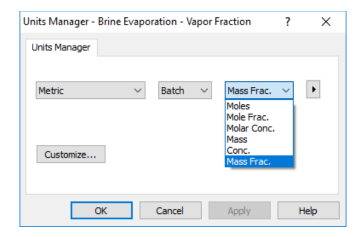
Add a new Stream

Click on the new Stream and press <F2> to change the name to Brine Evaporation – Vapor Fraction

Select the **MSE** thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the **Units Manager** Icon, and select Metric, Batch, Mass Frac. This will change all inflows to mass %.



Enter the gas composition given in the table above.

Go to the Add Calculation button and select Single Point calculation

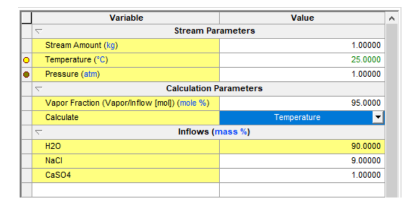
Change the SinglePoint name to Vapor Fraction Temperature using the <F2> key

Select Vapor Fraction as Type of Calculation

Enter 95 as the Vapor Fraction amount.

Note: The software will convert the 1 kg to mole units and will put 95% of this in the vapor phase.

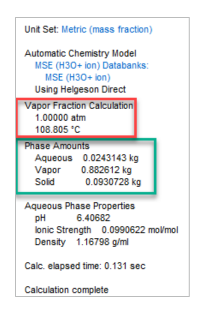
Keep the Calculate-Temperature option



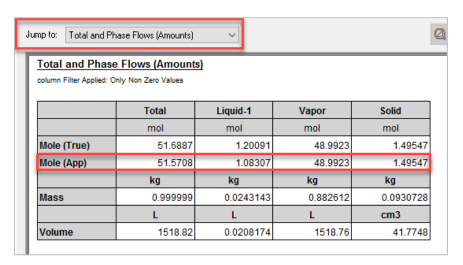
We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Single Point Calculations*.

Analyzing the Results

Review the Summary Box. The calculated temperature to put 95% (mole based) of the total stream amount in the vapor phase is 109°C, however the results in the summary box are given in mass. Let's go to the **Report**.



Select the Report tab and scroll down to Total and Phase Flow (Amounts) Table.



The apparent moles (**Mole (App)**) are the molecular representation of the system. There is a total of 51.5708 moles in this system, and exactly 95% of this system is in the vapor phase with a total of 48.9923 moles.

Set pH

The **Set pH** calculation is a useful tool in analyzing or designing a process. It allows you to create a solution that conforms to a premeasured pH. Other applications include designing a system to meet an operational pH or testing the impact of adding an acid or base to a system.

Example 7: Neutralizing Acetic Acid

In this example, you will compute the amount of base titrant required to neutralize a 1 molal acetic acid solution. You will assume a neutral system at 7 pH and 25°C.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Set pH Calculation | | | | |
|----------------------|--------------------------|-----------------------------------|-------------------|--|
| Calculation | on Settings | Stream Composition and Conditions | | |
| Calculation Type | Single Point Calculation | Stream Amount | Default | |
| Calculation Sub-type | Set pH | Temperature | 25 ℃ | |
| Stream Name | Neutralizing Acid | Pressure | 1 atm | |
| Name Style | Display Formula | H2O | 55.5082 (Default) | |
| Unit Set | Metric, Moles | СН3СООН | 1 mol | |
| Framework | MSE | Target pH | 7.0 | |

Setting the pH

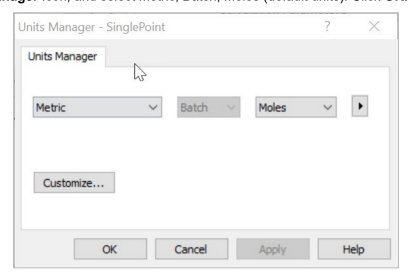
Add a new Stream

Click on the new Stream and press <F2> to change the name to Neutralizing Acid

Select the MSE (Default) thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Moles (default units). Click OK.



Enter 1 mol of CH3COOH as an inflow

Go to the Add Calculation button and select Single Point calculation

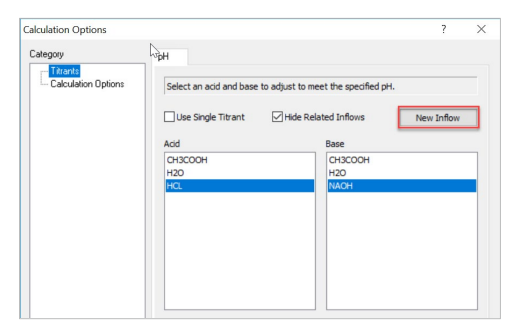
Select Set pH as Type of Calculation

Change the SinglePoint name to Neutralizing acetic acid using the <F2> key

A new grid section named **Calculation Parameters** appears. In this grid the **Target pH** can be defined, as well as the **Acid** and **Basic** titrants. The default titrants are **HCI** and **NaOH** (common) and so no additional specifications are required except to set the pH.

| | Variable | Value | |
|---|---------------------|------------|--|
| Г | | Parameters | |
| | Stream Amount (mol) | 56.5082 | |
| | Temperature (°C) | 25.0000 | |
| | Pressure (atm) | 1.00000 | |
| | | Parameters | |
| • | Target pH | 0.0 | |
| | Use Single Titrant | No | |
| | pH Acid Titrant | HCL | |
| | pH Base Titrant | NAOH | |
| | □ Inflows (mol) | | |
| | H2O | 55.5082 | |
| | СНЗСООН | 1.00000 | |

Note: The default titrants can be changed by other acids or bases by adding the desired titrant as an inflow. The titrants in the **Set pH** calculation can be also set using the **Specs** button.



The default acid and base are already set, so no additional work is required. If a different acid/base is desired, then the **New Inflow** button can be used to find additional components.

Enter 7 as the Target pH value

Use the default titrants: HCL and NAOH

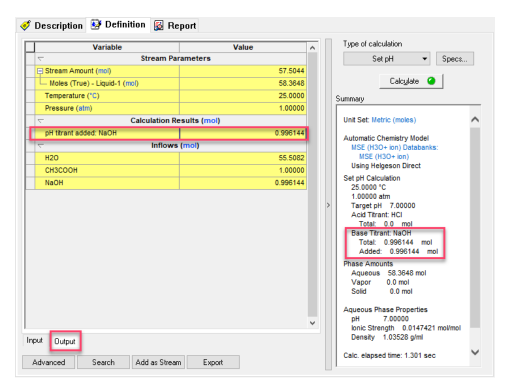
| | Variable | Value |
|---|---------------------|---------------|
| | | Parameters |
| | Stream Amount (mol) | 56.5082 |
| | Temperature (°C) | 25.0000 |
| | Pressure (atm) | 1.00000 |
| | | on Parameters |
| • | Target pH | 7.00000 |
| | Use Single Titrant | No 🔻 |
| | pH Acid Titrant | HCL |
| | pH Base Titrant | NAOH |
| | | ows (mol) |
| | H2O | 55.5082 |
| | СНЗСООН | 1.00000 |

We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Single Point Calculations*.

Analyzing the Results

Review the **Summary Box** or Click on the **Output-Minitab** to see the Results. About **0.996 moles** of NaOH is needed to neutralize 1 mole of acetic acid to 7 pH.



Precipitation Point

The **precipitation point** calculation computes the amount of material held in solution at given conditions and forces a very small amount of solid to exist. This calculation could be also called a **solubility calculation**.

Example 8: Determining the Solubility of Calcite (CaCO₃)

Equilibrium based simulators suffer from a potential problem, that the most stable solid will tend to be included over less stable (meta-stable) solids. Such is the case of calcium carbonate. Calcium Carbonate (CaCO₃) is found in nature in many forms. Two common forms are the more thermodynamically stable solids, **Calcite** and the less stable form **Aragonite**.

In this example, we will compute the solubility of CaCO₃ as **Calcite** in a solution containing Ca⁺², Mg⁺², and Cl-as well as dissolved CO₂.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Precipitation Point Calculation | | | |
|---------------------------------|---|-----------------------------------|------------------|
| | Calculation Settings | Stream Composition and Conditions | |
| Calculation Type | Single Point Calculation | Stream Amount | Default - 1 kg |
| Calculation Sub-type | Precipitation Point | Temperature | 25 °C |
| Stream Name | Solubility of CaCO3 – Precipitation Point | Pressure | 1 atm |
| Name Style | Display Formula | H2O | Calculated |
| Unit Set | Metric, Mass Fraction (ppm (mass)) | CaCl2 | 10870 ppm (mass) |
| Framework | MSE | MgCl2 | 9325 ppm (mass) |
| | | CO2 | 431 ppm (mass) |
| | | CaCO3 | 0 ppm(mass) |

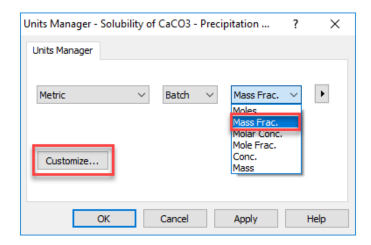
Calculating the Precipitation of Calcite (CaCO₃)

Add a new Stream

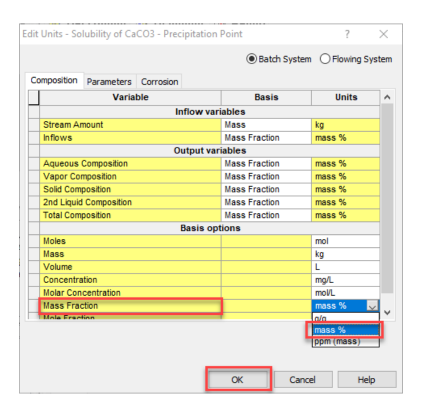
Click on the new Stream and press **<F2>** to change the name to *Solubility of CaCO3 – Precipitation Point* Select the **MSE** thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the **Units Manager** Icon, and select Metric, Batch, Mass Frac. This will change all inflows to mass%. However, you need to change the units to ppm (mass), to do this **Click** on the **Customize button.**



This will open the Edit Units window. Change the units from mass% to ppm (mass).



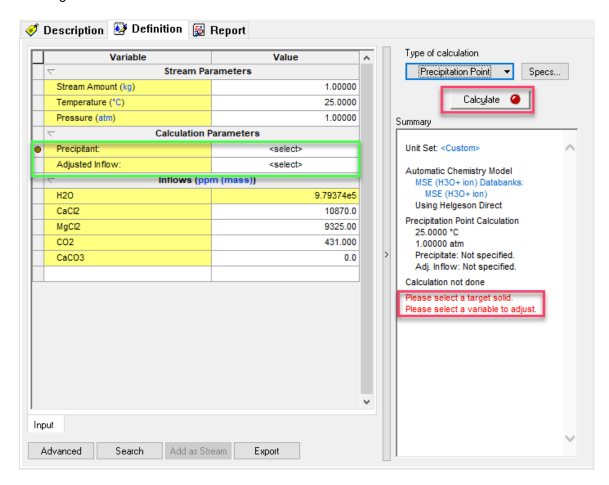
Enter the stream composition given in the table above.

Go to the **Add Calculation** button and select **Single Point** calculation

Select Precipitation Point as Type of Calculation

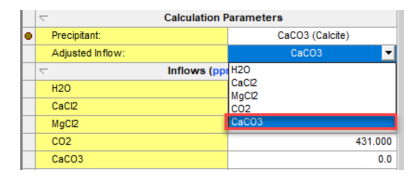
Change the SinglePoint name to Calcite Precipitation using the <F2> key

At this point you may have noticed that the Calculate button is red, and that there is a **red** text in the Summary Box with instruction that additional specs are needed. The specifications need to be defined in the **Calculation Parameters** grid section.



Select the CaCO3 (Calcite) as the Precipitant (note: you may need to scroll down)

Select CaCO3 as the Adjusted Inflow



The **Precipitant** refers to the independent variable – in this calculation, the solid phase selected will precipitate at amount of 1.0x10⁻¹⁰ times the stream amount.

The Adjusted Variable is the variable to be changed by the software until the solid target amount is achieved.

♦ OLI Studio (Version 11.5.1 Beta) - [1.7 Precipitation Point calculation_Example_8.oad*] III File Edit Streams Calculations Chemistry Tools View Window Help _ & × 🗅 😅 🖬 🖟 🖦 💼 🙆 🤻 🦎 🖟 L1 Va So L2 Re 🛨 🚾 🛨 🗯 🖢 🗯 🗯 🛨 👫 🚟 🛨 🛟 🔊 😭 🔏 Navigator д ***** × Г 1.7 Precipitation Point cal-Calculation Options ♦ Dew Point Tem New Point Pres Precipitation Point Type of calculation å lsochoric Calculation Options Precipitation Point 🔻 Specs... Hide Related Inflows New Inflow 🖶 💧 Brine Evaporation Adjusted Variable Precipitant Calculate 🥝 ---- 🔷 Vapor Fraction MgCl2.2CaCl2.6H2O H20 ... & Neutralizing Acid Summary MgCl2.12H2O Neutralizing A MaCI2 2H20 MaCI2 Solubility of CaCO MgCl2.4H20 C02 Unit Set: <Custom> Calcite Precipit MgCl2.6H2O (Bischofite) Automatic Chemistry Model MSE (H3O+ ion) Databanks MSE (H3O+ ion) MgCl2.8H20 Actions MgCl2.HCl.7H2O Actions Ma(OH)2 (Brucite) - Sol Using Helgeson Direct MgO (Periclase) - Sol Precipitation Point Calculation CaCO3 (Aragonite) - Sol 1.00000 atm Precipitate: CaCO3 (Calcite) Adj. Inflow: CaCO3 Total 313.080 ppm (mass) Total Phase Amounts
Aqueous 1.00032 kg
"nlid 1.00087e-13 kg Select the precipitant to precipitate and the adjusted variable to adjust until the solid forms. Aqueous Phase Properties 6.06957 Ionic Strength 0.0108617 mol/mol Density 1.01473 g/ml Plot Template Manager Calc. elapsed time: 1.577 sec Calculation complete

Note: The precipitation point parameters can be also set using the **Specs** button.

We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Single Point Calculations*.

OK Cancel Apply Help

Analyzing the Results

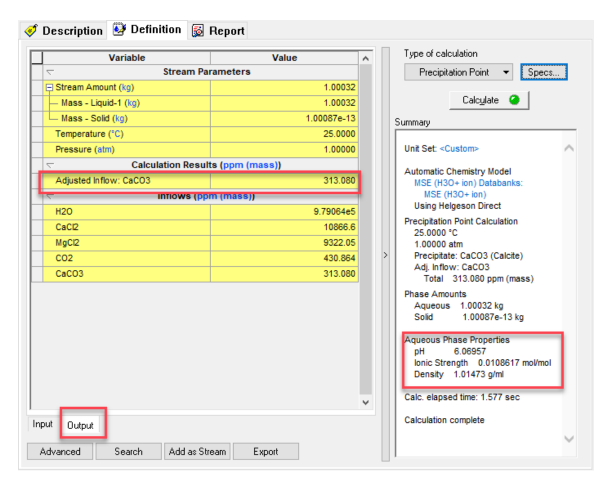
Calculation Complete!

For Help, press F1

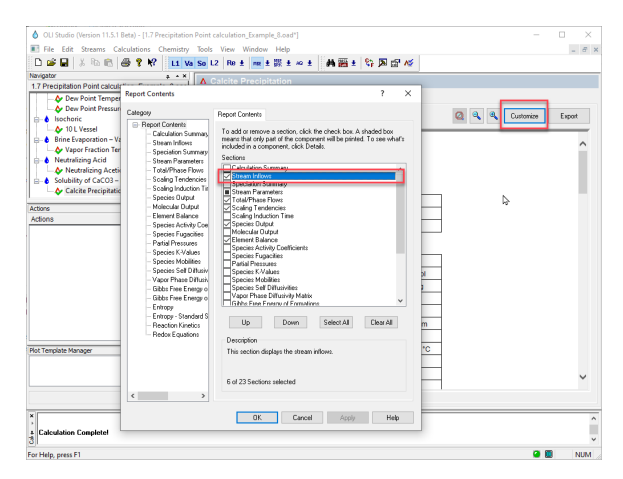
Review the **Summary Box** or Click on the **Output-Minitab** to see the Results.

The software calculated that the solubility of $CaCO_3$ as Calcite in the solution under study is around ~313.1 ppm (mass). The pH of this solution is ~6.07.

NUM



You can also check the results in the **Report** Tab. Click the **Customize** button and Select **Stream Inflows**.



Notice that the concentrations have changed slightly. This is to accommodate the additional 313.1 ppm CaCO₃ needed to meet the calculation specifications.

Stream Inflows

Row Filter Applied: Only Non Zero Values

| | Input | Output |
|---------|------------|------------|
| Species | ppm (mass) | ppm (mass) |
| H2O | 9.79374e5 | 9.79064e5 |
| CaCl2 | 10870.0 | 10866.6 |
| MgCl2 | 9325.00 | 9322.05 |
| CO2 | 431.000 | 430.864 |
| CaCO3 | 0.0 | 313.080 |

Example 9: Determining the Solubility of Aragonite (CaCO₃)

Calcite is the thermodynamically stable phase of calcium carbonate. **Aragonite**, calcite's orthorhombic polymorph, is about 1.5 times more soluble than calcite ⁴. At surface conditions, aragonite spontaneously turns into calcite over geologic time, but at higher pressures aragonite, the denser of the two, is the preferred structure.

In this example, we wish to determine the solubility of the less thermodynamically stable solid, **Aragonite**, independently of the more stable solid **Calcite**. To do this, we need to modify the **solid phases** in the chemistry model.

Starting the Simulation

Calculating the Precipitation of Aragonite (CaCO₃)

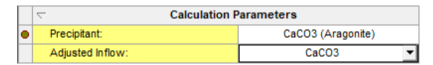
Under the Stream Solubility of CaCO3 - Precipitation Point, Add a Single Point calculation

Select Precipitation Point as Type of Calculation

Change the **SinglePoint** name to **Aragonite Precipitation** using the **<F2>** key

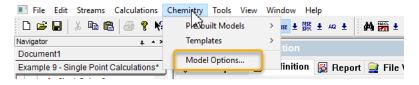
Select the CaCO3 (Calcite) as the Precipitant

Select CaCO3 as the Adjusted Inflow



For this example we need to modify the chemistry model.

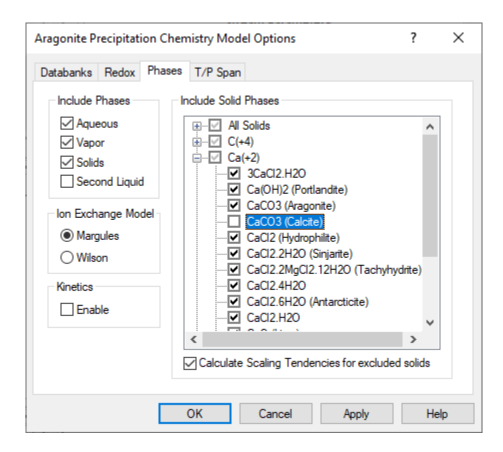
Click on Chemistry > Model Options



Select the Phases tab

Click on the "+" next to the Ca(+2) box to display all the calcium containing solids Scroll down to find CaCO3 (Calcite) and **uncheck** this solid Click **OK** continue.

⁴ https://www.sciencedirect.com/topics/earth-and-planetary-sciences/calcite



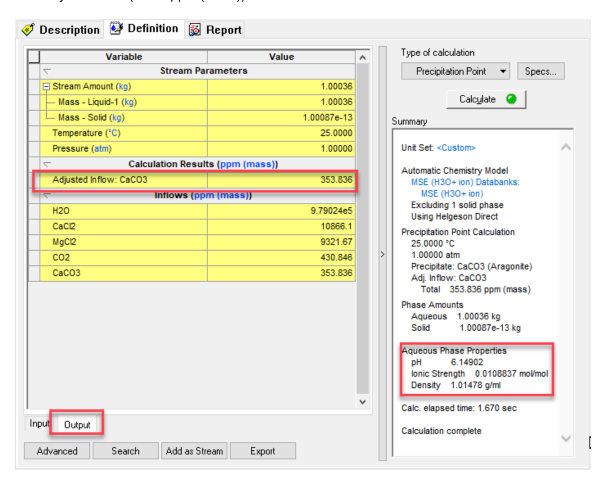
Note: The **Include Solid Phases box** lists the solids in the chemistry model. Unchecking these solids will mathematically eliminate the solid from consideration. The Scaling Tendency, however, will still be calculated.

We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Single Point Calculations*.

Analyzing the Results

Review the **Summary Box** or Click on the **Output-Minitab** to see the Results.

The software calculated that the solubility of CaCO₃ as Aragonite in the solution under study is around ~353.8 ppm (mass). The pH of this solution is ~6.15. You can see that the solubility of Aragonite is higher than the solubility of Calcite (313.1 ppm (mass)).



Example 10: Determining the Solubility of Dolomite (CaMg(CO₃)₂)

The third carbonate mineral to mention is **Dolomite**, which is something like calcite with a very high concentration of magnesium in it. Under some conditions there can be much more magnesium added in, and when the amount of magnesium becomes roughly the same as the calcium, the resulting mineral is called Dolomite with the following chemical formula: CaMg(CO₃)₂. The formation of Dolomite is believed to occur in geological time frames.

The species Dolomite is not present in the standard MSE database. However, it does exist in one of the specialty databases. This database is called *Geochemical (MSE)*.

Starting the Simulation

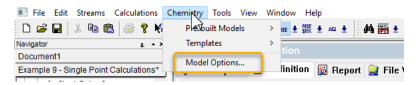
Calculating the Precipitation of Dolomite (CaMg(CO₃)₂)

Under the Stream Solubility of CaCO3 - Precipitation Point, Add a Single Point calculation

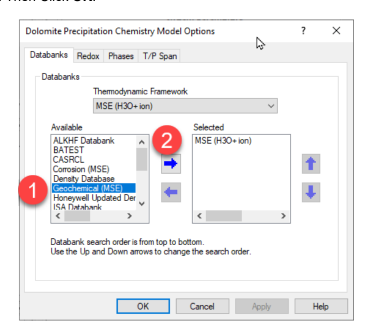
Change the SinglePoint name to Dolomite Precipitation using the <F2> key

Select Precipitation Point as Type of Calculation

Click on Chemistry > Model Options



In the Databanks Tab, select the **Geochemical (MSE)** #1Databank. You can double click or use the right arrow #2 to add the databank. Then Click **OK**.



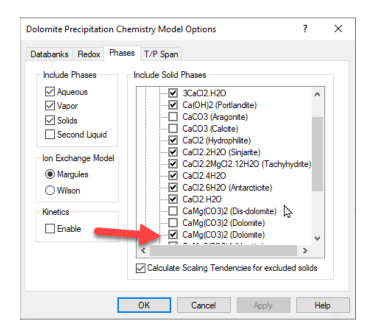
You may have additional or fewer databases depending on previous installations and imported databases. Some are from OLI and some may be private (user) databases.

Select the Phases tab

Click on the "+" next to the Ca(+2) box to display all the calcium containing solids

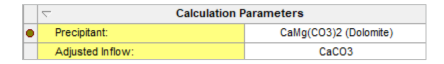
Scroll down to find CaCO3 (Calcite) and CaCO3 (Aragonite) and **uncheck** these solids. The MSE model also has ordered and disordered dolomite. These also should be unchecked. See the figure below. Make sure to keep the second dolomite species checked.

Click **OK** continue.



Note: The Include Solid Phases box lists the solids in the chemistry model. Unchecking these solids will mathematically eliminate the solid from consideration. The Scaling Tendency, however, will still be calculated.

Select the CaMg(CO3)2 (Dolomite) as the Precipitant Select CaCO3 as the Adjusted Inflow



We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

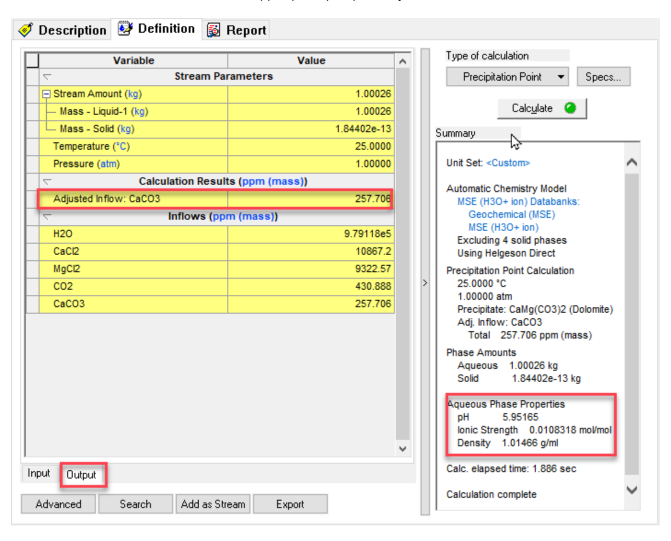
It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Single Point Calculations*.

Analyzing the Results

Review the **Summary Box** or Click on the **Output-Minitab** to see the Results.

The software calculated that the solubility of Dolomite in the solution under study is around \sim 257.7 ppm (mass). The pH of this solution is \sim 5.95.

So far, we know that the solubilities of Calcite, Aragonite, and Dolomite in a MgCl₂, CaCl₂ and CO₂ containing solution were 413.15, 552.87 and, 246.84 ppm (mass) respectively.



Composition Point

The **composition point** calculation is used to fix a species value. This calculation is useful when, for example, you want a target amount of a desired component or to fix an impurity or undesired component below a certain value.

Example 11: Targeting dissolved H₂S in water below 0.001 m

In this example, you will use the composition point calculation to target the amount of molecular H₂S dissolved in water below 0.001 m.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Composition Point Calculation | | | |
|-------------------------------|--------------------------|-----------------------------------|-------------------|
| Calculation Settings | | Stream Composition and Conditions | |
| Calculation Type | Single Point Calculation | Stream Amount | Default - 55.5082 |
| Calculation Sub-type | Composition Point | Temperature | 25 °C |
| Stream Name | Composition Point | Pressure | 1 atm |
| Name Style | Display Formula | H2O | 55.5082 (default) |
| Unit Set | Metric, Moles | H2S | 0 moles |
| Framework | MSE | Target H2S – Liq15 | 0.001 moles |
| | | Adjusted inflow | H2S |

Calculating the Vapor Fraction Temperature (Isobaric)

Add a new Stream

Click on the new Stream and press <F2> to change the name to Composition Point

Select the MSE thermodynamic Framework (this is the default)

Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the composition of the gas given in the table above

Go to the **Add Calculation** button and select **Single Point** calculation

Select Composition Point as Type of Calculation

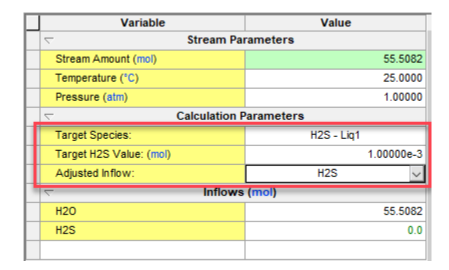
Change the SinglePoint name to H2S dissolved in water using the <F2> key

Set the **Target Species** to **H2S-Liq1** (i.e. the molecular H₂S dissolved in the Aqueous phase)

Set the Target H2S value to 0.001 moles

⁵ This used to be called the AQ or aqueous liquid value. With MSE this doesn't make sense so we have a waterrich phase = Liquid 1 and a water-poor phase = Liquid 2

Set the Adjusted Inflow to H2S

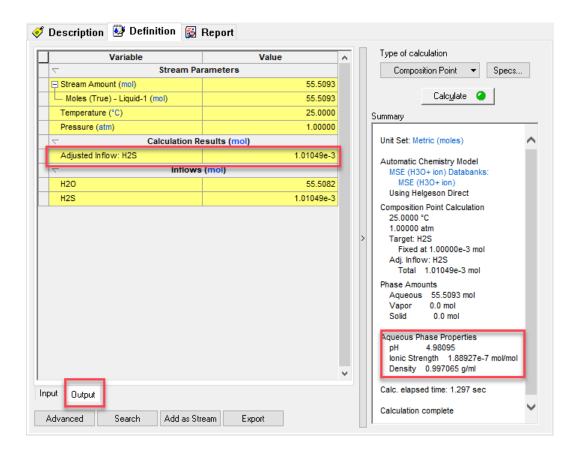


We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Single Point Calculations*.

Analyzing the Results

Review the Summary Box or Click on the Output-Minitab to see the results.



According to the calculation, \sim 0.00101 moles of H₂S were required to create a solution with 0.001 moles of H2S-aq. The pH of this solution is \sim **5.0**.

You can also go to the **Report** to see the distribution of species under the **Species Output (True Species)** table.

Species Output (True Species)

Row Filter Applied: Only Non Zero Values column Filter Applied: Only Non Zero Values

| | Total Limit 4 | |
|------------------|---------------|-------------|
| | Total | Liquid-1 |
| | mol | mol |
| H2O | 55.5082 | 55.5082 |
| H2S | 1.0e-3 | 1.0e-3 |
| H3O+1 | 1.04872e-5 | 1.04872e-5 |
| HS-1 | 1.04862e-5 | 1.04862e-5 |
| OH-1 | 9.71758e-10 | 9.71758e-10 |
| S-2 | 1.01628e-14 | 1.01628e-14 |
| Total (by phase) | 55.5093 | 55.5093 |

You can see that the molecular H2S is the value that was specified as a target. Notice that the concentration of H_3O^+ and HS^- are nearly identical. This is because the following reactions:

$$H_2S + H_2O \rightleftharpoons H_3O^+ + HS^-$$

H₃O⁺ is slightly higher because a second reaction, the dissociation of water also contributes to the H₃O⁺ formation.

$$2H_2O \rightleftharpoons H_3O^+ + OH^-$$

Autoclave

An **autoclave** is a high pressure, high temperature hydrometallurgy unit with carefully controlled conditions. From an OLI Software perspective, it is an **isochoric (constant volume) calculation** in which mass, pressure and temperature are allowed to vary.

Autoclave calculations are widely used in the upstream oil and gas applications. When performing corrosion testing, autoclave experiments are essential for simulating downstream conditions, i.e. conditions at high temperatures and pressures. Additionally, it is important to evaluate the corrosivity of production fluids by measuring solution properties, such as pH, and the concentration of aggressive species such as CO₂, H₂S, and chlorides. An autoclave, however, has a constant volume, so it is imperative to know how much material (NaCl solution, CO₂ and H₂S gases) is necessary to add at charging conditions (e.g. at room temperature) in order to reach the desired final specifications, such as partial pressure of CO₂ and H₂S gases.

We will use the Autoclave calculation to calculate partial pressures, fugacities and the final composition of key gases in the aqueous and vapor phases.

For the autoclave calculation 2 examples will be shown:

Using an Inert Gas

Calculating results at ambient conditions
Increasing the volume of the vessel and its implications
Increasing the final temperature and its implications
Increasing the final pressure and its implications

Using a Reactive Gas

<u>Calculating results at final conditions</u>
<u>Using a Reactive Gas and a NaOH solution</u>

Example 12: Using an Inert Gas

Calculating Results at Ambient Conditions

In this example, you will add 1 kg water to a 2 L autoclave and then fill the void (head space) with N₂. The ambient temperature is 25 °C. The N₂ is presumed to be the gas that remains in the autoclave headspace after sparging (O₂ removal) is completed.

The amount of N_2 added is the combination of N_2 vapor filling the headspace, and N_2 that dissolves in the water (N_2 aqueous). Since N_2 has a low water solubility, this second amount is negligible. H_2O also evaporates, so the headspace will be a mixed gas, N_2 and H_2O .

We can use the *Ideal Gas Law* to estimate the amount of N_2 needed to fill the headspace:

$$PV = nRT$$

$$1 atm \times 1 L = n \times 0.082057 \frac{L - atm}{K - mol} \times 298.15 K$$

$$n = 0.040874 \ moles$$

Although this estimate does not consider the H_2O that is in the 1 L headspace or the amount of N_2 dissolved in water, it is still an easy approach to get a reasonable estimate.

You will review the results in greater detail here because part of the goal of this first example it to show where the important variables and properties are.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Autoclave Calculation | | | | |
|-----------------------|--------------------------|-----------------------------------|-------------------|--|
| Calculation Settings | | Stream Composition and Conditions | | |
| Calculation Type | Single Point Calculation | Stream Amount | Default - 55.5082 | |
| Calculation Sub-type | Autoclave | Ambient Temperature | 25 °C | |
| Stream Name | Autoclave | Final Temperature | 25 °C | |
| Name Style | Display Formula | Final Pressure 1 atm | | |
| Unit Set | Metric, Moles | Vessel Volume | 2 L | |
| Framework | MSE-SRK | H2O | 55.5082 (Default) | |
| Results for | Ambient conditions | N2 | 0 moles | |
| | | Compute results | Ambient | |

Add a new Stream

Click on the new Stream and press **<F2>** to change the name to *Autoclave*

Select the MSE-SRK thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

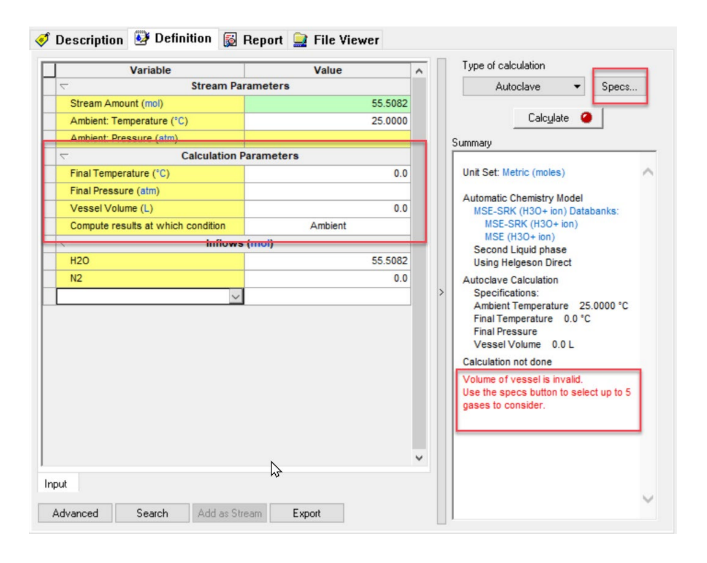
Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the N2 as an inflow

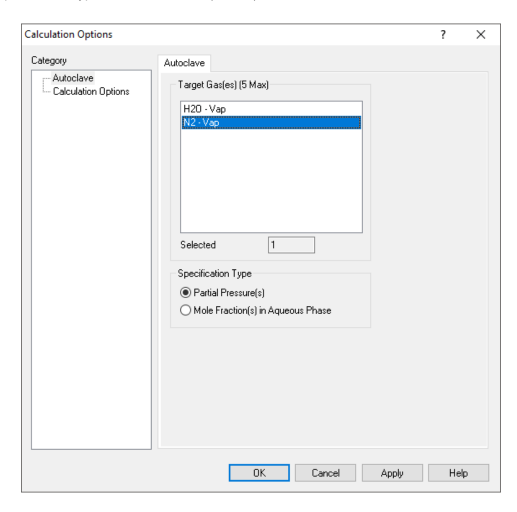
Go to the **Add Calculation** button and select **Single Point** calculation Change the **SinglePoint** name to **Autoclave – Inert Gas** using the **<F2>** key Select **Autoclave** as Type of Calculation

Note: At this point you may have noticed that the Calculate button is red, and that there is a **red** text in the Summary Box with instructions that additional specs are needed. The specifications need to be defined in the **Calculation Parameters** grid section.

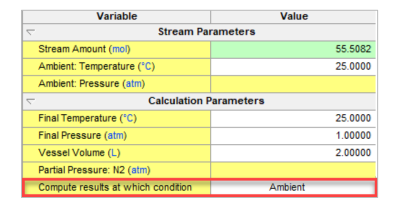
Click on the **Specs** button.



This will open the *Calculation Options* window to set up the target gas. Select **N2-Vap** and make sure to select the specification type **Partial Pressure** (default). Click **OK**.



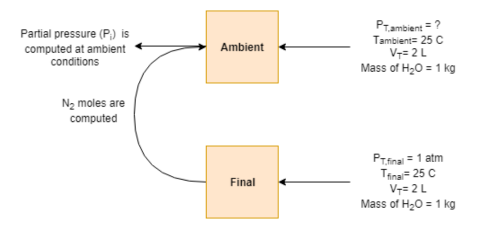
Set the Final Temperature to 25°C, the Final Pressure to 1 atm and the Vessel Volume to 2 L Set the Compute results at which Conditions to Ambient



At this point the system is defined as 1 kg H_2O (55.5082 moles) at an ambient temperature of 25 °C. The partial pressure of N_2 will be calculated at a final temperature, 25 °C and pressure, 1 atm in a 2 L vessel.

Note: The Autoclave calculation pathway

When performing an **autoclave calculation**, the software will perform two computations: First, at the final conditions, and the second one at ambient (initial) conditions. It will compute the amount (in moles) of N_2 needed to create a 1 atm pressure at final conditions. It will then use this N_2 to compute the pressure at ambient conditions. The calculation pathway is shown in the image below.

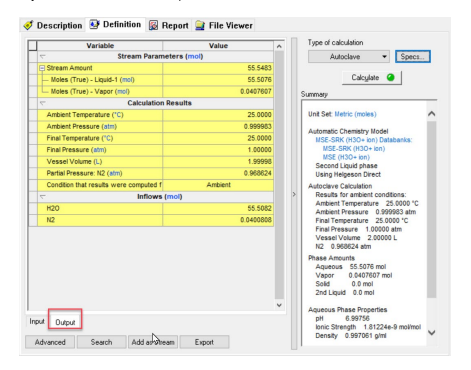


We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Single Point Calculations*.

Analyzing the Results

Review the Summary Box or Click on the Output-Minitab to see the results.



The calculated ambient pressure is 1 atm. It is identical to the final pressure, which is not surprising, since the ambient and final temperatures are the same, and by design, the autoclave calculation uses the identical inflows for ambient and final conditions.

The amount of N_2 added is 0.0400808 moles, which produces a \sim 0.9686 atm partial pressure. The computed N_2 inflow compares to the Ideal Gas value of 0.040874 moles, a \sim 2% deviation.

Let's review the distribution of N_2 in the liquid and vapor phase in the **Report** tab. Scroll down to the last table named **Element Distribution**.

The last row is N (N is molecular nitrogen). The distribution is 98.5% in the vapor phase and 1.5% in the liquid phase. Also notice that around 0.0023% of the water has evaporated (see the H(+1) and O(-2) rows).

Element Distribution

| | Total | Total | Liquid-1 | Vapor |
|-------|-----------|-----------|------------|------------|
| | mol | mole % | % of Total | % of Total |
| H(+1) | 111.016 | 66.6346 | 99.9977 | 2.30275e-3 |
| O(-2) | 55 5082 | 33 3173 | 99 9977 | 2 30275e-3 |
| N(0) | 0.0801616 | 0.0481148 | 1.49271 | 98.5073 |

These results infer two effects:

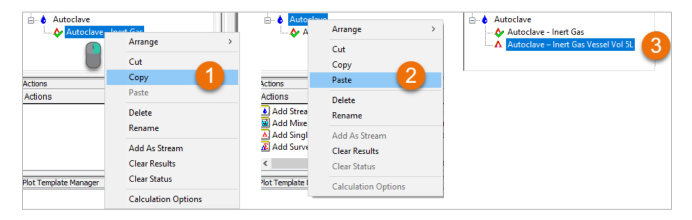
As the final pressure (P_T) increases, additional N_2 will dissolve in the water, shifting the fraction of nitrogen to the liquid.

As the total autoclave volume (V_T) increases relative to the input liquid (which stays constant), additional water will evaporate, shifting the water to the vapor. This case will be shown next.

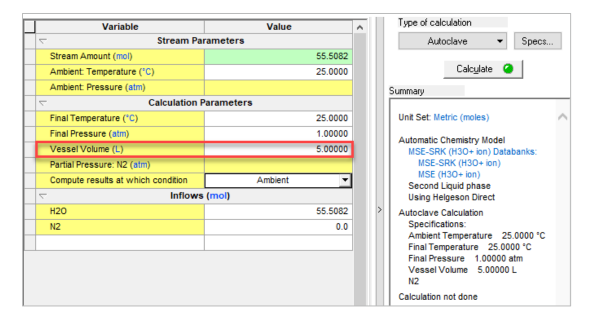
Increasing the Vessel Volume to 5 L

You are going to use the same case that you built up in the example above, with the only difference that you will change the **Vessel Volume**.

Right click on the *Autoclave – Inert Gas* single point calculation and select **copy** Selecting the stream named **Autoclave**, right click on it and select **paste** Rename the copied stream *Autoclave – Inert Gas Vessel Vol 5L*



Change the Vessel Volume to 5 L and Calculate <F9>

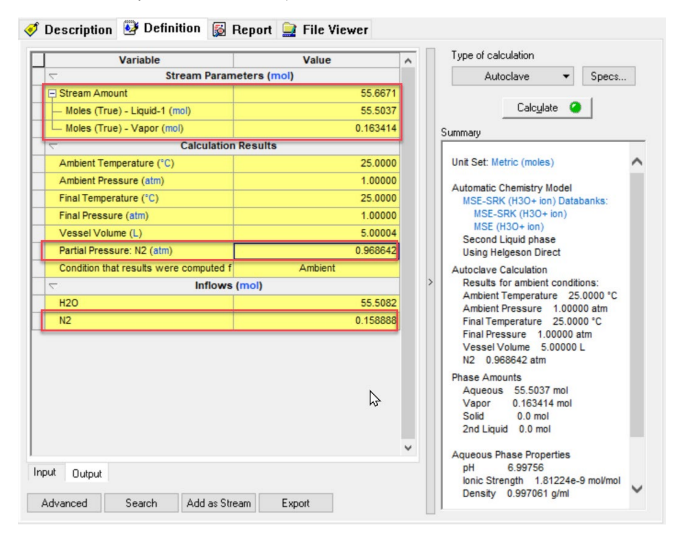


At this point the system is defined as 1 kg H_2O (55.5082 moles) at an ambient temperature of 25 °C. The partial pressure of N_2 will be calculated at a final temperature, 25 °C and pressure, 1 atm in a 5 L vessel.

It is time to save your file (File >Save as...) or using the save icon in the tool bar.

Analyzing the Results

Review the Summary Box or Click on the Output-Minitab to see the results.



The moles of liquid are now 55.5037. The reduction of 0.0039 moles (compared to the case of 2 L which was 55.5076 moles) is because this amount of H_2O evaporated into the larger headspace. This phase distribution is important to modeling the autoclave system properly, especially at elevated conditions as will be seen in the next example.

The amount of N_2 required to fill the 4-L headspace is 0.1589 moles; this amount is ~4-times greater than the first case, which is expected for gas with a low water solubility.

Increasing the Final Temperature

In this example, you will add 1 kg water to a 5 L autoclave and then fill the void (head space) with N₂. The ambient temperature is 25 °C, and the final conditions will be 1 atm and 100 °C.

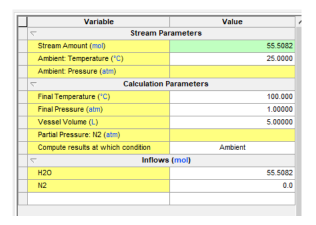
The purpose of this example is to raise the H_2O partial pressure relative to N_2 . In the 5 L case above, the N_2 inflow is 0.1634 moles. As temperature increases, the H_2O vapor pressure increases, and the amount of N_2 required to fill the vapor void will decrease.

You are going to use the same case that you built up in the example above, with the only difference that you will change the **Final Temperature** to **100 °C** and the **Final Pressure** to **50 Atm**.

Copy the Autoclave – Inert Gas Vessel Vol 5L single point calculation and paste under the Autoclave Stream.

Name the new calculation object Autoclave – Inert Gas Tf=100C

Change the Final Temperature to 100°C and Calculate <F9>

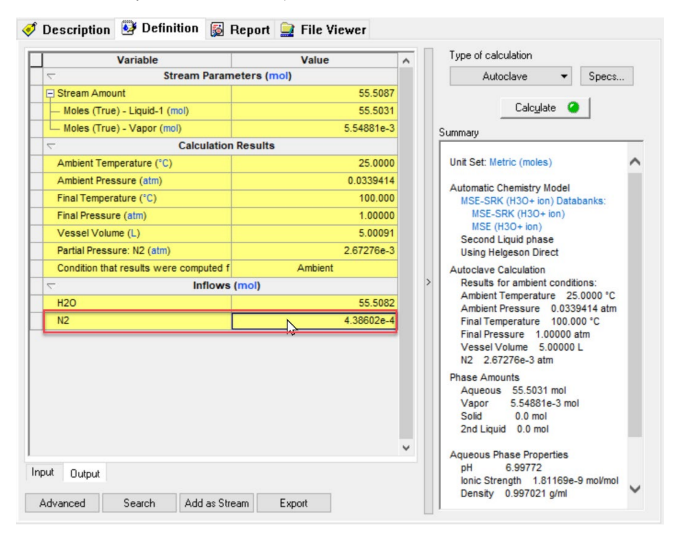


At this point the system is defined as 1 kg H_2O (55.5082 moles) at an ambient temperature of 25 °C. The partial pressure of N_2 will be calculated at a final temperature, 100 °C and pressure, in a 5 L vessel.

It is time to save your file (File >Save as...) or using the save icon in the tool bar.

Analyzing the Results

Review the Summary Box or Click on the Output-Minitab to see the results.

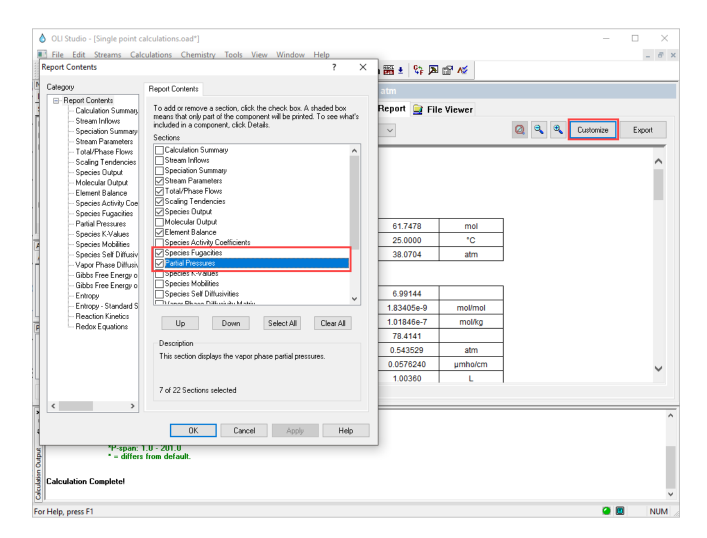


The amount of N_2 added is very small (~0.0004 moles). This is to be expected, because the vapor pressure of pure water at 100 °C is 1 atm. Therefore, a negligible amount of N_2 is needed to raise the pressure. Notice also that the calculated ambient pressure is 0.0339 atm. The vapor pressure of pure water at 25° C is 0.0313.

The results above imply that the autoclave will have a vacuum pressure at the ambient temperature, and when heated to 100 °C will reach atmospheric pressure.

In autoclave testing, it is also of utmost importance to study and understand the partial pressures and fugacity of key gases at the final conditions. You need to enable these properties to be shown in the **Report**.

Go to the **Customize** button and enable both the **Species Fugacities** and the **Partial Pressures** options, and then click **OK**.



Notice that the **Fugacity** and **Partial Pressure** of N_2 are nearly identical. This is expected since at low pressure the fugacity approaches its partial pressure.

Species Fugacities

 Species
 atm

 H2O
 0.0312529

 N2
 2.67376e-3

Partial Pressures

Row Filter Applied: Only Non Zero Values

| Species | atm |
|---------|------------|
| H2O | 0.0312686 |
| N2 | 2.67276e-3 |

Increasing the Final Pressure

In this example, you will add 1 kg water to a 5 L autoclave and then fill the void (head space) with N_2 . The ambient temperature is 25 °C, and the final conditions will be 100 atm and 100 °C.

You are going to use the same case that you built up in the example above, with the only difference that you will change the **Final Pressure** to **100 atm**.

Copy the Autoclave – Inert Gas Tf=100C single point calculation and paste under the Autoclave Stream.

Name it Autoclave – Inert Gas Pf=100 atm

Change the Final Pressure to 100 atm and Calculate <F9>

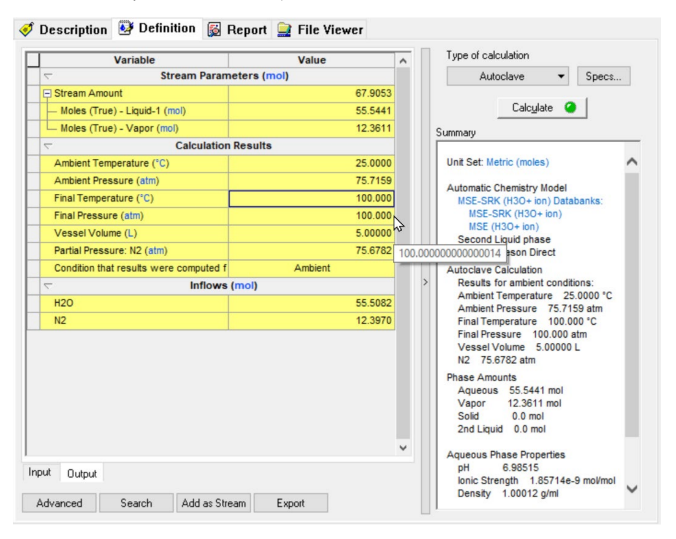
| Variable | Value | | |
|------------------------------------|-------------------------|--|--|
| | Stream Parameters | | |
| Stream Amount (mol) | 55.5082 | | |
| Ambient: Temperature (°C) | 25.0000 | | |
| Ambient: Pressure (atm) | | | |
| Calculation Parameters | | | |
| Final Temperature (°C) | 100.000 | | |
| Final Pressure (atm) | 100.000 | | |
| Vessel Volume (L) | 5.00000 | | |
| Partial Pressure: N2 (atm) | | | |
| Compute results at which condition | which condition Ambient | | |
| ▽ Inflows (mol) | | | |
| H2O | 55.5082 | | |
| N2 | 0.0 | | |

At this point the system is defined as 1 kg H_2O (55.5082 moles) at an initial temperature of 25 °C. The partial pressure of N_2 will be calculated at a final temperature of 100 °C and a final pressure of 100 atm in a 5 L vessel.

It is time to save your file (File >Save as...) or using the save icon in the tool bar.

Analyzing the Results

Review the Summary Box or Click on the Output-Minitab to see the results.



The computed ambient pressure $P_{T,A}$ is ~75.72 atm. Of this total, 75.68 atm is N_2 (P_{N2} =75.68 atm) and 0.0377 atm is H_2O (P_{H2O} =0.0377 atm). Thus, to create a 100 atm final pressure, the autoclave would need to be charged with N_2 at a regulator pressure of 75.68 atm.

According to the calculation, the amount of N_2 in the autoclave is 12.397 moles. This amount of N_2 plus the 55.5082 moles of H_2O produces a final pressure of 100 atm in a 5 L autoclave at 100 °C.

Let's study the partial pressure and fugacity of N_2 at these final conditions. Remember, you need to enable these properties to be shown in the **Report**.

Go to the **Customize** button and select both the **Species Fugacities** and the **Partial Pressures** tables, and then click **OK**.

For comparison purposes, the **Species Fugacities** and the **Partial Pressures** for 100 °C and 1 atm, and 100 °C and 100 atm are shown.

100 °C and 1 atm

Species Fugacities

Row Filter Applied: Only Non Zero Values

| Species | atm |
|---------|------------|
| H2O | 0.0312529 |
| N2 | 2.67376e-3 |

Partial Pressures

Row Filter Applied: Only Non Zero Values

| Species | atm |
|---------|------------|
| H20 | 0.0312686 |
| N2 | 2.67276e-3 |

100 °C and 100 atm

Species Fugacities

Row Filter Applied: Only Non Zero Values

| Species | atm |
|---------|-----------|
| H2O | 0.0330213 |
| N2 | 76.0121 |

Partial Pressures

Row Filter Applied: Only Non Zero Values

| Species | atm |
|---------|-----------|
| H2O | 0.0376589 |
| N2 | 75.6782 |

Notice that at low pressures, the partial pressures and fugacities values of both H_2O and N_2 are nearly identical. At the higher pressure, i.e., 100 atm, the partial pressures and fugacities of H_2O and N_2 start to differ.

The cause of this deviation is the *non-ideal effects of high pressures on gas molecules as they are forced closer together*. At low pressures, gas molecules are too far apart to interact. As pressure increases, the average distance between two molecules decreases until at short range, intermolecular forces begin to impact the molecular properties. The impact of these forces is quantified in the fugacity term. Notice that fugacity and partial pressure have the same units, atm.

Thus, when reviewing these two tables, the partial pressure can be seen as the idealized gas property at high pressures (which does not take into account intermolecular forces). The fugacity is the real gas property at this pressure. The non-ideal effect of pressure is therefore encapsulated in the following relationship:

$$\varphi = \frac{Fugacity}{Partial\ Pressure} = \frac{f}{p_i}$$

Where,

 φ is the fugacity coefficient

At low pressures, a gas molecule fugacity (f) approaches its partial pressure (p_i) , i.e. $f \to p_i$ or $\varphi = 1$.

Example 13: Using Reactive Gases

Calculating Results at Final Conditions

In this example, you will add 1 kg water to a 2 L autoclave and then fill the void (head space) with CO₂. The ambient temperature is 25 °C.

You will then modify the case further by adding NaOH so that additional CO₂ dissolves in water to form bicarbonate.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Autoclave Calculation | | | | |
|-----------------------|--------------------------|-----------------------------------|-------------------|--|
| Calculation Settings | | Stream Composition and Conditions | | |
| Calculation Type | Single Point Calculation | Stream Amount | Default - 55.5082 | |
| Calculation Sub-type | Autoclave | Ambient Temperature | 25 °C | |
| Stream Name | Autoclave – Reactive Gas | Final Temperature | 25 °C | |
| Name Style | Display Formula | Final Pressure | 1 atm | |
| Unit Set | Metric, Moles | Vessel Volume | 2 L | |
| Framework | MSE-SRK | H2O | 55.5082 (Default) | |
| Results for | Final Conditions | CO2 | 0 moles | |
| | | NaOH | 0 moles | |

Add a new Stream

Click on the new Stream and press <F2> to change the name to Autoclave Reactive Gas

Select the MSE-SRK thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the CO2 and NaOH as inflows

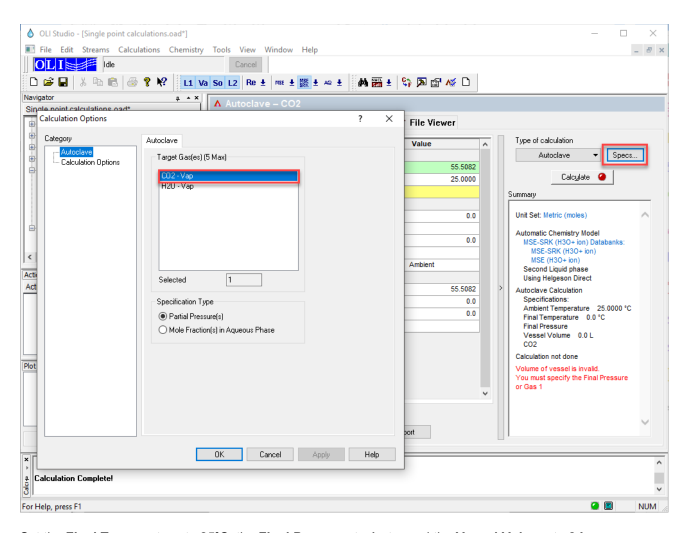
Go to the **Add Calculation** button and select **Single Point** calculation

Change the SinglePoint name to Autoclave - CO2 using the <F2> key

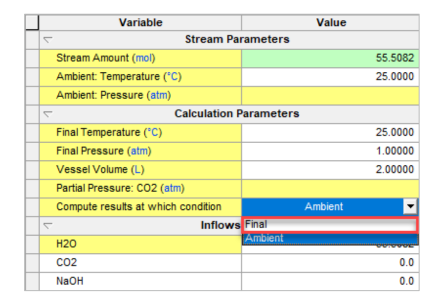
Select Autoclave as Type of Calculation

Click on the **Specs** button. This will open the *Calculation Options* window to set up the target gas.

Select CO2 and make sure to select the specification type Partial Pressure (default). Click OK.



Set the Final Temperature to 25°C, the Final Pressure to 1 atm and the Vessel Volume to 2 L Set the Compute results at which Condition to Final.



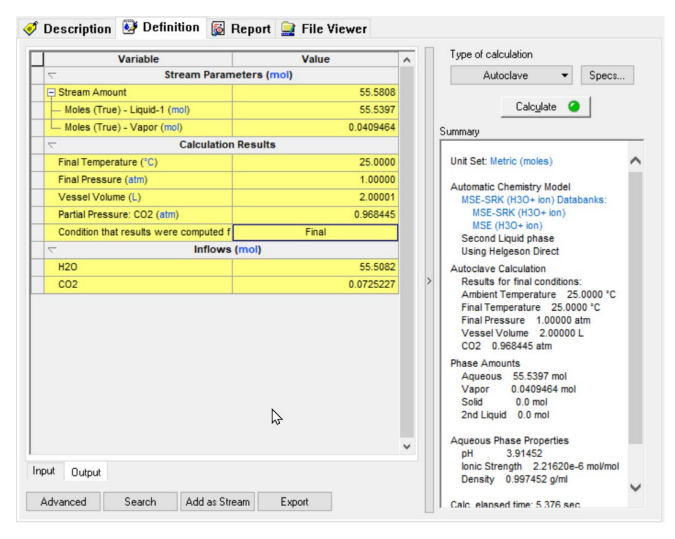
At this point the system is defined as 1 kg H₂O (55.508 moles) at an ambient temperature of 25 °C. The partial pressure of CO₂ will be calculated at a final temperature, 25 °C and pressure, 1 atm in a 2 L vessel.

We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Single Point Calculations*.

Analyzing the Results

Review the Summary Box or Click on the Output-Minitab to see the results.



A total amount of 0.0725 moles of CO_2 added was computed by the software to set the autoclave total pressure to 1 atm. For comparison, the same scenario in Example 12-1 Analyzing the Results required 0.04 moles of N_2 . The difference is the amount of CO_2 that dissolved in the water. CO_2 is more soluble than N_2 in water.

To analyze the CO₂ distribution in more detail, go to the **Report** Tab, and Scroll down until you find the **Element Distribution** table.

Element Distribution

| | Total | Total | Liquid-1 | Vapor |
|-------|-----------|-----------|------------|------------|
| | mol | mole % | % of Total | % of Total |
| H(+1) | 111.016 | 66.5797 | 99.9977 | 2.32772e-3 |
| O(-2) | 55.6533 | 33.3768 | 99.8552 | 0.144827 |
| C(+4) | 0.0725227 | 0.0434939 | 45.3215 | 54.6785 |

The **Element Distribution** table contains three rows, H(+), O(-2), and C(+4). These are the elements of H_2O and CO_2 . If you compare the C(+4) in the liquid and vapor phases, you will see that about 45.3% of the CO_2 dissolved in the water phase. By comparison, 1.5% of the added N_2 dissolved in water at the same condition (see Example 12-1).

Using Reactive Gas and NaOH solution

In this example, you will further modify the chemistry by adding NaOH so that additional CO₂ dissolves in water to form bicarbonate. This will increase the gas requirement.

Starting the Simulation

You are going to use the same case that you built up in the example above, with the only difference that you will add the **0.1 moles or NaOH.**

Copy the Autoclave – CO2 single point calculation and paste under the Autoclave Stream.

Name it Autoclave - CO2 - NaOH

Change the composition of NaOH to 0.1 moles and Calculate <F9>

| Variable | Value | |
|------------------------------------|----------|--|
| | | |
| Stream Amount (mol) | 55.6082 | |
| Ambient: Temperature (°C) | | |
| Ambient: Pressure (atm) | | |
| Calculation Parameters | | |
| Final Temperature (°C) | 25.0000 | |
| Final Pressure (atm) | 1.00000 | |
| Vessel Volume (L) | 2.00000 | |
| Partial Pressure: CO2 (atm) | | |
| Compute results at which condition | Final | |
| □ Inflows (mol) | | |
| H2O | 55.5082 | |
| CO2 | 0.0 | |
| NaOH | 0.100000 | |

At this point the system is defined as 1 kg H_2O (55.508 moles) and 0.1 moles of NaOH at an ambient temperature of 25 °C. The partial pressure of CO_2 will be calculated at a final temperature, 25 °C and final pressure, 1 atm in a 2 L vessel.

It is time to save your file (File >Save as...) or using the save icon in the tool bar.

Analyzing the Results

Review the Summary Box or Click on the Output-Minitab to see the results.

| ▽ Inflows (mol) | | |
|-----------------|----------|--|
| H2O | 55.5082 | |
| CO2 | 0.171600 | |
| NaOH | 0.100000 | |

The CO_2 inflow is calculated to be 0.1716 moles. This is exactly 0.1 moles more than the calculation with only CO_2 (Example 13-1). The difference results from the acid-base reaction with the 0.1 moles of NaOH.

To see the different species that have formed in solution go to the **Report** tab and select the **Species Output** (**True Species**).

Species Output (True Species)

Row Filter Applied: Only Non Zero Values column Filter Applied: Only Non Zero Values

| | Total | Liquid-1 | Vapor |
|------------------|-------------|-------------|------------|
| | mol | mol | mol |
| H2O | 55.5083 | 55.507 | 1.28481e-3 |
| Na+1 | 0.1 | 0.1 | |
| HCO3-1 | 0.0998917 | 0.0998917 | |
| CO2 | 0.0716555 | 0.0320887 | 0.0395668 |
| CO3-2 | 5.42505e-5 | 5.42505e-5 | |
| H3O+1 | 2.3994e-7 | 2.3994e-7 | |
| OH-1 | 6.87229e-8 | 6.87229e-8 | |
| NaOHCO3-2 | 2.76574e-14 | 2.76574e-14 | |
| NaOH | 3.90126e-15 | 3.90126e-15 | |
| Total (by phase) | 55.7799 | 55.7391 | 0.0408516 |

You can see that \sim 0.1 moles of bicarbonate (HCO_3^-) is formed. This reaction is increasing the CO₂ gas requirement to reach a total final pressure of 1 atm.

Custom

In the calculations that we have seen so far, variables were predefined. For instance, we selected either temperature or pressure as a fixed value in order to find the dew point pressure or dew point temperature. With the **Custom** calculation, however, you have the option to choose which parameters to set at fixed values and which parameters to keep as variables, to calculate the desired information, such as, for example, to investigate the solubility of gases in solution.

Example 14: Solubility of Oxygen in Water

With a custom calculation, we are going to calculate the solubility of O₂ in water at 25°C and 1 atm.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Custom Calculation | | | |
|----------------------|------------------------------------|-----------------------------------|---------------|
| Calculation Settings | | Stream Composition and Conditions | |
| Calculation Type | Single Point Calculation | Stream Amount | Default – 1kg |
| Calculation Sub-type | Custom | Temperature | 30 °C |
| Stream Name | Custom Calculation | Pressure | 4 atm |
| Name Style | Display Formula | H2O | Calculated |
| Unit Set | Metric, Mass fraction – ppm (mass) | O2 | 0 mol |
| Framework | MSE | | |

Setting the pH

Add a new Stream

Click on the new Stream and press <F2> to change the name to Custom Calculation

Select the MSE thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the **Units Manager** Icon, and select Metric, Batch, Mass Frac. This will change all inflows to mass %

Click on the Customize button. This opens the Edit Units window.

Under Basis Options change mass fraction from mass % to ppm (mass)

Enter O2 as an inflow and 0 moles

Change the T to 30 °C and P to 4 atm

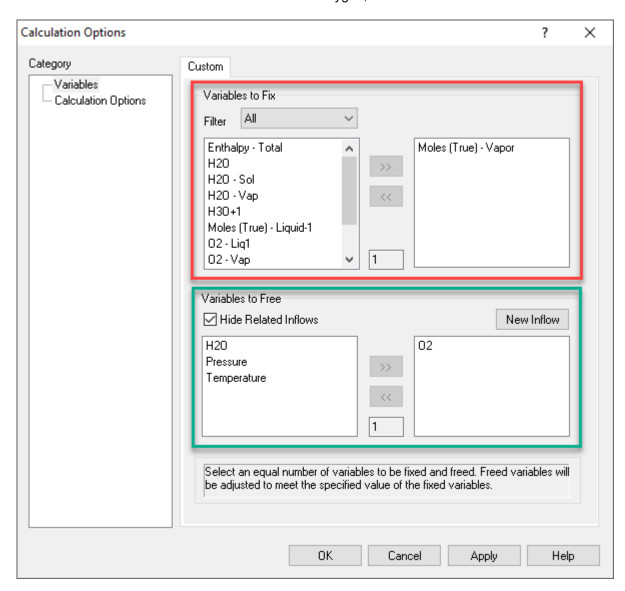
Go to the Add Calculation button and select Single Point calculation

Change the SinglePoint name to Oxygen Solubility using the <F2> key

Select Custom as Type of Calculation

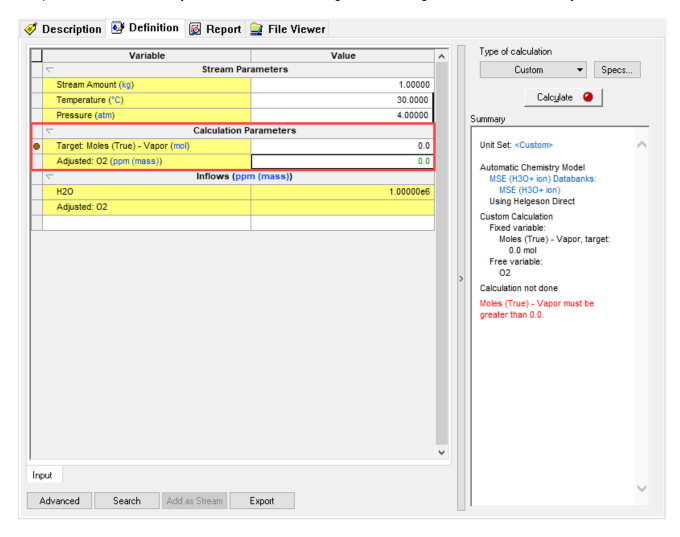
Select the Specs button. This will open the Calculation Options window

Note: In the Calculation Options window, you have the option to select the **Variables to Fix** and the **Variables to Free**. In this example, the Variable to Fix is the number of Moles (True) of Vapor which will be given a small value of 1×10^{-7} moles. You can interpret this as the first bubble of vapor formed, indicating that the aqueous phase is saturated. The Variable to Free is the inflow of oxygen, in order to achieve saturation.



In the **Variables to Fix** section, select **Moles (True) Vapor**, by double clicking or using the >> button. In the **Variables to Free** section, select **O2** as the inflow. Then click OK.

This will add a **Calculation Parameters** section, where the Target (Variable to Fix) and the Adjusted Inflow (Variable to Free) are shown. The Target has a brown dot in front of it indicating that the Target: Moles (True) - Vapor is fix value. The Adjusted Inflow value font is green indicating that the software will adjust this value.



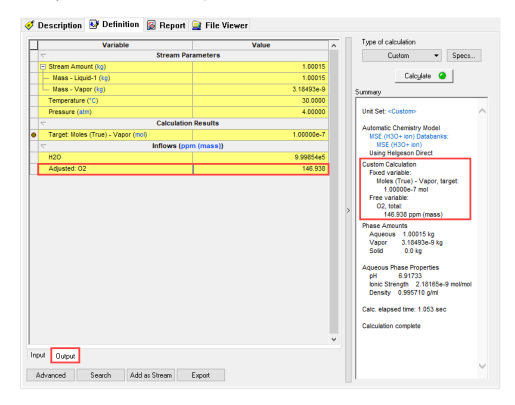
Fix the Moles (True) Vapor to $1e^{-7}$

We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Single Point Calculations*.

Analyzing the Results

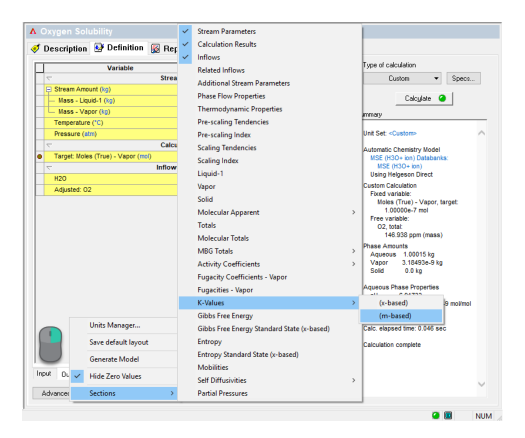
Review the Summary Box or Click on the Output-Minitab to see the results.



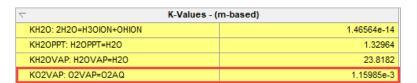
At 30°C and 4 atm the solubility of O₂ in water is 146.94 ppm (mass)

To get an estimate of the Henry's constant, you can study the K-values reported by the software.

Right click on the gray area, go to Sections>K-values>m-based.



The reciprocal of KO2VAP: O2VAP=O2AQ will give you an estimate of the Henry's constant.



Reaction Kinetics

Most of the chemistry and calculations that we have presented so far assume equilibrium. However, chemical reactions vary in speed as they occur. Some can occur instantaneously, while others can take years to reach equilibrium. The speed of a chemical reaction is given by the **Reaction Rate**, which is a measure of the change in concentration of the reactants or the change in concentration of the products per unit time.

For the stoichiometric reaction below:

$$aA + bB + \cdots \rightarrow cC + dD + \cdots$$

The reaction rate is defined as:

$$rate = -\frac{1}{a} \left(\frac{d[A]}{dt} \right) = -\frac{1}{b} \left(\frac{d[B]}{dt} \right) = \frac{1}{C} \left(\frac{d[C]}{dt} \right) = \frac{1}{d} \left(\frac{d[D]}{dt} \right)$$

We can describe the kinetics of a chemical reaction by using a **Rate Law**, which is an expression that relates the concentration of each reactant raised to an exponent that reflects the reaction order (which is determined experimentally) and the **rate constant**, k, (a proportionality constate between reaction rate and concentration).

The general rate law is generally expressed as:

$$Rate = k [A]^s [B]^t$$

Where, s + t gives us the reaction order of the reaction. s and t are not the same as the stoichiometric coefficients a and b.

The reaction **rate constant** units are specific for the overall reaction order, since we always want the units of the rate to be concentration units per unit of time, e.g., M/s.

| Reaction order | Units |
|----------------|---------|
| 0 | M/s |
| 1 | 1/s |
| 2 | 1/(M•s) |

The rate constant, k, is dependent on the temperature at which the reaction takes place. Its temperature dependence can be studied using the Arrhenius Equation, as shown below.

$$k = A \exp\left(-\frac{E_a}{RT}\right)$$

where: k= Reaction rate constant (units depend on reaction order)

A = Arrhenius frequency factor (has same units as k)

 E_a = Activation energy $\left(\frac{joule}{gmole}\right)$

 $R = \text{Universal gas constant} \left(8.314 \frac{joule}{amole K}\right)$

T = Temperature (K)

There are two different ways to enter rate laws reaction kinetics in OLI Studio: Stream Analyzer

Standard Rate Law which uses the Arrhenius equation for the reaction rate constant k, referred to as STD type within the software.

Non-Standard Rate Law, referred to as SPEC type within the software.

In this section, we will show you how to set up both types of reaction kinetics.

The reaction kinetics is enabled Under the Chemistry menu > Model Options > Phases > Kinetics

Standard Rate Law

The Standard Rate Law (STD) considers both the forward and reverse reaction rates, the individual species reaction orders, and the forward and reverse reaction constants (determined using the Arrhenius Equation).

For the reaction below:

$$aA + bB + \cdots \leftrightarrow cC + dD + \cdots$$

where: $a, b, \dots c, d$ are stoichiometric coefficients

 A, B, \dots are reactant species C, D, \dots are product species

The rate law is of the form:

$$Rate = (k_f a_A^{r_1} a_B^{r_2} ... - k_r a_C^{p_1} a_D^{p_2})$$

where:

Reaction rate $\frac{mol}{h}$ Rate = k_f = Forward reaction rate constant (units depend on reaction order) $k_r =$ Reverse reaction rate constant (units depend on reaction order) Activities of reactant species $\left(\frac{mol}{m^3}\right)$ a_A , a_B , ... = $r_1, r_2 \dots =$ Reaction order of individual reactant species (normally from experimental data. Default is stoichiometric coefficients: a, b, ...) Activities of product species $\left(\frac{mol}{m^3}\right)$ $a_C, a_D, \dots =$ Reaction order of individual product species $p_1, p_2, \dots =$ (normally from experimental data. Default is stoichiometric coefficients: c, d, \ldots)

Within the software the naming of the constants defined above is the following:

| OLI Keyword | Description |
|-------------|--|
| KF | Forward reaction rate constant |
| KR | Reverse reaction rate constant |
| AF | Forward reaction Arrhenius factor |
| AR | Reverse reaction Arrhenius factor |
| BF | Forward reaction activation energy divided by the universal gas constant (K) |
| BR | Reverse reaction activation energy divided by the universal gas constant (K) |
| ER_i | Reaction order of reactant species i |
| EP_i | Reaction order of product species i |
| | |

Thus, when specifying a standard rate expression, the user must define one of the following: (1) The Arrhenius frequency factors (AF and AR), and reaction activation energies divided by the universal gas constant (BF and BR) or, (2) the reaction rate constants (KF and KR) directly. In addition, the user can specify the individual

species order coefficients for the forward and reverse reactions if these differ from the stoichiometric coefficients (which are the default).

When defining the reaction order for a species, the order in which the species appears in the reaction equation must be defined (i.e., subscript *i*) with a sequential number, for either the reactant or product species. Hence, the first reactant is identified with the number 1, the second, 2 and so on. Similarly, the product species are identified with the integers 1, 2, 3, etc.

If any of the keywords are not defined, the software assumes a default value for that variable. These default values are assumed to be zero for the reaction rate constants, For the species reaction order coefficients, the reaction stoichiometric values are assumed. To complete the standard rate expression definition, the reaction temperature and initial reactant molality are included in the process stream composition definition.

Example 15: Reaction of Ammonia and Carbon Dioxide

In this example, we are going to consider the reaction of ammonia and carbon dioxide to form urea and water, according to the following reaction:

$$2NH_{3(aa)} + CO_{2(aa)} \leftrightarrow NH_2CONH_{2(aa)} + H_2O$$

This reaction will take place in a plug flow reactor at 200°C and 100 atm, with a total residence time of 100 hours.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Single Point Calculation with Reaction Kinetics Enabled | | | |
|---|--------------------------|-----------------------------------|------------|
| Calculation Settings | | Stream Composition and Conditions | |
| Calculation Type | Single Point Calculation | Stream Amount | Calculated |
| Calculation Sub-type | Isothermal | Temperature | 200 °C |
| Stream Name | Standard Rxn Kinetics | Pressure | 100 atm |
| Single Point Name | Rxn Ammonia/CO2 | Water | 55.5082 |
| Name Style | Display OLI Name (TAG) | CO2 | 2 moles |
| Unit Set | Metric, Moles | NH3 | 2 moles |
| Framework | MSE | UREA | 0 moles |
| Kinetics | Enabled | | |
| Kinetics Holdup Time | 100 hours | | |
| Number of Kinetic Steps | 10 | | |

Add a new Stream

Click on the new Stream and press **<F2>** to change the name to *Standard Rxn Kinetics* Select the **MSE** thermodynamic Framework

Click on the Names Manager Icon, and select the OLI Name (TAG) option, and click OK.



Click on the Units Manager Icon, and select Metric, Batch, Moles.

Enter the stream composition, temperature and pressure given in the table above.

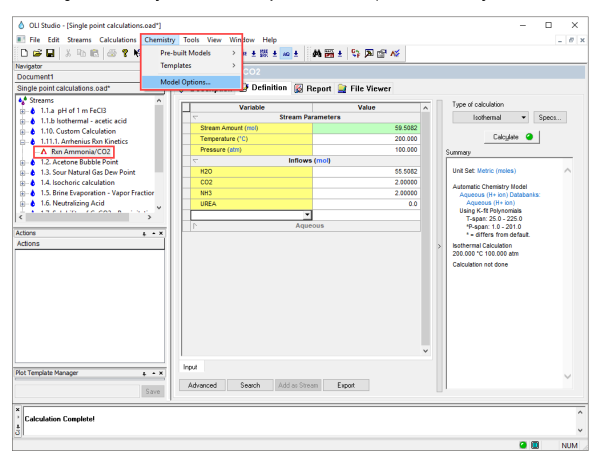
Go to the Add Calculation button and select Single Point calculation

Select Isothermal (default) as Type of Calculation.

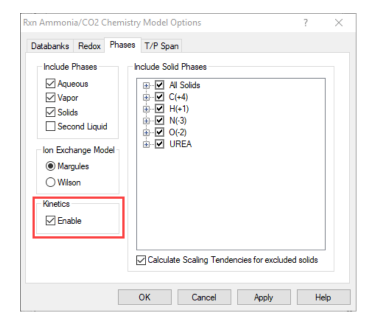
Change the SinglePoint name to Rxn Ammonia/CO2 using the <F2> key

So far, we have created an Isothermal calculation. Now, we are going to enable the **Reaction Kinetics** option at the *Single Point Level*.

To enable reaction kinetics, make sure that you are at the *Single Point Level* by clicking on the Single Point Icon, and then go to **Chemistry menu > Model Options**. This will open the **Chemistry Model** window.

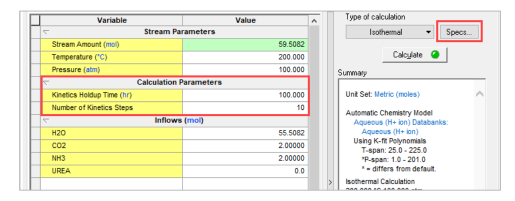


Select the Phases tab and check the Kinetics box. Then click OK.



A new grid section named **Calculation Parameters** appears. In this grid the **Kinetics Holdup Time**, and the **Number of Kinetic Steps** can be defined. The default values are 2 h and 2 steps respectively.

Change the Kinetics Holdup Time to 100 hours and the Number of Kinetic Steps to 10.



Kinetics Holdup time is equivalent to Residence Time. The Number of Kinetic Steps is equivalent to Δt . Thus, Kinetics Holdup time = $\Sigma \Delta t$.

Note: Kinetics Holdup Time vs Number of Kinetic Steps

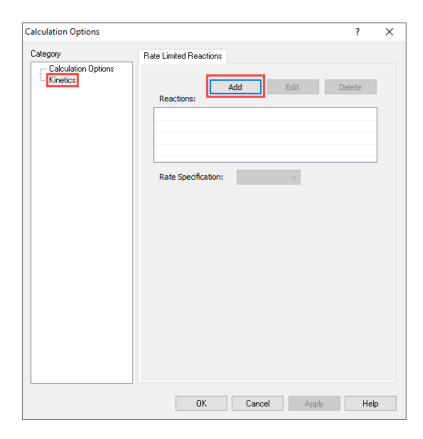
For processes that have reaction kinetics, the software needs to know how the concentration of the reactants (or products) change over time during the reaction, which is given by the **rate law**. It is necessary to divide the total residence time into small steps. Adding more steps decreases the error; however, as you increase the number of stages, the computational time increases.

Now, our next step is to define the reaction kinetics.

Click on the **Specs** button. This will open the **Survey Options Window**.

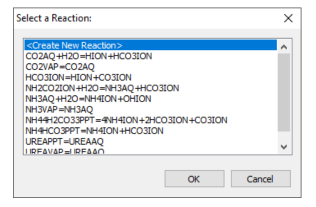
Select Kinetics under the Category window

Click the Add button to add a new reaction. This will open the Select a Reaction Window.



You will be given a list of reactions which are already in the chemistry model. For our example, we need to create a new reaction.

Select the option Create a New Reaction, and then click OK.



This enables the option to enter a new reaction.

Type the following reaction: 2NH3AQ+CO2AQ=UREAAQ+H2O and then hit Enter. The window will update.

Note: You must use the OLI Tag Name for this step, and additionally specify the phase of the reactants and products. Water is a special case; it is written only as H2O.

As a general rule:

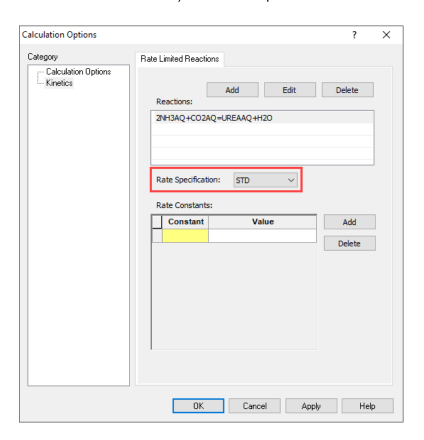
For an aqueous phase: AQ For a vapor/gas phase: VAP For a solid phase: PPT

For a hydrated solid: SOLIDNAME.nH2O, where n is the hydration number

For an ion: ION

You can find the OLI TAG Name of your specific species using the Component Search Tool.

Select the STD (Standard Rate Reaction Kinetics) as the Rate Specification



Now, let's start entering the rate constants.

Remember, the forward and reverse reactions have the same format. The rate is the following:

$$R = k_f [NH_3]^2 [CO_2] - kr [NH_2CONH_2]$$

For this example, the forward rate constant (KF) is a constant value equal to 2000.

And, the reverse rate constant (KR) is defined as follows:

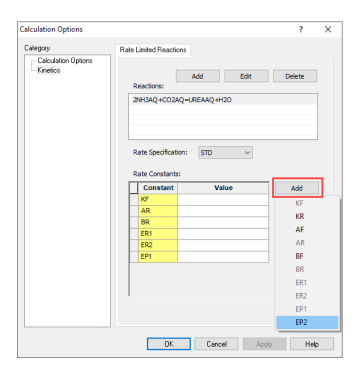
$$KR = AR e^{\frac{-BR}{T}}$$
 $KR = 1.2 \times 10^{-10} e^{\frac{-3480.78}{T}}$

A summary of the Reaction Rate Parameters is given in the table below (These parameters where created for illustration purposes, do not use them for any real design work).

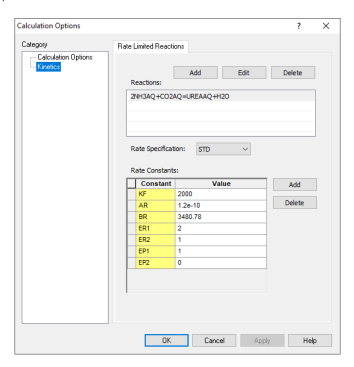
| Parameters | Value | Comment |
|------------|--|--|
| KF | 2000 | Forward Rate Constant |
| KR | Calculated | Reverse Rate Constant |
| AF | | Not needed |
| AR | | 1.2×10^{-10} |
| BF | | Not needed |
| BR | 3480.78 | Determined from $BR = \frac{E_a}{RT} = \frac{28939.9}{8.3142}$ |
| ER1 | 2 | This is the exponent of reactant 1 [NH3] ² |
| ER2 | 1 | This is the exponent of reactant 2 [CO2] |
| EP1 1 | This is the exponent of product 1 in the | |
| | ı | mechanism. In this case [NH2CONH2] |
| EP2 | 0 | H2O does not participate in the reaction |

Now that we have identified the parameters needed, we can add them.

Click on the Add button and select the parameters needed.



Type the constant values, and then click **OK**.



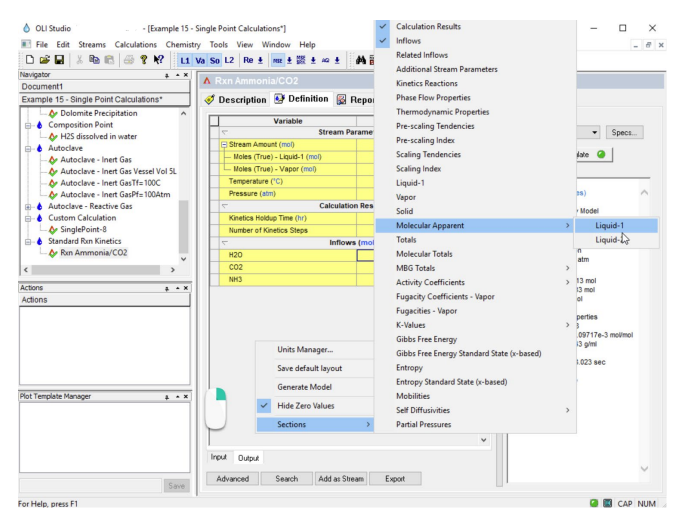
We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key.

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Single Point Calculations*.

Analyzing the Results

Click on the Output-Minitab to see the Results.

Right click on the gray area and select Sections> Molecular Apparent - Liquid-1



About **0.00062 moles** of Urea had been formed with a total residence time of 100 hours.



You can make changes on the residence time, reaction conditions or the rate of reaction to maximize the production of urea.

Non-standard Rate Law

Non-Standard Rate Law, also referred to as SPEC type within the software. These reactions do not follow the Arrhenius kinetics.

Example 16: Ammonia Hydrolysis

In this example we are using simpler chemistry. We are going to study the hydrolysis of ammonia.

$$NH_{3(aq)} + H_2O \rightleftharpoons NH_4^+ + OH^-$$

For this reaction we know the forward rate constant (k_f) but need to constrain the reverse reaction to the thermodynamic equilibrium constant. This implies that we are going to replace an existing equation within the software.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Single Point Calculation with Reaction Kinetics Enabled | | | | | |
|---|---------------------------|-----------------------------------|------------|--|--|
| Calculation Settings | | Stream Composition and Conditions | | | |
| Calculation Type | Single Point Calculation | Stream Amount | Calculated | | |
| Calculation Sub-type | Isothermal | Temperature | 25 °C | | |
| Stream Name | Non-Standard Rxn Kinetics | Pressure | 1 atm | | |
| Single Point Name | Ammonia Hydrolysis | Water | 55.5082 | | |
| Name Style | Display OLI Name (TAG) | NH3 | 0.1 moles | | |
| Unit Set | Metric, Moles | | | | |
| Framework | MSE | | | | |
| Kinetics | Enabled | | | | |

Add a new Stream

Click on the new Stream and press <F2> to change the name to Non-Standard Rxn Kinetics

Select the MSE thermodynamic Framework

Click on the Names Manager Icon, and select the OLI Name (TAG) option, and click OK.

Click on the Units Manager Icon, and select Metric, Batch, Moles.

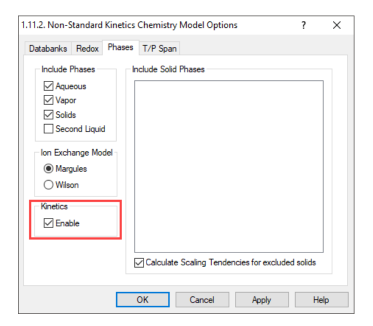
Enter the stream composition, temperature and pressure given in the table above.

Go to the Add Calculation button and select Single Point calculation

Change the SinglePoint name to Ammonia Hydrolysis using the <F2> key

Select **Isothermal** (default) as Type of Calculation.

To enable reaction kinetics, make sure that you are at the *Single Point Level* by clicking on the Single Point Icon, and then go to **Chemistry menu > Model Options**. This will open the **Chemistry Model** window. Select the **Phases** tab and check the **Kinetics** box. Then click **OK**.



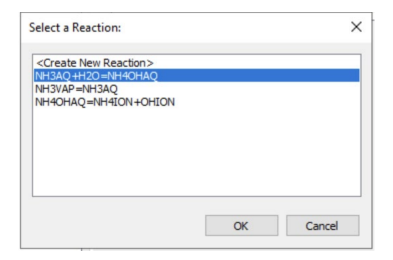
Now, our next step is to define the reaction kinetics.

Click on the Specs button. This will open the Options Window.

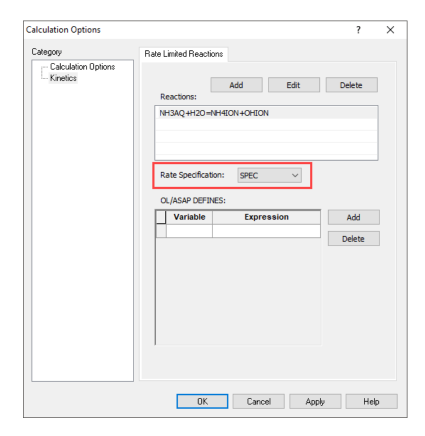
Select Kinetics under the Category window

Click the Add button to add a new reaction. This will open the Select a Reaction Window.

Select the first reaction: NH3AQ+H2O=NH4OHAQ and then click OK.



Select the SPEC as the Rate Specification.



For this example, the forward and reverse reaction rates are the following:

$$Rate_f = k_f \cdot \gamma_{NH_{3,aq}}[NH_{3,aq}] \cdot \gamma_{H_2O}[H_2O]$$
 where $k_f = 3$

$$Rate_r = k_r \cdot \gamma_{NH_4^+}[NH_4^+] \cdot \gamma_{OH^-}[OH^-]$$
 where $k_r = \frac{k_f}{K_{eq}}$

Thus, the total rate is given by:

$$Rate = Rate_f - Rate_r$$

$$Rate = k_f \cdot \gamma_{NH_3,aq} \big[NH_{3,aq} \big] \cdot \gamma_{H_2O} [H_2O] - k_r \cdot \gamma_{NH_4^+} [NH_4^+] \cdot \gamma_{OH^-} [OH^-]$$

This is where things get difficult. You now need to speak "OLI". We have some rules for SPEC type equations.

We now need to turn these values into "OLI" terms 6.

⁶ Commonly referred to as ASAP variables.

We can define any variable we want. We have some variables that you will need to use. Any concentration variable such as $[NH_{3aq}]$ is defined as the natural log and is designated with the letter "L". So $[NH_{3aq}]$ is used as LNH3AQ.

Similarly, activity coefficients γ are also taken as the natural log. So, γ_{OH^-} is written as Log_e γ_{OH^-} = AOHION.

KEQ is the thermodynamic equilibrium constant for the equation. AH2O is special in the Aqueous thermodynamic framework in that it is the variable $Ln(a_{H_2O})$.

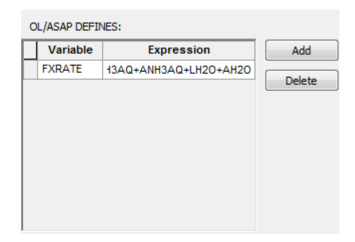
We now need to add these variables to the kinetics window 7:

| Variable | Expression |
|----------|---|
| FXRATE | LNH3AQ+ANH3AQ+LH2O+AH2O |
| RXRATE | LNH4ION+ANH4ION+LOHION+AOHION |
| KF1 | 3 |
| KR1 | KF1/KEQ |
| RATE1 | (KF1*EXP(FXRATE)-KR1*EXP(RXRATE))*VOLLIQ/1000 |

Note: For a reaction rate to be considered in the program, your set of variables should include a RATEn statement where the "n" is the reaction rate equation number.

Note: The variable VOLLIQ is the volume of the liquid phase in Liters. OLI requires the rate to be in mol/m³ we need to divide by 1000.

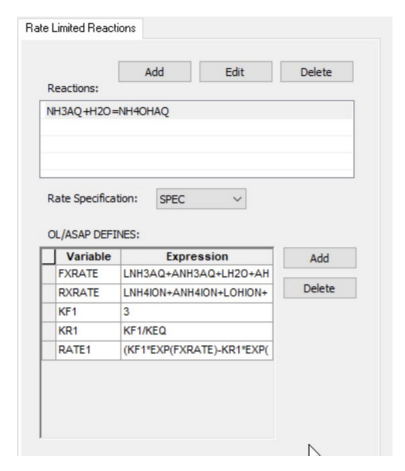
Change the Kinetics Holdup Time to 100 hours and the Number of Kinetic Steps to 10.



⁷ Copy and paste from this document can make this step easier

You will notice in the image above that we have scrolled to the end. That is ok. Click the <u>Add</u> button to continue with the equations. It is important that you keep the order correct. You see what we are using previously defined variables in subsequent equations.

Complete the remainder of the variables.



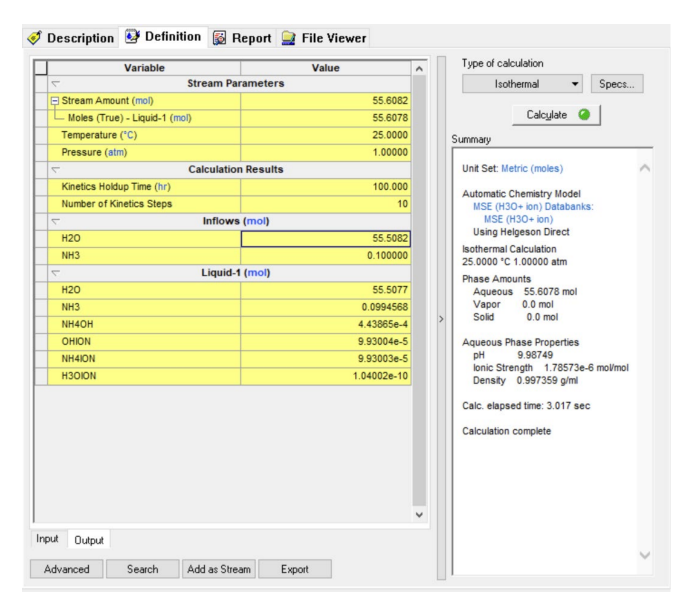
The completed equations

Click the **OK** button.

Like the previous example, this is a plug-flow reactor with a holdup time of 100 hours and 10 steps.

Now **Calculate**

Once the calculation is complete you can click the **Output** mini tab and add the <u>Liquid-1</u> section (not molecular aqueous):



You can see that some ammonium ions have been created. It is interesting to see what the equilibrium condition would be.

Create another single point isothermal calculation and run it without any reaction kinetics. Here are the results below:

| Variable | Value | | | |
|-------------------------------|-------------|--|--|--|
| | | | | |
| Stream Amount (mol) | 55.6082 | | | |
| Moles (True) - Liquid-1 (mol) | 55.5287 | | | |
| Temperature (°C) | 25.0000 | | | |
| Pressure (atm) | 1.00000 | | | |
| □ Inflows (r | nol) | | | |
| H2O | 55.5082 | | | |
| NH3 | 0.100000 | | | |
| | nol) | | | |
| H2O | 55.4274 | | | |
| NH4OH | 0.0795157 | | | |
| NH3 | 0.0191164 | | | |
| OHION | 1.36788e-3 | | | |
| NH4ION | 1.36788e-3 | | | |
| H3OION | 7.99036e-12 | | | |

The reaction kinetics have forced the back reaction to be dominant. In the equilibrium case, approximately

 $^{1.4}x10^{-3}$ moles of NH₄⁺ have been created. By limiting the forward reactions, we allow the back reaction to take place and that only formed ~1.0x10⁻⁴ moles of NH₄⁺.

Section 2. Survey Calculations

Survey calculations are single point calculations strung together in series. They are also referred to as multiple point calculations.

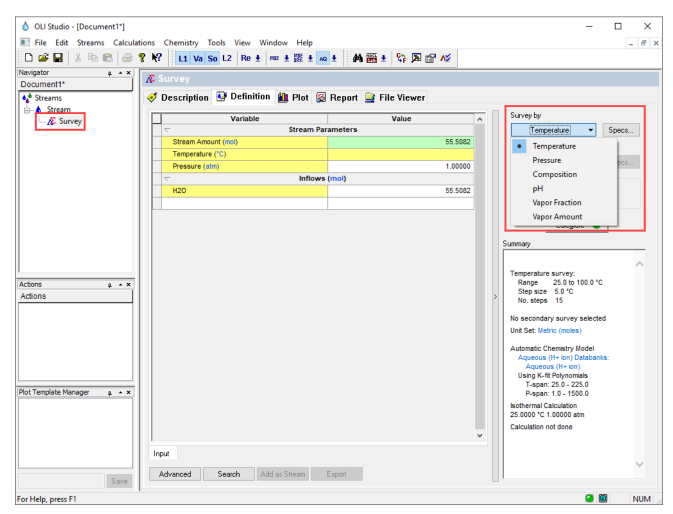
There are three different ways to set up surveys:

<u>Single Survey:</u> These calculations allow the user to designate one independent variable. The predefined single surveys include surveys by Temperature, Pressure, Composition, pH, Vapor Fraction and Vapor Amount.

Dual Survey: These calculations allow the user to designate two independent variables.

<u>Survey by changing the single point calculation type:</u> By default, the isothermal calculation is selected for survey calculations. If the user wants to study the solubility of a salt in solution, it is possible to change the survey from **Isothermal** to another calculation type like **Precipitation Point**.

In this section, you will learn how to set up **survey types** and introduce some additional features to get the most out of your simulation results.



Single Surveys

An introduction to single surveys is given here. We will illustrate, in detail, the steps necessary to conduct this type of calculation. The example below was designed to be used as a guide for future reference.

Example 17: Removal of Nickel from Wastewater

This case study is a typical wastewater treatment problem, the removal of a trace heavy metal ion (nickel) from a water stream in which the presence of another chemical (cyanide) significantly alters the treatment strategy. In this case, we are considering precipitation as an approach to removal of the nickel.

The wastewater in this case study contains nickel ions at a concentration of 0.002 m (or moles/ 1 kg H_2O). The existing treatment strategy is to precipitate the nickel ions as Nickel Hydroxide (Ni(OH)₂). The soluble nickel remaining after precipitation needs to be less than 1 ppm, which is the maximum contaminant level allowed.

During the course of the plant operation, some cyanide ion is inadvertently added to the waste stream. The soluble nickel is now more than 1ppm. Sulfide salts were then added to hopefully precipitate the nickel and once again achieve the design specification.

This example will be divided into three different scenarios:

Scenario 1: Wastewater without Additives
Scenario 2: Wastewater with Cyanide Content
Scenario 3: Wastewater with Sulfide Salts

The power of the Stream Analyzer becomes apparent when we are seeking to study the chemistry of individual streams. For this specific example a Survey by **pH** will be used. We will also be using the **AQ** thermodynamic model instead of the default MSE model.

Scenario 1: Wastewater without Additives

We begin by starting the Stream Analyzer Program. This may be accomplished by *clicking* on the OLI Studio icon or by using the Start button and finding OLI Studio under *Programs*, which will take you to the New window where you start creating your calculations.

Select the AQ-Databank (

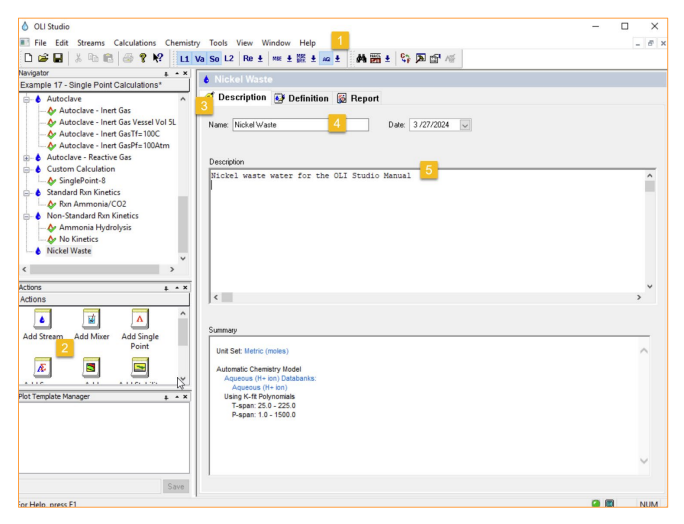
Click on **Add Stream** icon (in the actions panel). This will display the **Definition** window. We should add some descriptive information about this stream, so we can later identify the stream

Click on the **Description** tab. This will display the description information

Replace the name *Stream* with the name *Nickel Waste*. It is advisable to change the name of the stream from the default name. You may be entering many streams and will need to sort them out later

Add the following text to the Description box: "Nickel wastewater for the OLI Aqueous Modeling Course"

Note: The summary box will contain additional information as the calculations proceed. This information may be the name of additional databases or chemistry models imported from other OLI software packages. The filled-out window should look like the figure below.



Click on the **Definition** tab to start defining the wastewater stream

Click in the white box in the grid below the Inflows line. Add the formula Ni(OH)2

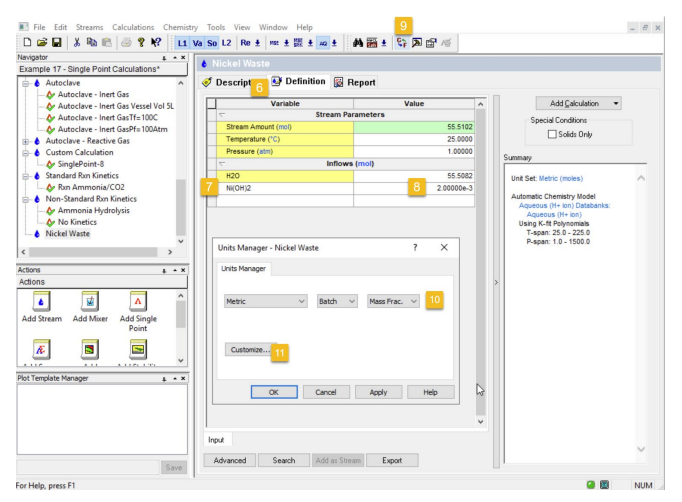
Click in the white box next to the species you just entered and enter the value **0.002**. Press **<Enter>** to update the list

Note: Stream Analyzer will automatically change the name of the species to the selected display name. In this instance, the name you entered was **Ni(OH)**₂ but it may have changed to a different form. If it changed you can specify which display name to use in the Names Manager icon

For the output results, we want output units in **ppm**. To change the output units, click on the Units Manager icon ^{\$\frac{1}{2}\$}. This will open a new window.

Change moles to Mass Frac. Using the drop-down arrow

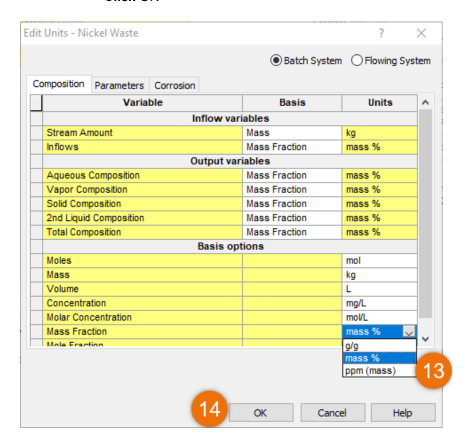
Click on the **Customize** button. The **Units Manager** window will appear.



Make sure you are on the Composition tab

Change the Mass Fraction Units to ppm (mass)

Click OK



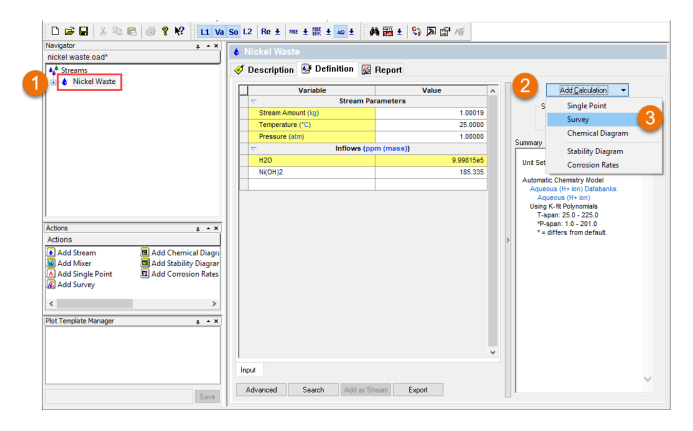
Our primary interest in this application is finding the optimum pH for nickel removal. To create a plot of Ni⁺² as function of pH, we will need to run a pH survey.

Creating a pH Survey to find the optimum pH for nickel removal

Click on the **Nickel Waste** stream icon in the tree view on the left-hand side of the window. This will bring you back to the top of the series of calculations by displaying just the stream information.

Click on the Add Calculation button

Select Survey



You can now add descriptive information about this calculation.

Enter a new Survey name: Base Survey. You can also double click on

'Survey' to change the name

Enter a **Description**: Base pH survey without additives.

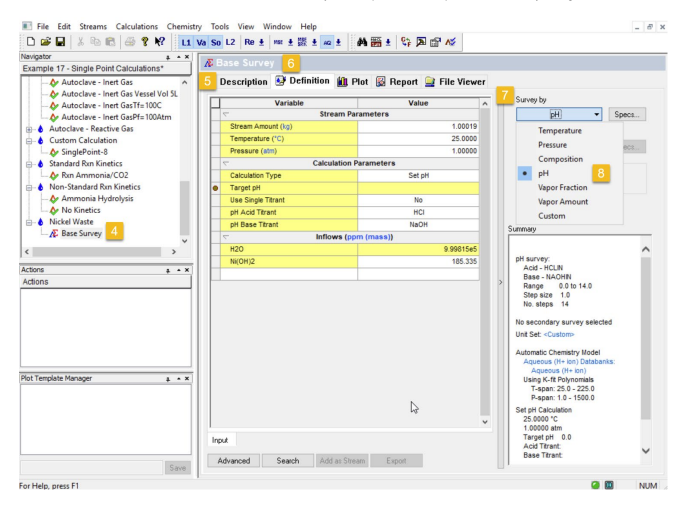
Since we do not want a temperature survey which is the default, we will need to change the survey type.

Click on the **Definition** tab to do the survey calculation.

Click on the Survey by button

Select pH

Note: The default acid titrant and the base titrant are already defined (HCl and NaOH). We are now ready to begin the calculations.



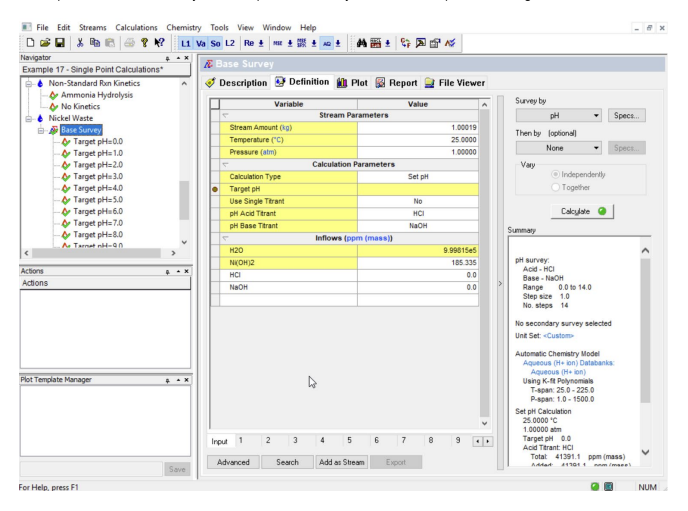
We should now save our work. It is very frustrating to work for a long period of time and forget to save our work. So please save.

Select File

Select **Save As** from the menu. Give the name *Removing nickel from* wastewater⁸

Click on the Calculate button

The program will run for a short time. When the orbit disappears, check the summary box to see if the calculation is complete. In the tree-view, you can expand the survey to see if all the points converged.

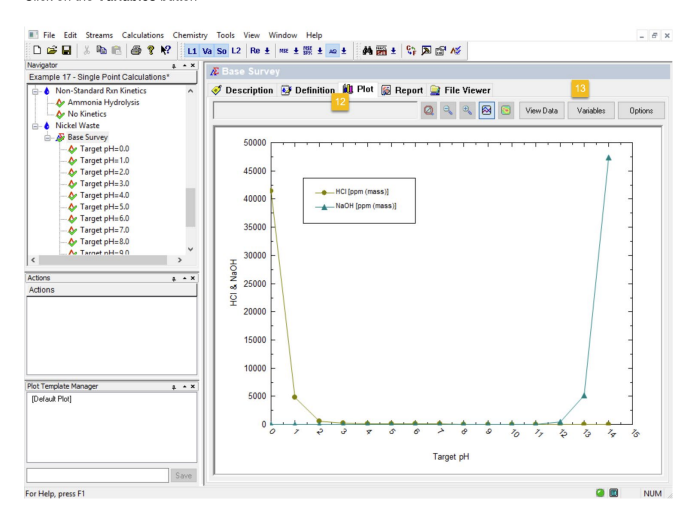


⁸ This may also be referred to as Example 17

We can now obtain some graphical results.

Click on the Plot tab

Click on the Variables button

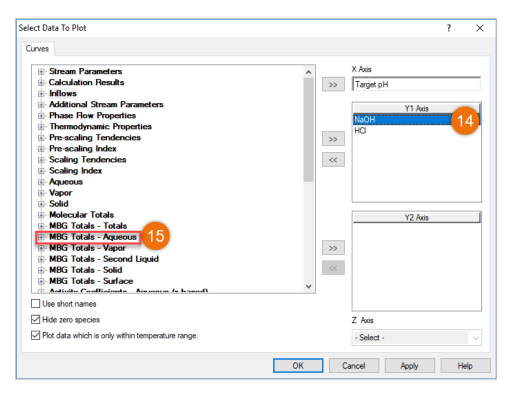


This will open a new window. The NaOH and HCI variables in the Y-Axis box should be displayed.

Select them and then **Click** on the left double-arrow (<<) button which will remove **NaOH** and **HCI** from the list. You can also select and double click to remove them.

Scroll down the left-hand window to find **MBG Totals - Aqueous** and expand the list by clicking the \blacksquare box.

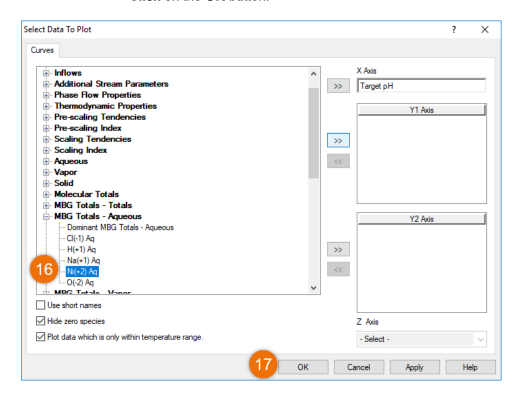
Note: MGB is an abbreviation for Material Balance Groups. The **MGB** variable is a sum of all the species for that material in the phase requested.



The grid updates to show the **MGB** totals available to display. In this case we desire the Nickel(+2) species. in this case all the **Ni(+2) Aq** variable is a sum of all nickel containing ions in the aqueous phase in solution. Any solids are excluded from the summation.

Double-Click the Ni(+2) item or select it and use the >> button.

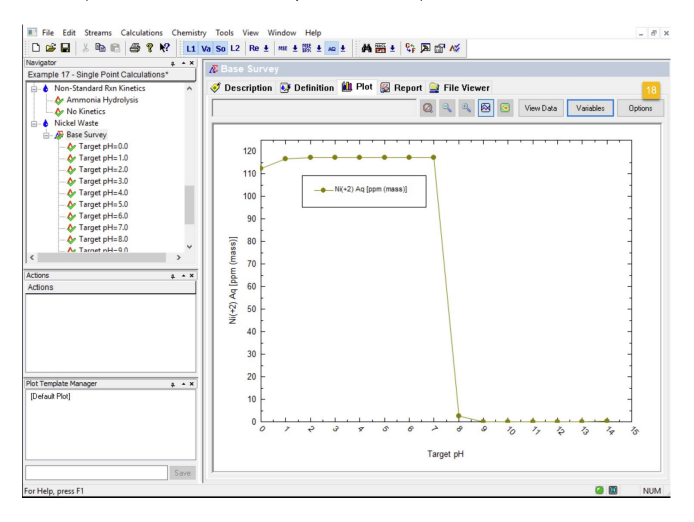
Click on the OK button.



You will see the new plot.

For many calculations, the values on the plot extend over a very large range of numbers. The default linear axis may not capture all the details we require.

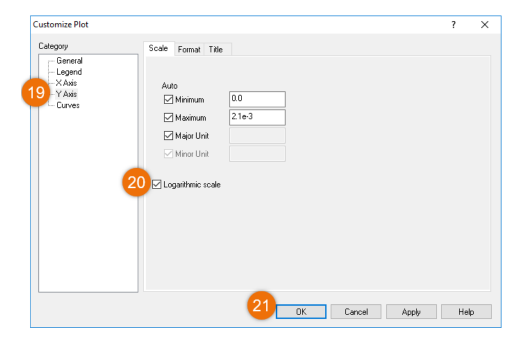
Above the plot window locate and select the **Options** button. This opens a new window.



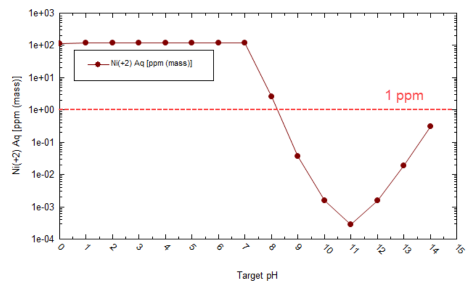
Select Y-Axis from the Category List

Check the Logarithmic Scale Box

Click on the OK box



The modified plot is then displayed. The limit of 1 ppm for Ni⁺² is approximately 2 x 10⁻⁵ moles. Above pH=9, we are several orders of magnitude below this limit. Additionally, you can see that a minimum amount of Ni²⁺ in the aqueous phase seems to occur in the pH=11 range. This is the result of nickel solids forming and leaving the aqueous phase.

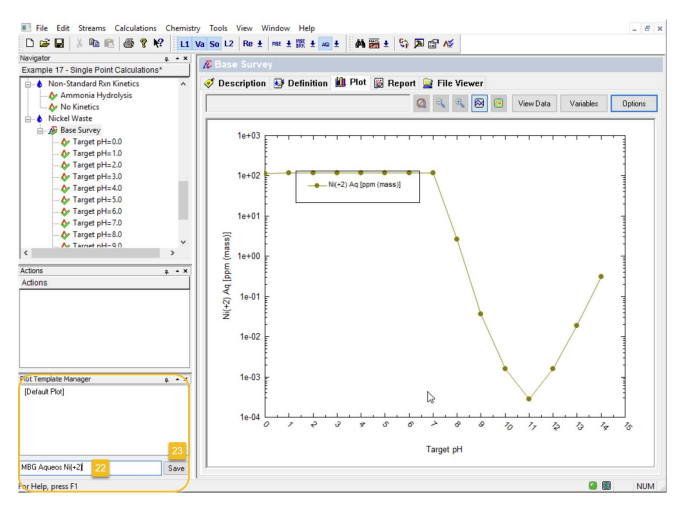


Note: The red line was put as a reference. it was not plot with the software.

Let's make the plot shown above the default plot. To do that we are going to use the Plot Template Manager Tool.

Save the description in the left-hand corner panel named **Plot Template Manager**. Type **MBG Aqueous Ni(+2)** as the name in this case.

Click Save



Now there will be two plots on that list now. The first is the default plot, and the second is **MBG Aqueous Ni(+2)**. You are now able to see these plot conditions for any other analyzer object in this case when you add survey for that object.

What else is important in this solution?

Click once more on the Variables button

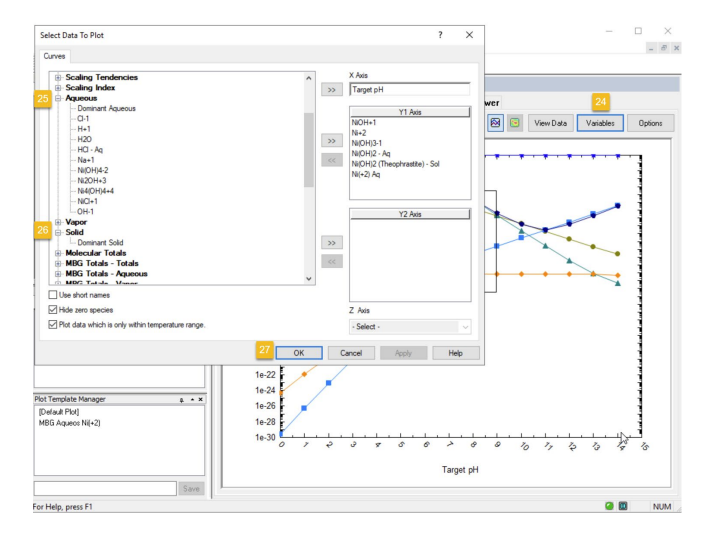
Add the following Aqueous species to the plot. (You may need to scroll up or down to find all the species):

Ni(OH)2 NiOH+1 Ni+2 Ni(OH)3-1

Add the following Solids species to the plot:

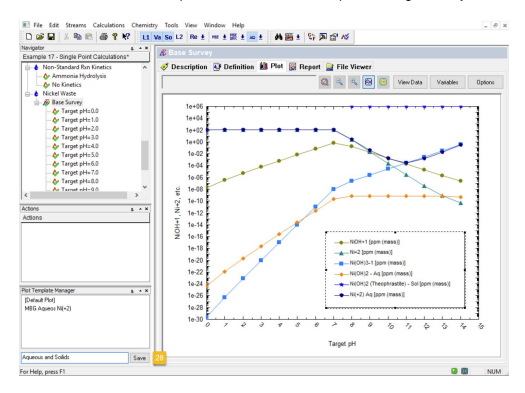
Ni(OH)2

Click on the OK button when done



You can see that the soluble nickel (**Ni(+2) Aq**) is a summation of the other species. The large drop in the value is because most of the nickel leaves the aqueous solution as Ni(OH)2-Solid at pH values greater than 7.0 with a maximum near pH=11.

Let's save this as a default plot as well in the Plot Template Manager as Aqueous and Solids



Scenario 2: Wastewater with Cyanide Content

The real importance of aqueous speciation modeling of this treatment is only really appreciated if we introduce cyanides, which brings us to the real waste treatment problem.

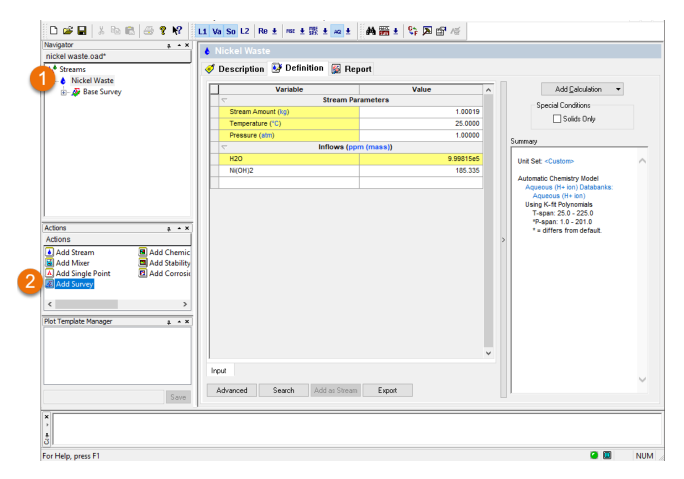
For this scenario, you will repeat many of the same steps as in <u>Scenario 1</u>. We recommend that you create new calculations below the **Nickel Waste** stream. This will keep the core composition the same without affecting the results of other calculations.

Please follow these steps for this scenario. Please note that we will only show the screens that are substantially different from those that you have already seen.

Creating a pH Survey to find the optimum pH for nickel removal

Click on the Nickel Waste stream in the tree view in the left-hand window

This will display the **Actions pane** in the bottom left corner of the Stream Analyzer window. **Click** on the **Add Survey** icon in the Actions pane.



Click on the Description Tab, and change the Name and Description in the Description tab.

Name: Waste Survey with CN

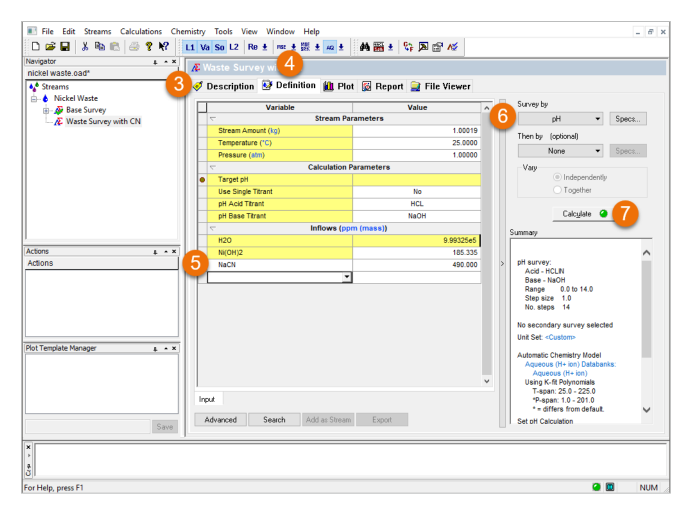
Description: pH survey with both Nickel and CN

Click on the Definition Tab

Add NaCN to the grid with a value of 490 ppm

Click on the Survey By button and select pH. Note that HCl and NaOH are the default titrants and are automatically added.

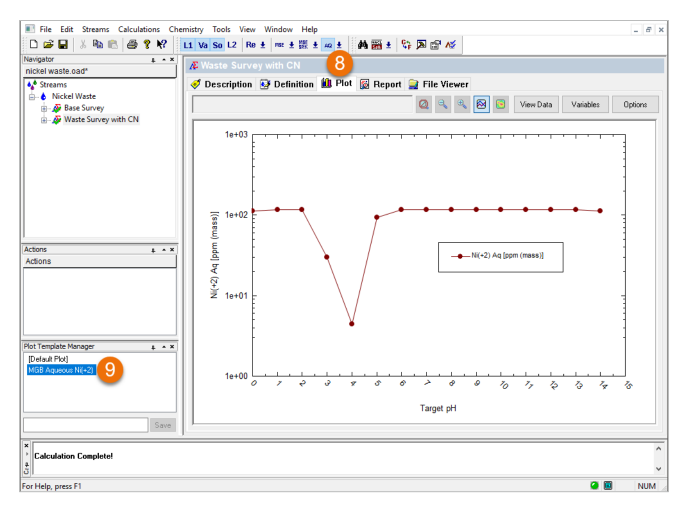
Click on the Calculate button.



Let's review the results.

Click on the Plot tab

Go to the Plot Template Manager tool, and click on MGB Aqueous Ni(+2)



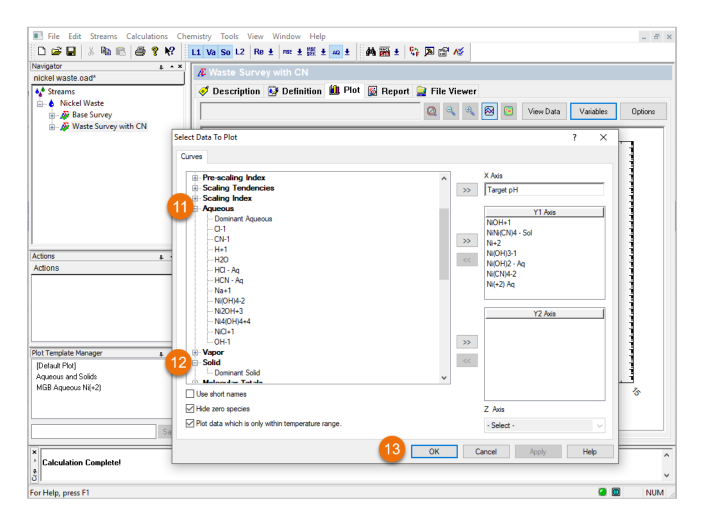
The results have changed very dramatically. The new optimum pH for Ni removal is around 4.0, rather than 11. However, the lowest total Ni remaining in solution is now around 4 ppm which is well over 1 ppm.

Let's analyze why this is happening. We can now modify the plot to display more variables. Select the **Aqueous** and **Solids** plot saved in the **Plot Template Manager**.

Click on Variables

Add the following new Aqueous species: Ni(CN)4-2
Add the following new Solid species: NiNi(CN)4-Solid

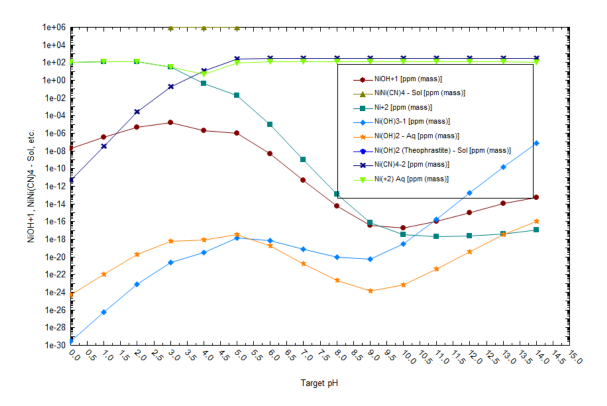
Click OK



Let's analyze the results.

The culprit is the Ni(CN)₄-2 complex of nickel and cyanide. Basically, the plot of the total Ni in solution and the Ni(CN)₄-2 complex overlap over the interval pH=5 to 12. This means that virtually all nickel in solution is in the form of this complex.

This complex thus holds the Ni in solution and does not allow the nickel hydroxide to even form. Instead, a much weaker precipitate, the NiNi(CN)₄ salt forms over a narrow range of pH with 4.0 being the optimum.



Scenario 3: Wastewater with Sulfide Salts

We can now try to influence nature by introducing a source of sulfide. We do this because many metal sulfide salts are highly insoluble.

Create a new single point calculation and a survey as you did in the previous two scenarios.

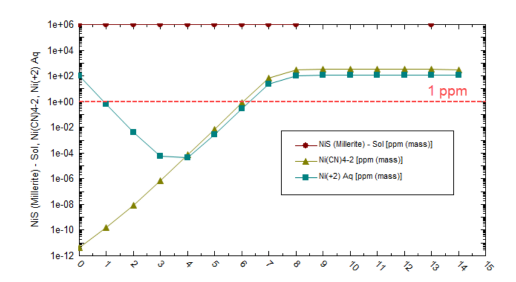
Add 490 ppm of NaCN

Add 340 ppm of H2S

Select the pH survey and then calculate.

For the survey case, use the plot template manager to start off with the MBG– Aqueous (Ni+2) and then add the following variables:

Aqueous Species: Ni(CN)4-2 Solid Species: NiS-Solid



The results reflect a "power struggle" between the Ni(CN)-2 which is holding the nickel in solution and the NiS solid which clearly has a greater tendency to form than the NiNi(CN)4 solid. As a result, our optimum pH is still around 4.0 and we are now around 10-5 ppm total nickel in solution which is a bit below 1 ppm.

This would be a good time to **save** your work. You may use the **File/Save As...** menu item or use the **Save icon** on the toolbar.

Final Thoughts...

Aqueous speciation modeling can teach us a great deal about complex chemical systems and the interactions of individual species.

The actual removal achieved with sulfide may not be quite enough to satisfy the regulations. This is useful information to have. In addition, with the power of OLI Studio: Stream Analyzer, one could now explore alternative treatment methods such as ion exchange.

Although such a simulation is beyond the scope of this demonstration, consider how vital it is to know that the dominant species to be exchanged (removed from solution) is an anion Ni(CN)⁻² and not the cation (Ni+2) as the conventional wisdom might dictate.

Temperature Survey

The default single survey calculation is the Survey by **Temperature.** The software computes solution properties based on a known composition and pressure at different temperatures.

Example 18: Studying a Four-phase mixture and its Partitioning

The purpose of this first example is to present the basic Temperature survey. You will study how a four-phase mixture partitions with temperature. The system composition will be basic, containing one primary phase component, H₂O (liquid), CH₄ (gas), Decane (organic), and NaCl (solid). Each component will partition each of the other three phases.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Temperature Survey Calculation | | | | | | |
|--------------------------------|----------------------|-----------------------------------|-----------------------------|--|--|--|
| Calculation Settings | | Stream Composition and Conditions | | | | |
| Calculation Type | Survey | Stream Amount | Calculated | | | |
| Stream Name | Temperature Survey | Temperature Range | 25-100 °C (Default) | | | |
| Survey Name | 4-Phase Partitioning | Step Size | Increment by 5 °C (Default) | | | |
| Name Style | Display Formula | Pressure | 1 atm | | | |
| Unit Set | Metric, Moles | H2O | 50 moles | | | |
| Framework | MSE-SRK | CH4 | 10 moles | | | |
| | | C10H22 | 10 moles | | | |
| | | NaCl | 10 moles | | | |

Add a new Stream

Click on the new Stream and press <F2> to change the name to Temperature Survey

Select the MSE-SRK thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Moles

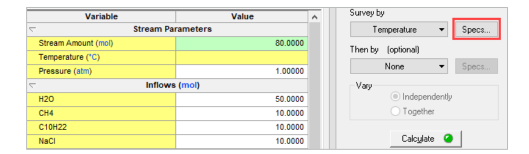
Enter the composition of the stream given in the table above

Go to the Add Calculation button and select Survey calculation

Change the Survey name to 4-Phase Partitioning using the <F2> key

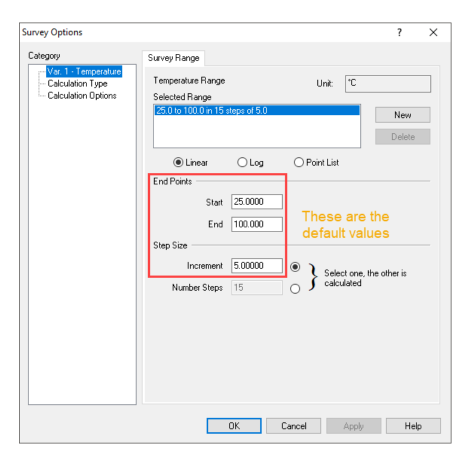
Select **Temperature** as Type of Survey - Default

Click on the Specs button. This will open the Survey Options Window



The Temperature Range is entered in the **End Points** section. The default values for the Temperature Range are 25°C (Start) to 100°C (End). Leave the default values.

The **Step Size** can be given as **Increments** or **Number of Steps**. By default, in the Temperature Survey the **Step Size** by **5** increments is selected. Leave the default values. Then click **OK**.



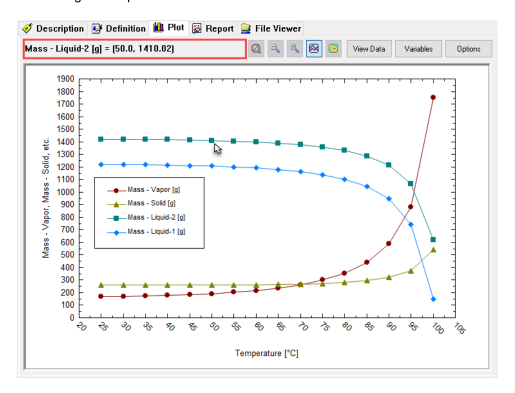
We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Survey Calculations*.

Analyzing the Results

Click on the Plot tab (Plot).

The default plot is the phase amounts in grams of Liquid-1 (Aqueous Phase), Liquid-2 (Organic Phase), Solid and Vapor.

The plot below shows that as temperature increases, the mass of Liquid-1 and Liquid-2 (organic) decreases, as they evaporate, and hence the Vapor phase increases. The solid phase mass remains constant up to 80°C, and then increases at higher temperatures.



Note: When the mouse-pointer is positioned over a point on one of the curves, the message box at the top of the plot indicates the variable and the coordinates of that point.

Pressure Survey

In the Survey by **Pressure**, the software computes solution properties based on a known composition and temperature at different pressures.

Example 19: Dissolution of CO₂ in water as a Function of Pressure

Many thermodynamic properties are less dependent on pressure than they are on temperature. Vapor-Liquid-Equilibrium (VLE) is affected by pressure.

In this example, the dissolution of CO₂ in water (which is a direct reflection of VLE) will be studied as a function of pressure.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Pressure Survey Calculation | | | | | | |
|-----------------------------|------------------------|-----------------------------------|--------------------|--|--|--|
| Calculation Settings | | Stream Composition and Conditions | | | | |
| Calculation Type | Survey | Stream Amount | Calculated | | | |
| Stream Name | Pressure Survey | Pressure Range | 5-100 atm | | | |
| Survey Name | CO2 dissolved in water | Step Size | Increment by 5 atm | | | |
| Name Style | Display Formula | Temperature | 25 °C | | | |
| Unit Set | Metric, Moles | H2O | 55.5082 moles | | | |
| Framework | MSE-SRK | CO2 | 10 moles | | | |

Add a new Stream

Click on the new Stream and press <F2> to change the name to Pressure Survey

Select the MSE-SRK thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Moles

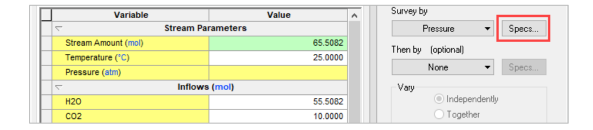
Enter the composition of the stream given in the table above

Go to the Add Calculation button and select Survey calculation

Change the Survey name to CO2 dissolved in water using the <F2> key

Select **Pressure** as Type of Survey

Click on the Specs button. This will open the Survey Options Window



The default pressure range is from 1-10 atm with an increment of 1 atm. Change the **Pressure Range** to 5-100 atm. Set the **Step Size** to **5 atm** increments. Then click **OK**.

We are ready to perform the calculation. Click on the Calculate button or press the <F9> key

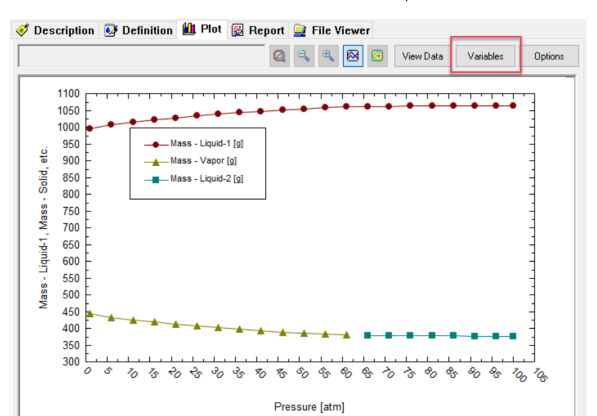
It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Survey Calculations*.

Analyzing the Results

Click on the **Plot** tab (Plot). The default plot is the phase amounts in grams of Liquid-1 (Aqueous Phase), Liquid-2 (Organic Phase), Solid and Vapor.

You can see that the Liquid-1 phase increases up to around 60 atm, and then it follows a constant trend as the pressure continues to increase. Regarding the Vapor phase, you can see that it shows a sudden decrease in mass from around 400 g at 60 atm to 0 g at 65 atm. The pressure of the system has increased in such a way that causes a phase change from vapor phase to Liquid-2 phase.

We are interested in the solubility of CO₂ in water, i.e. in the Liquid-1 phase. We can change the default plot to show the molecular CO2 dissolved in water (CO2-Liq1).

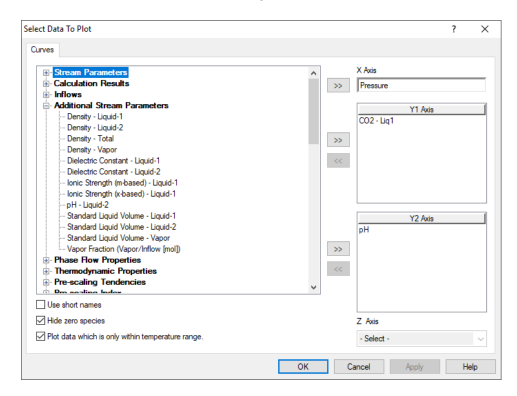


Click on the Variables button. This will open the Select Data to Plot window.

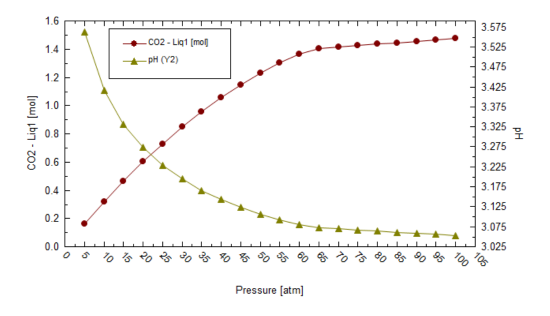
Double click or use the << button to remove the **Phase Amount** variables.

Look for **Liquid-1** and click on the **■** box to show all the available variables. **Select** CO2-Liq1 and put it in the Y1 Axis using the >> button.

Look for **Additional Stream Parameters** section and click on the ⊞ box to show all the available variables. Select **pH** and put it in the Y2 Axis using the >> button. Then click **OK**.



The default plot now shows the selected variables: The dissolved CO₂ in the water (Liquid-1 Phase) in the Y1 Axis, and the pH in the Y2 Axis. Both variables are presented as a function of pressure.



The dissolved CO_2 in solution increases as the pressure increases. When the pressure goes above 65 atm, the dissolved CO_2 remains nearly constant.

A similar behavior is seen in the pH. As the pressure increases, a decrease in pH is observed. This is attributed to the following equilibria dissociation reactions:

$$CO_{2,vap} \leftrightarrows CO_{2,aq}$$

$$CO_{2,aq} + 2H_2O \leftrightarrows H_3O^+ + HCO_3^-$$

$$HCO_3^- + H_2O \leftrightarrows H_3O^+ + CO_3^{2-}$$

Thus, as the amount of CO_2 in the aqueous phase increases, the dissociation reactions create more hydronium ions H_3O^+ that are available in solution, subsequently decreasing the pH 9 . As the pressure reaches values above 65 atm, the pH value remains constant. Hence, the pH is a direct reflection of the dissolved CO_2 in solution.

think simulation | getting the chemistry right

⁹ How can you have pH with no hydrogen ions (H⁺) in solution in the MSE framework? See: pH in MSE

Composition Survey

In the Survey by **Composition**, the software computes solution properties based on a given composition of species (or set of species) at a specific temperature and pressure.

Example 20: Dissolution of Calcite (CaCO₃) as a function of CO₂

Calcite is a scale that forms very easily in the production of oil and gas. It is known that adding CO₂ to water dissolves Calcite. Likewise, removing CO₂ from water precipitates calcite. In this example, you are going to evaluate the effect of different CO₂ concentrations on the dissolution of Calcite.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Composition Survey Calculation | | | | |
|--------------------------------|---------------------|-----------------------------------|-------------------------|--|
| Calculation Settings | | Stream Composition and Conditions | | |
| Calculation Type | Survey | Stream Amount | Calculated | |
| Stream Name | Composition Survey | Temperature | 25 °C | |
| Survey Name | Calcite Dissolution | Pressure | 75 atm | |
| Name Style | Display Formula | H2O | 55.5082 moles | |
| Unit Set | Metric, Moles | CaCO3 | 0.05 moles | |
| Framework | MSE | CO2 concentration range | 0 - 1 moles | |
| | | Step Size | Increment by 0.05 moles | |

Add a new Stream

Click on the new Stream and press <F2> to change the name to Composition Survey

Select the MSE thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the composition of the stream given in the table above

Go to the Add Calculation button and select Survey calculation

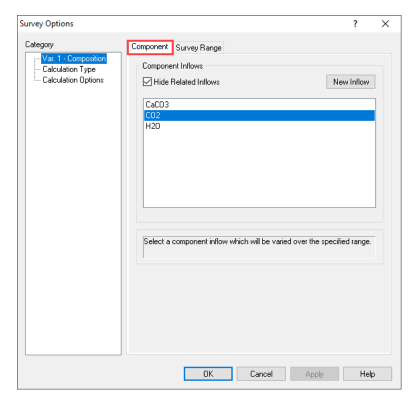
Change the Survey name to Calcite Dissolution using the <F2> key

Select **Composition** as Type of Survey.

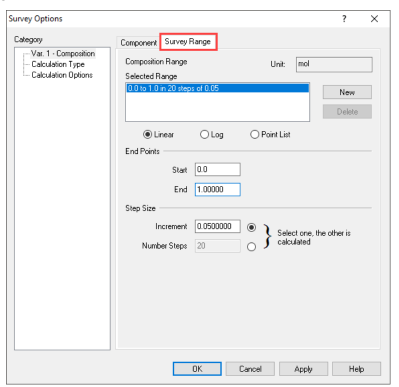
Note: In the summary box a message in red appears: 'Component not selected for composition survey variable'. This means that we need to select the component that we want to vary. In this example CO2.

To specify the component, click on the Specs button. This will open the Survey Options Window.

Under the Component tab, select CO2



Now, click on the **Survey Range** tab. Enter the CO2 composition range from 0 to 1 mol, by increments of 0.05 moles. Then click **OK**.

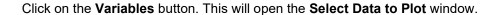


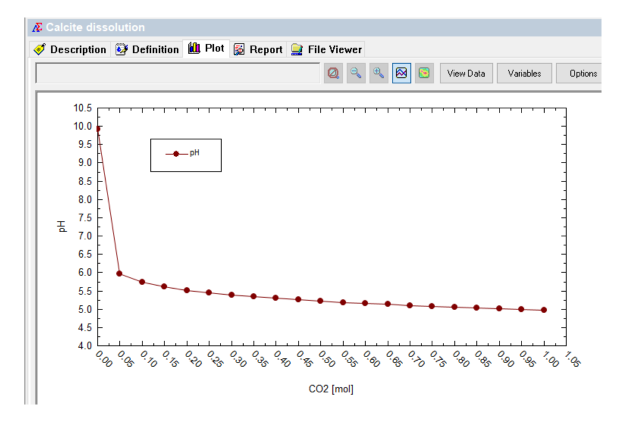
We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Survey Calculations*.

Analyzing the Results

Click on the **Plot** tab (Plot). The default plot is the pH as a function of total CO₂ in moles. As the concentration of CO₂ increases, the pH decreases (as it was explained in the example above).

We are interested in understanding the dissolution of Calcite as a function of CO₂. To see these results, we need to study how the moles of solid calcite change as the concentration of CO₂ increases.

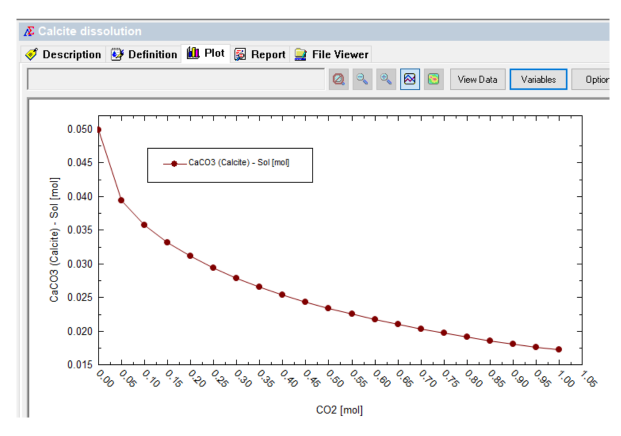




Double click or use the << button to remove the pH variable

Look for **Solid** section and click on the ⊞ box to show all the available variables. **Select CaCO3 (Calcite) – Sol** and put it in the Y1 Axis using the >> button. Click **OK**.

The default plot now shows the selected variable: CaCO3 (Calcite) - Sol as a function of CO2.



As expected, the amount of CaCO₃ (Calcite) decreases with increasing CO₂. In an oil production setting, when there is a significant pressure drop, CO₂ will be lost. This will decrease the solubility of CaCO₃ and will increase the likelihood of scale formation.

pH Survey

In the Survey by **pH**, the software adjusts the flowrate of acid or basic titrants to at a predefined set of pH values and computes the solution properties of a solution based on a given composition of species (or set of species) at a specific temperature and pressure.

Example 21: A Speciation Diagram for the Carbonic Acid System as a Function of pH

In this survey calculation, a 0.001 molal CO₂ solution is titrated with HCl and NaOH. Instead of defining the amount of acid/base to add, the pH will be specified, and the software will compute the amounts of HCl and NaOH needed.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| pH Survey Calculation | | | |
|-----------------------|----------------------|-----------------------------------|------------------|
| Calculation Settings | | Stream Composition and Conditions | |
| Calculation Type | Survey | Stream Amount | Calculated |
| Stream Name | pH Survey | Temperature | 25 °C |
| Survey Name | Carbonic Acid System | Pressure | 1 atm |
| Name Style | Display Formula | H2O | 55.5082 moles |
| Unit Set | Metric, Moles | CO2 | 0.001 moles |
| Framework | MSE | pH range | 4-12 |
| | | Step Size | Increment by 0.1 |

Add a new Stream

Click on the new Stream and press <F2> to change the name to pH Survey

Select the MSE (Default) thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the composition of the stream given in the table above

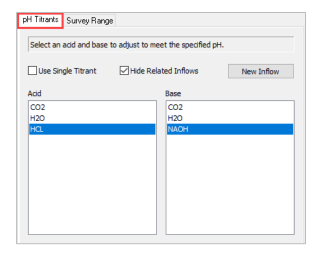
Go to the Add Calculation button and select Survey calculation

Select **pH** as Type of Survey

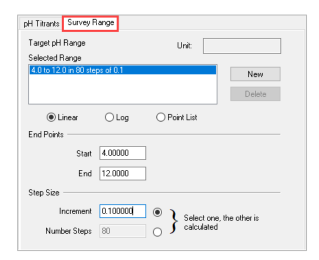
Change the **Survey** name to **Carbonic Acid System** using the **<F2>** key

The calculation can be run without modification. The default titrants are **HCI** and **NaOH**, and the default pH range is set from **0** to **14** with a step size of **1** increment. However, this pH range is too broad for the carbonate system, in which most of the speciation occurs between pH=4 and 12. Additionally, it is ideal to specify finer increments to see a smoother curve.

Click on the **Specs** button. This will open the **Survey Options Window** In the **pH Titrants** tab, leave the default pH Titrants: **HCI** and **NaOH**



Now, click on the **Survey Range** tab. Enter the pH range from 4 to 12, with increments of 0.1. Then click **OK**.



We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Survey Calculations*.

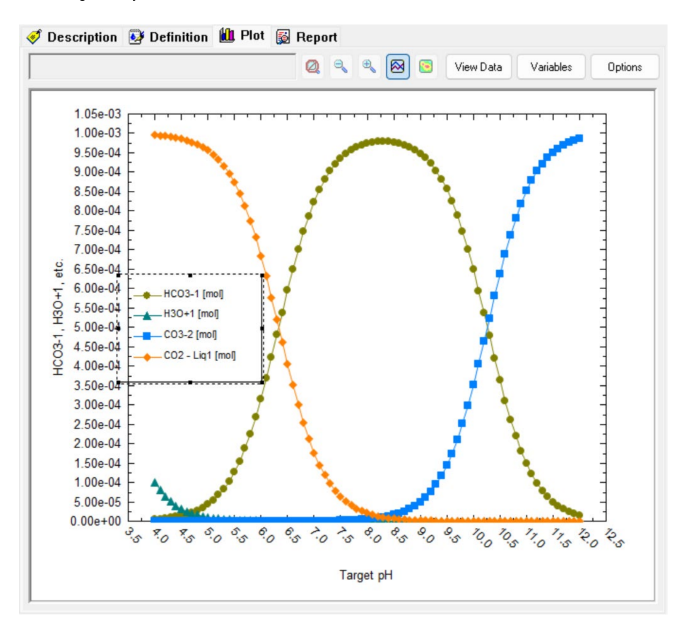
Analyzing the Results

Click on the **Plot** tab (Plot). The default plot is the moles of added HCl and NaOH vs Target pH. This is not the target variable, and you will change it to display the carbonate-containing species.

Click on the Variables button

Remove the NaOH and HCI from the Y1 axis (double click or use the << button)

Look for **Liquid-1** section and click on the \blacksquare box to show all the available species. **Select** the following species: CO2-Aq, HCO3-1, CO3-2, and H₃O⁺ and put them in the Y1 Axis using the >> button or by double clicking. When you are done, click **OK**.

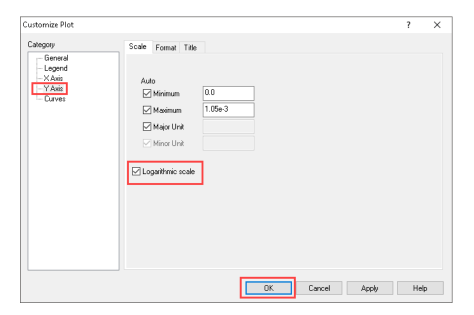


You can also present this plot on a semi-log scale.

Click on the Options button. This will bring the Customize Plot window

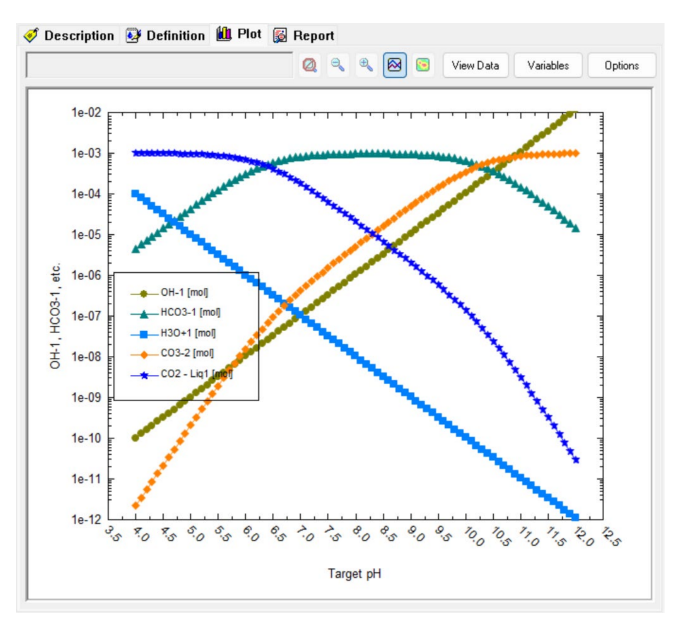
Select Y Axis in the Category window

Check the Logarithmic scale box, and then click **OK**.



Before analyzing the plot, let's add the OH- species.

Click on the **Variables** button, look for OH-1 in the **Liquid-1** section, add it to the Y1 Axis, and then click **OK**. Now we are ready to analyze the plot.



At lower pH, CO2-Aq dominates the system, and notice how the HCO3-1 slope is 1 and the CO3-2 slope is 2. Likewise, in the pH region where CO3-2 dominates (high pH), the HCO3-1 slope is -1 and the CO2-Aq slope is -2. These $\frac{mole}{pH}$ slopes are based on the number of H+ ions added or removed in the chemical reactions. See reactions below.

$$CO_{2,aq} + 2H_2O \leftrightarrow H_3O^+ + HCO_3^-$$

 $HCO_3^- + H_2O \leftrightarrow H_3O^+ + CO_3^{2-}$

Note the pH where the CO2_(aq) and HCO₃-1 lines intersect (pH~8.3), and where the HCO3-1 and CO3-2 lines intersect (pH~10.3). These pH values are the same as the pKa values (the equilibrium equations the specific equilibrium reactions).

As NaOH is added to convert HCO₃- to CO₃-2, a portion of the NaOH remains as free base, OH-1. This fraction increases at higher pH values, and its concentration is reflected in the calculated pH. A similar pattern is seen at pH below ~3 for HCl and H+.

Vapor Fraction / Vapor Amount Survey

The purpose of the **Vapor Fraction/Vapor Amount Survey** calculation is to set a specific fraction of the stream to the vapor phase, and compute the temperature or pressure required to achieve the specified Vapor Fraction/Vapor amount.

Example 22: Seawater Evaporation

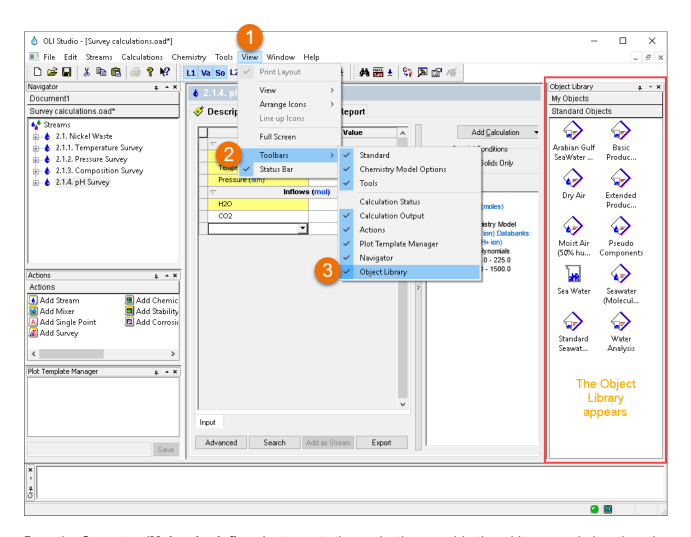
Seawater contains a variety of salts, and when seawater evaporates, these solids are left behind. The most abundant salt in seawater is sodium chloride (NaCl) which is commonly referred to by its mineral name halite.

You will evaporate seawater to dryness and look at the solids that form. What is interesting is the number of phases that can be produced when seawater evaporates, giving you a sense of how certain natural systems build up sediment of a particular nature.

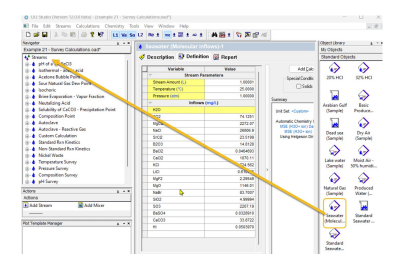
In this example we will also introduce the Object Library tool.

Starting the Simulation

Open the **Object Library** window – from the menu, View > Toolbar > Object Library. The Object Library appears on the right-hand side of the window.



Drag the **Seawater (Molecular inflows)** stream to the navigation panel in the white space below the other streams and calculation objects or on top of the "Streams" at the top of the Navigator tree



The inputs and parameters are automatically populated when selecting the Seawater (molecular inflows) from the Object Library.

Click on the new Stream and press <F2> to change the name to Vapor Fraction Survey

Go to the Add Calculation button and select Survey calculation

Change the **Survey** name to **Seawater Evaporation** using the **<F2>** key

Select Vapor Fraction as Type of Survey

Click on the Specs button. This will open the Survey Options Window

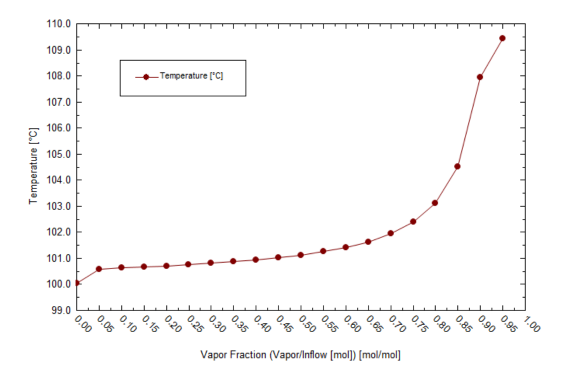
The default Vapor Fraction range is from 0-1 with an increment of 0.1. Change the **Vapor Fraction Range** to **Survey End** to 0.95. Set the **Step Size** to **0.05** increments. Then click **OK**.

We are ready to perform the calculation. Click on the Calculate button or press the <F9> key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Survey Calculations*.

Analyzing the Results

Click on the **Plot** tab (Plot). The default plot is the Vapor Fraction plot (as the dependent variable) vs Temperature. It shows that seawater boils initially at 100°C, and at near complete evaporation (95%) the temperature is 109.4°C.



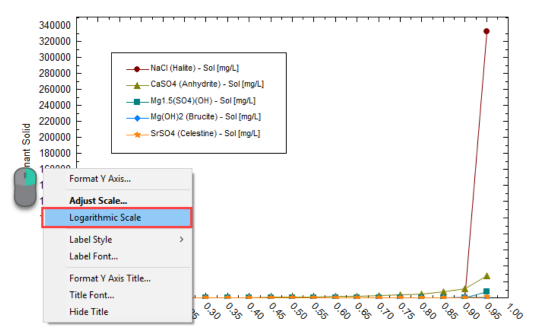
We are also interested to know which solid phases precipitate as the water evaporates.

Click on the Variables button

Remove **Temperature** from the Y1 axis (double click or use the << button)

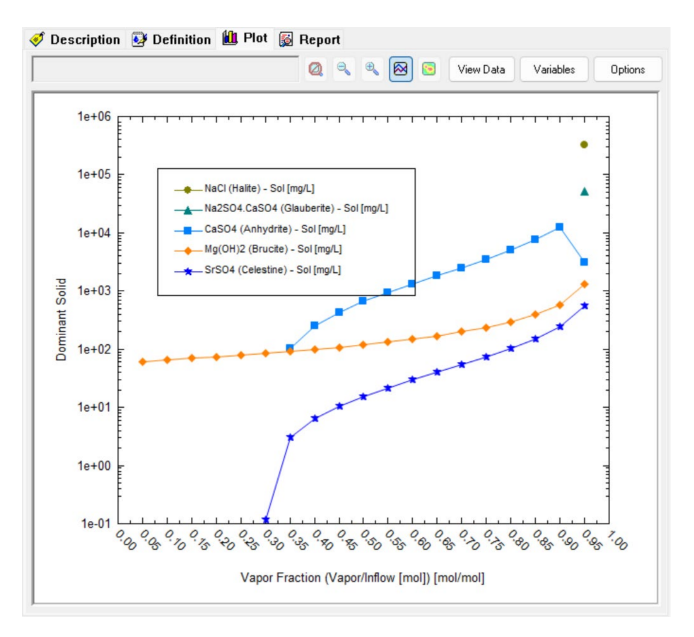
Look for **Solid** section and click on the 🗈 box to show all the available species. **Select** the option **Dominant Solids** and put it in the Y1 Axis using the >> button or by double clicking. When you are done, click **OK**.

The default plot is not very easy to read because halite dominates the solids and because the units are in mg/l and the liquid volume decreases from left-to-right on the plot. To see the plot more clearly, **right click** on any number in the Y axis, and select the **Logarithmic Scale** option.



Vapor Fraction (Vapor/Inflow [mol]) [mol/mol]

Plot in semilogarithmic scale



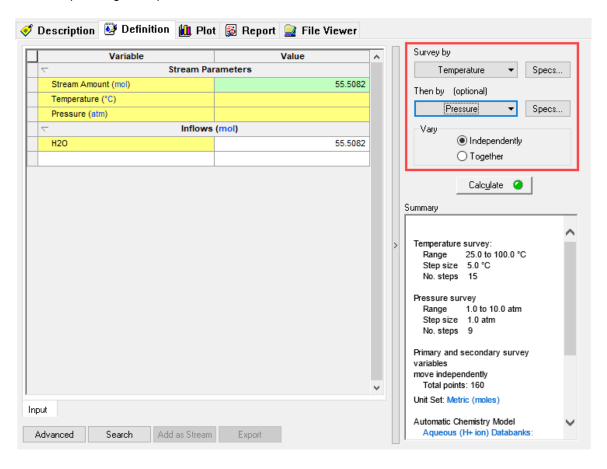
As the water evaporates $CaSO_4$, $Mg(OH)_2$ and $SrSO_4$ salts precipitate at all temperatures. NaCl and a Mg/SO4/OH double salt starts to precipitate at 95% vapor fraction.

Dual Surveys

Dual Surveys allow you to study the effect of two independent variables on your system under study.

The option for the dual survey is highlighted in the red box in the image below. There are three options available in the optional survey, Temperature, Pressure and Composition.

You also have the option of varying the selected variables independently or together. We will explore this in more detail in upcoming examples.



Temperature and Pressure Survey

In this type of survey two variables are adjusted simultaneously, temperature and pressure. This type of survey creates a matrix of results, which can then be interpreted using the plot function in different ways.

Example 23: Dissolution of CO2 in water as a Function of Temperature and Pressure

In Example 19, the dissolution of CO₂ in water was studied only as a function of pressure. In this case, we are going to study the effect of both temperature and pressure on the CO₂ dissolution.

In this example the Contour Plot will be used to analyze the results.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Temperature and Pressure Survey Calculation | | | |
|---|-----------------------|-----------------------------------|--------------------|
| Calculation Settings | | Stream Composition and Conditions | |
| Calculation Type | Survey | Stream Amount | Calculated |
| Stream Name | T/P Survey | Pressure Range | 5-100 atm |
| Survey Name | CO2 dissolution – T/P | Step Size | Increment by 5 atm |
| Name Style | Display Formula | Temperature Range | 5-100 °C |
| Unit Set | Metric, Moles | Step Size | Increment by 5 °C |
| Framework | MSE-SRK | Vary | Independently |
| | | H2O | 55.5082 moles |
| | | CO2 | 10 moles |

Add a new Stream

Click on the new Stream and press **<F2>** to change the name to *T/P Survey*

Select the MSE-SRK thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the composition of the stream given in the table above

Go to the **Add Calculation** button and select **Survey** calculation

Change the Survey name to CO2 dissolution T/P using the <F2> key

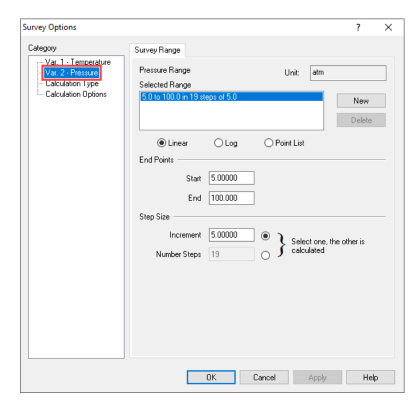
Select Temperature and then by Pressure

Click on the Temperature Specs button. This will open the Survey Options Window

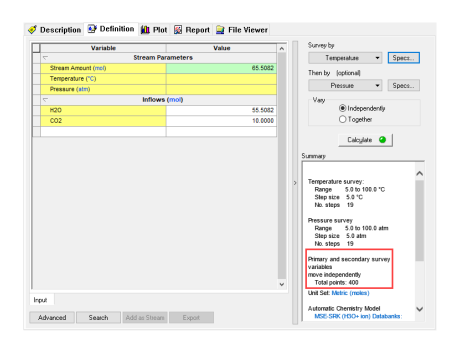
Change the **Temperature Range** to 5-100 °C. Set the **Step Size** to **5** °C increments.

Click on Var. 2 - Pressure, in the Category Window to define the Pressure Range.

Change the Pressure Range to 5-100 atm. Set the Step Size to 5 atm increments. Then click OK.



In the summary box, notice that a total of 400 points will be calculated. We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key



It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Survey Calculations*.

Analyzing the Results

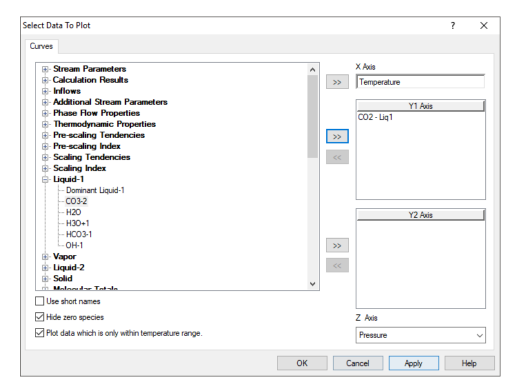
Click on the **Plot** tab (Plot). The default plot is the phase amounts in grams of Liquid-1 (Aqueous Phase), Liquid-2 (Organic Phase), Solid and Vapor.

We are interested in the solubility of CO₂ in water, i.e., in the Liquid-1 phase. We can change the default plot to show the molecular CO₂ dissolved in water (CO₂-Liq₁).

Click on the Variables button. This will open the Select Data to Plot window.

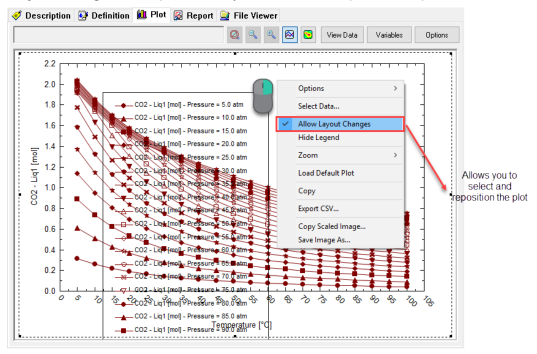
Double click or use the << button to remove the **Phase Amount** variables.

Look for **Liquid-1** and click on the ⊕ box to show all the available variables. **Select** CO2-Liq1 and put it in the Y1 Axis using the >> button. Then click **OK**.

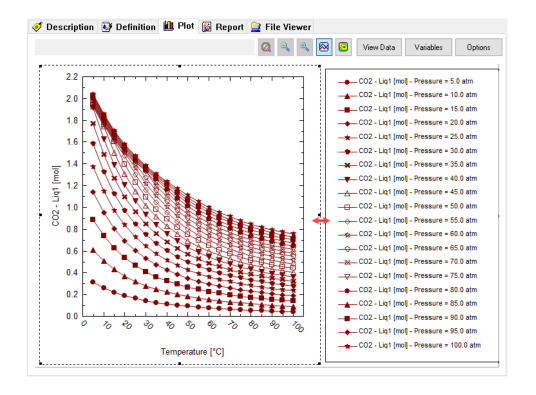


The default plot now is showing the dissolved CO₂ in the water (Liquid-1 Phase) in the Y1 Axis as function of pressure. Unfortunately, the legend is covering the plot.

To move the legend to the side without interfering with the results, right click on any white space within the plot and **Allow Layout Changes**. This option allows you to select and reposition the plot.



Resize the plot to make it smaller and create space for the legend. Drag the legend to the right.

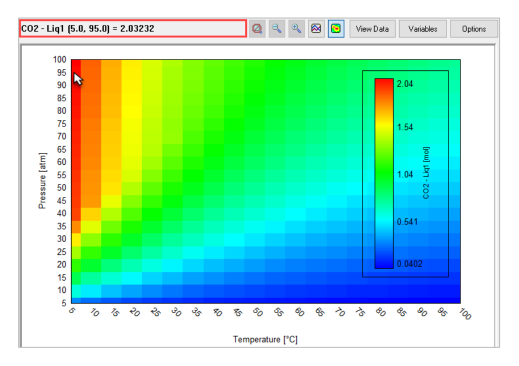


As the temperature increases (at a fixed pressure) the solubility of CO_2 decreases. This is to be expected since at higher temperatures it is easy to overcome the vapor pressure and CO_2 is released as gas. As the pressure increases (at a fixed temperature) the solubility of CO_2 in solution increases.

The plot above can be represented better using a **Contour Plot**. To convert the results into a contour plot view, simply click on the contour plot icon ().

The resulting plot is now a pixilated color plot showing the moles of molecular CO2 dissolved in the Liquid-1 (Aqueous) phase, at each T and P value.

The number of moles of CO₂ is characterized by a different color that is shown in the legend (maximum value is given in red, and minimum value is given in blue). You can mouse over any location in the plot and the moles of CO₂ dissolved in the Liquid-1 phase at each T and P value will be shown in the left corner box.

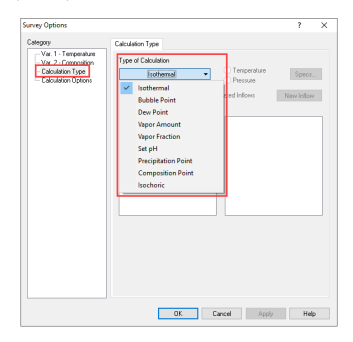


Note: The legend scale can also be optimized by either right-mouse-clicking on the Legend and selecting Options>Plot Options>Contour or clicking on the Options button in the upper right and selecting Contour. Both open to the Contour options window, where the color and range can be modified.

In the contour plot, you can see the effect of both Temperature (x-axis) and Pressure (y-axis) on the solubility of CO_2 in water. Higher solubilities are obtained at high pressures and low temperatures, while lower solubilities are obtained at high temperatures and low pressures.

Survey by changing the single point calculation type

The sections presented so far have presented single and dual surveys. In addition to adjusting these survey variables, you can select one of several single point calculations embedded within the survey. The embedded single point calculations include Isothermal (default), bubble point, dew point, vapor amount, vapor fraction, set pH, precipitation point, composition point, and isochoric.



Temperature Survey with a Precipitation Point Calculation

In this **Temperature Survey**, the Calculation Type will be changed from Isothermal Calculation to **Precipitation Point**. This will allow us to model the solid solubility vs temperature.

Example 24: Solubility of Halite as a Function of Temperature

In this example, we are going to study the solubility of NaCl (halite) as a function of temperature.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Temperature Survey with a Precipitation Point Calculation | | | | |
|---|------------------------|-----------------------------------|-------------------|--|
| Calculation Settings | | Stream Composition and Conditions | | |
| Calculation Type | Survey | Stream Amount | Calculated | |
| Stream Name | Solubility vs T | Pressure | 1 atm | |
| Survey Name | Halite Solubility vs T | Temperature Range | 5-100 °C | |
| Calculation Type | Precipitation Point | Step Size | Increment by 5 °C | |
| Name Style | Display Formula | H2O | 55.5082 moles | |
| Unit Set | Metric, Moles | NaCl | 0 moles | |
| Framework | MSE | | | |

Add a new Stream

Click on the new Stream and press **<F2>** to change the name to *Halite Solubility vs T and P* Select the **MSE** thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the composition of the stream given in the table above

Go to the Add Calculation button and select Survey calculation

Change the Survey name to Halite Solubility vs T using the <F2> key

Select **Temperature** as the type of survey

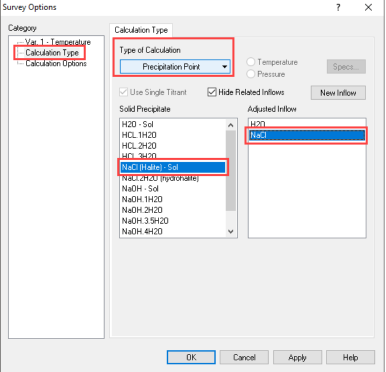
Click on the Temperature Specs button. This will open the Survey Options Window

Change the **Temperature Range** to 0-100 °C. Set the **Step Size** to **5** °C increments.

Click on Calculation Type, in the Category Window to define the Type of Calculation

Under Type of Calculation Change the Default Isothermal to **Precipitation Point** (use the drop-down arrow)

Select NaCl (Halite)-Sol as the Solid Precipitate and NaCl as the Adjusted Inflow. Then click OK.



The **Calculation Parameters** section is now shown in the Definition Tab, highlighting the Precipitant (NaCl (Halite)) and the Adjusted Inflow (NaCl).

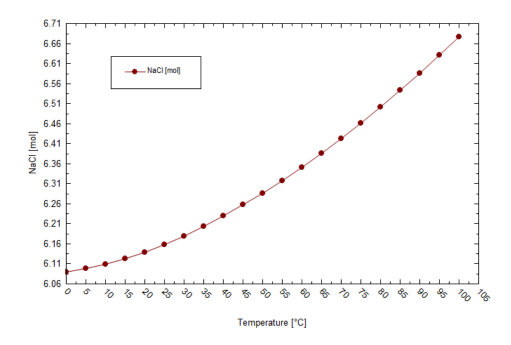
| Variable | Value |
|---------------------|---------------------|
| ▽ Strea | am Parameters |
| Stream Amount (mol) | 55.5082 |
| Temperature (°C) | |
| Pressure (atm) | 1.00000 |
| | ation Parameters |
| Calculation Type | Precipitation Point |
| Precipitant: | NaCl (Halite) |
| Adjusted Inflow: | NaCl |
| - In | flowe (mel) |
| H2O | 55.5082 |
| NaCl | 0.0 |

We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Survey Calculations*.

Analyzing the Results

Click on the **Plot** tab (Plot). The default plot is the amount of NaCl in moles added to 1 kg of water before Halite precipitates as a function of Temperature.



| Halite solubility increase at 100°C. | ses slightly with temp | perature, from 6.16 m | oles/1kg H₂O at 25º0 | C to 6.68 moles/1kg H₂C |
|--------------------------------------|------------------------|-----------------------|----------------------|-------------------------|
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Example 25(Part II): Solubility of Halite as a Function of Temperature and Pressure

The effects of pressure on mineral solubility is generally less important than temperature. There is still an effect that can be observed. The same stream created on as in example 25 will be used.

Starting the Simulation

Under the **Solubility vs T** stream add a new Survey (see Stream information in Example 24)

Change the Survey name to Halite Solubility vs T and P using the <F2> key

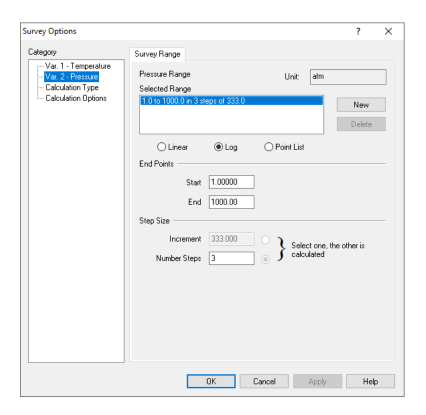
Select Survey by Temperature and then by Pressure (This is a dual survey)

Click on the Temperature Specs button. This will open the Survey Options Window

Change the **Temperature Range** to 0-100 °C. Set the **Step Size** to **5** °C increments.

Click on Var. 2 – Pressure, in the Category Window to define the Pressure Range.

Change the Scale to **Log**, change the **Pressure Range** from 1 to 1000 atm, and set the **Number of Steps** to **3** (i.e. it will plot the following pressures: 1, 10, 100 and 1000 atm). Then click **OK**.



Click on Calculation Type, in the Category Window to define the Type of Calculation

Under Type of Calculation Change the Default Isothermal to **Precipitation Point** (use the drop-down arrow)

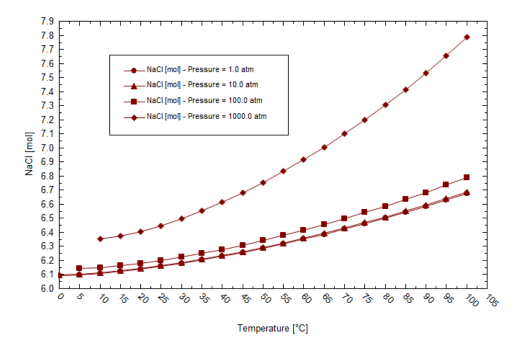
Select NaCl (Halite)-Sol as the Solid Precipitate and NaCl as the Adjusted Inflow. Then click OK.

We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key.

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Survey Calculations*.

Analyzing the Results

Click on the **Plot** tab (Plot). The default plot is the amount of NaCl in moles added to 1 kg of water before Halite precipitates as a function of Temperature.



The effect of pressure is significant between 100 and 1000 atm relative to the effects between 1 and 100 atm.

We can also present the Halite Solubility using a Contour Diagram. It would be ideal to define a finer range of temperature and pressure.

Return to the **Definition Tab**

Click on the **Temperature Specs** button.

Change the **Temperature Range** to 0-150 °C. Set the **Step Size** to **2** °C increments.

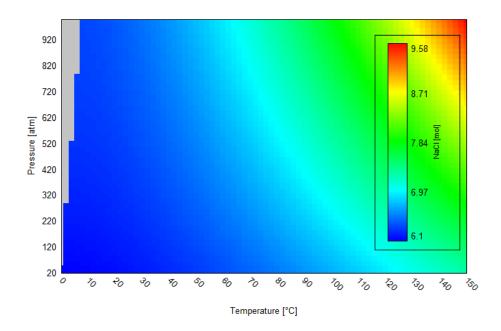
Click on Var. 2 - Pressure, in the Category Window to define the Pressure Range.

Change the survey to **Linear** scale, change the **Pressure Range** from 20 to 1000 atm, and set the **Step Size** to **20 atm** increments. Then click **OK**.

Note: In the Summary box we can see that survey creates a matrix of 75 steps x 49 steps for a total of 76 temperature and 50 pressure points, or 3800 points in total.

Click the Calculate button. This calculation will take a few minutes to compute.

Click on the **Plot** tab and change the Plot to contour ().



The solubility is shown to increase as temperature and pressure increase. The increase is up to 50% from the low values. A few of the points failed to converge at the low temperature and high-pressure conditions (shown in gray).

Composition Survey with a Bubble and Dew Point Calculation

In this **Composition Survey**, the Calculation Type will be changed from Isothermal Calculation to **Bubble and Dew Point calculations**. This will allow us to model the vapor liquid equilibria (VLE) of mixtures.

Example 25: Ethanol-Water Azeotrope

In this example, you are going to calculate both the bubble point and dew points as a function of Ethanol-Water composition.

The two data sets will then be transferred to a spreadsheet where the curves will be plotted together to locate the azeotrope. You will need a spreadsheet/plotting software to complete this case.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Composition Survey with a Bubble and Dew Point Calculation | | | |
|--|--------------------------------|-----------------------------------|-----------------------|
| Calculation Settings | | Stream Composition and Conditions | |
| Calculation Type | Survey | Stream Amount | Default – 1kg |
| Stream Name | Composition survey - Azeotrope | Pressure | 1 atm |
| Survey Name | Ethanol-Water Azeotrope | Temperature | It will be calculated |
| Calculation(s) Type | Bubble Point / Dew Point | H2O | 100 mass % - default |
| Name Style | Display Name | Ethanol | 0 mass% |
| Unit Set | Metric, Mass Frac. | Composition Range | 0 -100 mass% |
| Framework | MSE | Step Size | Increment by 2 mass% |

Calculating the Bubble Point

Add a new Stream

Click on the new Stream and press <F2> to change the name to Composition survey - Azeotrope

Select the MSE thermodynamic Framework

Click on the Names Manager Icon and select the Display Name option

Click on the Units Manager Icon, and select Metric, Batch, Mass Frac.

Enter the composition of the stream given in the table above

Go to the Add Calculation button and select Survey calculation

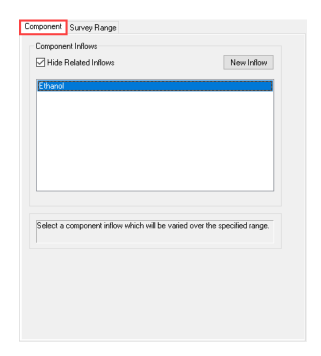
Select Composition as the type of survey

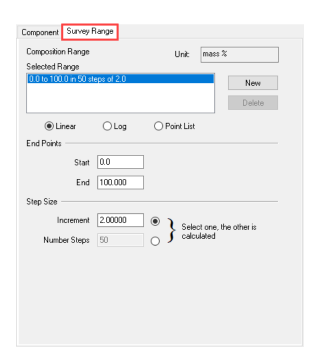
Change the **Survey** name to **Ethanol-Water Azeotrope** using the **<F2>** key

Click on the Composition Specs button. This will open the Survey Options Window

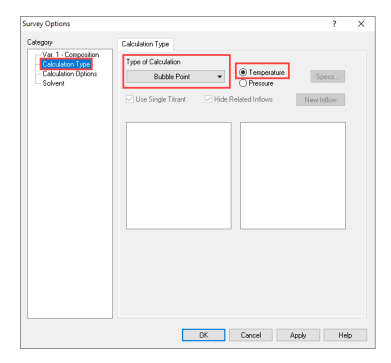
Under the Component tab, select Ethanol

Click on the **Survey Range tab.** Change the **Composition Range** to 0-100 mass%. Set the **Step Size** to **2** mass% increments.





Click on **Calculation Type**, in the Category Window to define the Type of Calculation
Under Type of Calculation Change the Default Isothermal to **Bubble Point** (use the drop-down arrow). Then click **OK**.

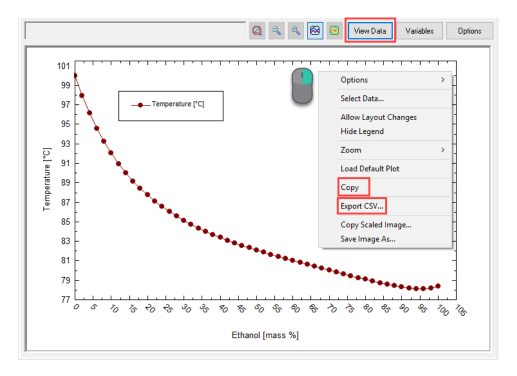


We are ready to perform the calculation. Click on the Calculate button or press the <F9> key.

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Survey Calculations*.

Analyzing the Results

Click on the **Plot** tab (Plot). The default plot is the bubble point temperature vs the mass fraction ethanol. As the mass% of ethanol increases the bubble point temperature decreases.

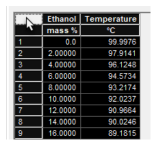


You have 3 different options to retrieve this data to use in your preferred plotting software.

Option 1: Right-click on the white region of the plot to open the drop-down menu and select Copy.

Option 2: Right-click on the white region of the plot to open the drop-down menu and select Export CSV.

Option 3: Click on the **View Data** button, select all data by clicking on the upper left corner of the table, and copy the data using **Ctrl+C**.



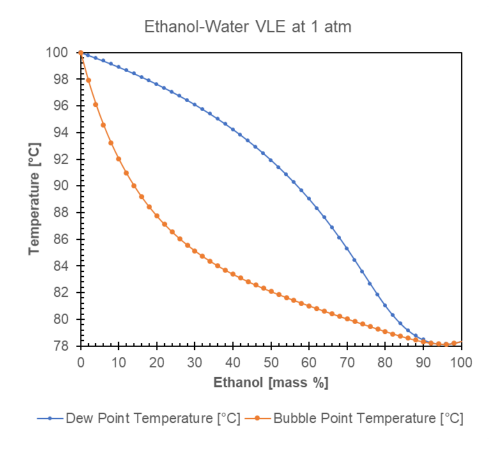
In this case let's use **Option 1**. Right click and **Copy** the data, then open your preferred plotting tool and **Paste** the data.

Calculating the Dew Point

Return to the **Definition tab**

Click on the **Composition Specs** button. This will open the **Survey Options Window** Click on **Calculation Type**, in the Category Window to define the Type of Calculation Change the calculation type to **Dew Point** (use the drop-down arrow). Then click **OK**. **Recalculate**, and click on the **Plot tab**

Right click and **Copy** the data, then open your preferred plotting tool and **Paste** the data **Plot** the two temperature curves vs mass% of ethanol and format as needed

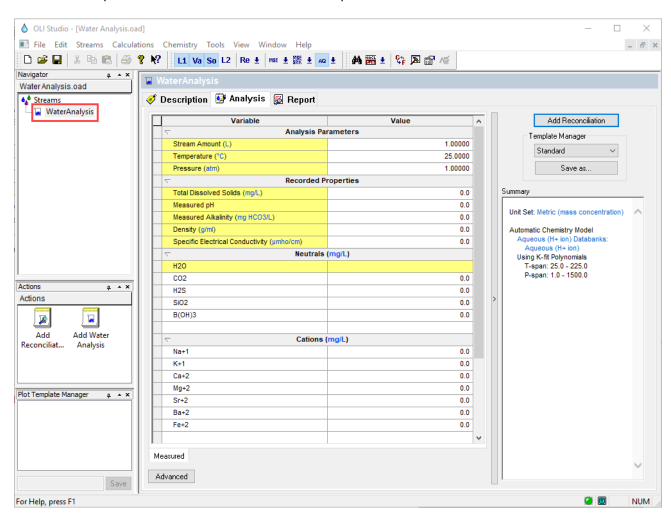


The azeotrope occurs at about 95% ethanol where the two curves intersect, at 78.15 °C.

Section 3. Water Analysis (Ionic Inputs)

OLI Studio: Stream Analyzer can accept two input types: molecular inflows and ionic inflows.

Sections 2 and 3 focused on molecular inflows. In this section, we will focus on **ionic inflows**, and we will learn how to enter cations and anions using the object called **Water Analysis**. Additionally, we will show how the reconciled sample can be converted into a molecular representation.



Basic Terminology

When reviewing laboratory analysis of water samples, it is quite common for the positive ions (cations) and the negatively charged ions (anions) in solution to not balance. This may be due to the precision limits of the various experimental procedures used to measure the ions - i.e., some ions may not have been analyzed. These solutions must have a neutral charge. Stream Analyzer will adjust/modify inflows in order to balance the charges and make the solution neutral. This adjusting procedure is referred to as **Reconciliation**.

The pH and the alkalinity of the solution are frequently measured. However, since the analysis is experimental and subject to errors, the pH and alkalinity values that are calculated by the Water Analysis tool may be different from what is measured experimentally. Stream Analyzer can also reconcile this difference.

Scaling

Scaling is the deposition of a mineral salt on processing equipment. Scaling is a result of supersaturation of mineral ions in the process fluid. Through changes in temperature, or solvent evaporation or degasification, the concentration of salts may exceed the saturation, leading to a precipitation of solids (usually crystals). The saturation level of a salt in water is a good indicator of the potential for scaling.

The potential for scaling is calculated using the solubility product constant (K_{sp}) and Ion Activity Product (IAP) definitions.

Solubility Product Constant, K_{sn}

The solubility of ionic compounds of salts and minerals in water are governed by a solubility equilibrium expression and a solubility product constant known as K_{sp} . It is important to note that the solubility product, K_{sp} is a function of both temperature and pressure. Consider the general dissolution reaction below (in aqueous solutions):

$$aA_{(s)} \rightleftharpoons bB_{(aq)} + dD_{(aq)}$$

With equilibrium constant K_{sp} defined as:

$$K_{sp} = (a_B)^b \cdot (a_D)^d$$

Where, a_B and a_D are the activities of the aqueous species. The activity of any species i is defined as the product of its concentration in molality by its corresponding activity coefficient:

$$a_i = m_i \gamma_i$$

Ion Activity Product, IAP

A real solution may not be in the state of equilibrium. This non-equilibrium state is described by the ion activity product (IAP). It has the same form as the equilibrium constant K_{sp} , but involves the actual activities of the species in solution.

$$IAP = (a_B)_{actual}^b \cdot (a_D)_{actual}^d$$

Scaling Tendency and Scale Index

The Scaling Tendency (ST) is defined as the ratio of the Ion Activity Product (IAP) divided by the equilibrium constant (K_{SD}).

$$ST = \frac{IAP}{K_{SD}}$$
 Equation (1)

Scaling tendencies are essentially saturation ratios. Thus, if

 $\mathit{ST} < 1$ Indicates sub-saturation, and the solid is not expected to form

ST = 1 Indicates saturation, and the solid is in equilibrium with water

ST > 1 Indicates supersaturation, and solids will form

The Scale Index (SI) (aka: Saturation Index in the literature), is given by the following relationship:

$$SI = \log_{10} \left(\frac{IAP}{K_{sp}} \right)$$
 Equation (2)

Thus, if

SI < 0 Indicates sub-saturation, and the solid is not expected to form

SI = 0 Indicates saturation, and the solid is in equilibrium with water

SI > 0 Indicates supersaturation, and solids will form

Pre-scaling Tendency and Scale Index

Pre-Scaling tendency is defined as the scaling tendency before any solids are formed (this can be seen as all the species suspended in solution). The same equations (Equations 1 and 2) are used for calculating ST and SI.

The Pre-Scaling tendency is reported in the software as Pre-Scale, with its respective SI, index.

Difference between Post-scale and Pre-scale

Pre-Scale: The saturation ratio before solids precipitate.

Post-Scale: The saturation ratio **after** solids precipitate (if solids are selected).

Another way to interpret these two definitions is:

Pre-Scale represents the condition before any solids are allowed to form. This is a non-equilibrium condition and can be viewed as the condition where time = 0.

Post-Scale Tendency is the saturation ratio after all potential solids come to equilibrium with water. This is the true equilibrium condition ($time = \infty$).

Example 26: Calculation the Scaling Tendency and the Scale Index of $CaSO_4 \cdot 2H_2O$

Let's calculate the Scaling Tendency and the Scale Index of 0.01 moles of gypsum $(CaSO_4 \cdot 2H_2O)$ dissolved in 1 kg of water at 25°C and 1 atm.

The equilibrium expression for the dissolution reaction is:

$$CaSO_4 \cdot 2H_2O \rightleftharpoons Ca^{2+} + SO_4^{2-} + 2H_2O$$

Where, $K_{sp} = 3.2 \times 10^{-5}$

The molal concentration and the activity coefficients for each one of the species are:

$$[Ca^{+2}] = 0.0093 m$$
 $\gamma_{Ca^{+2}} = 0.4663$

$$[SO_4^{-2}] = 0.0093 m$$
 $\gamma_{SO_4^{-2}} = 0.4663$

$$a_{H_2O} = 0.9997$$

Calculating the IAP

With the information given above, we can calculate the IAP as follows:

$$IAP = a_{Ca^{+2}} \cdot a_{SO_4^{-2}} \cdot (a_{H_2O})^2$$

$$IAP = (m_{Ca^{+2}} \gamma_{Ca^{+2}}) \cdot (m_{SO_4^{-2}} \gamma_{SO_4^{-2}}) \cdot (a_{H_2O})^2$$

$$IAP = (0.0093 \times 0.4663) \cdot (0.0093 \times 0.4663) \cdot (0.9997)^2$$

$$IAP = 1.879 \times 10^{-5}$$

Calculating the Scaling Tendency

$$ST = \frac{1.879 \times 10^{-5}}{3.2 \times 10^{-5}}$$
$$ST \approx 0.587$$

This result indicates that the solution is under-saturated with respect to calcium sulfate.

Calculating the Scaling Index

$$SI = \log_{10} \frac{IAP}{K_{sp}} = \log_{10} ST$$
$$SI = \log_{10} (0.587)$$
$$SI \approx -0.231$$

Entering Data for Water Analysis

The grid for water analysis has 5 different grids where we need to enter information: Analysis Parameters, Recorded Properties, and Neutrals, Total Ions, Cations and Anions concentration in solution.

The **Analysis Parameters** grid is where you enter the conditions at which the sample properties were measured. The default values are 1 L, 25°C and 1 atm.

| Stream Amount (L) | 1.00000 | | | |
|-------------------|---------|--|--|--|
| Temperature (°C) | 25.0000 | | | |
| Pressure (atm) | 1.00000 | | | |

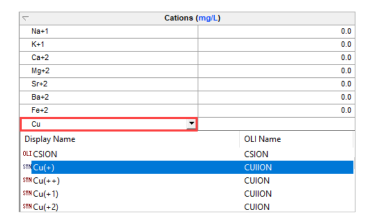
The **Recorded Properties** grid is where you enter the laboratory measured properties of the solution. These are: Total Dissolved Solids (TDS), Measured pH, Measured Alkalinity, Measured Total Inorganic Carbon (TIC), Density, and Specific Electrical Conductivity. You can always change the units of these properties by clicking on the units highlighted in blue. This action will open the Units Manager Window.

| Recorded Properties | | | | |
|--|-----|--|--|--|
| Total Dissolved Solids (mg/L) | 0.0 | | | |
| Measured pH | 0.0 | | | |
| Measured Alkalinity (mg HCO3/L) | 0.0 | | | |
| Measured TIC (mol C/L) | 0.0 | | | |
| Density (g/ml) | 0.0 | | | |
| Specific Electrical Conductivity (µmho/cm) | 0.0 | | | |

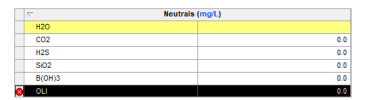
The concentration of Neutrals, Total lons, Cations and Anions is entered in the section. The Water Analysis grid comes already prepopulated with some of the most common cations, anions, and neutrals species found in laboratory water analyses. If your species is not present in the prepopulated grid, simply click on the white grid and type the ion or neutral of interest. If it is a **cation**, type the element followed by a plus (+) sign and the corresponding oxidation state, e.g. Cu+2. If it is an **anion**, type the element followed by a minus (-) sign and the corresponding oxidation state, e.g. Br-1. If it is a **neutral**, simple type the species either using the formula name or its name, as has been shown in the previous sections.

| Neutrals (m | g/L) |
|-------------|--------------|
| H2O | |
| CO2 | 0.0 |
| H2S | 0.0 |
| SiO2 | 0.0 |
| B(OH)3 | 0.0 |
| | |
| | ng/L) |
| P as PO4-3 | 0.0 |
| Si as Si02 | 0.0 |
| B as B(OH)3 | 0.0 |
| | |
| Cations (mg | g/L) |
| Na+1 | 0.0 |
| K+1 | 0.0 |
| Ca+2 | 0.0 |
| Mg+2 | 0.0 |
| Sr+2 | 0.0 |
| Ba+2 | 0.0 |
| Fe+2 | 0.0 |
| | |
| | ı/L) |
| CI-1 | 0.0 |
| S04-2 | 0.0 |
| HCO3-1 | 0.0 |
| HS-1 | 0.0 |
| C2H3O2-1 | 0.0 |

The Water Analysis grid also contains search aids to find a specific cation or anion. The first search aid is the drop list located in each the cation, anion, and neutral grid sections. The list is alphabetic and is activated using the drop-down arrow within the cell, after the first few letters of the ion is typed.



If a name is misspelled or if the text is unrecognized, then a red 'X' appears to the left of the name. This name needs to be corrected or the row deleted before proceeding. To delete the row, simply select the wrong entry (which will turn black) and hit the key **<Delete>**.



Water Analyses - Reported Elements as Total Ions

Water analysis data obtained from ICP measurements will contain concentrations for B, P, S, and Si. These elements do not exist in the water, rather they exist as dissolved ions. If they are part of your analysis, then you should convert them to the following before entering them into the Water Analyses object. Some of these ions are already entered into the **Total lons** section, and the software will do the conversion automatically.

Converting element concentration to species for Brine or Water Analysis

| ICP data | Aqueous Species | Formula to enter | Formula weight multiplier |
|---------------|----------------------|---------------------------------|--|
| B, boron | Boric Acid | H3BO3 | B (mg/l) × 5.72 = H3BO3 (mg/l) |
| Si, Silicon | Silica | SiO2 | Si (mg/l) × 2.14 = SiO2 (mg/l) |
| P, Phosphorus | Dihydrogen Phosphate | H2PO4-1 | $P (mg/l) \times 3.13 = H2PO4 (mg/l)$ |
| | | HS-1 or SO4-2 | S (mg/l) \times 1.03 for HS-1 (mg/l) |
| S, Sulfur | Sulfate or Sulfide | (Cannot tell from total S only) | or |
| | | | S (mg/l) \times 3.0 for SO4-2 (mg/l) |

A Basic Water Analysis

A brief introduction to the water analysis tool will be shown in the example below. As we go through the example, the basic definitions, functionalities, and reporting for the *Water Analysis tool* will be introduced.

Example 27: Ground Water Analysis

Will calculate the **pH** and **density** of a Ground Water sample based upon its measured composition at 1 atm and 25 °C. Then we will create a molecular stream based on the ionic inflows.

Starting the Simulation

Use the inputs and parameters from the table below to create the water analysis. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Ground Water Analysis | | | | | | | |
|------------------------------|-----------------------|-----------|---------------|--------------------------|---------------|---------------------|--------|
| Analysis Parameters/Settings | | | | Recorded Properties | | | |
| Stream Amount | | 1 L (Def | ault) | ault) Total Dissolved So | | Solids Not recorded | |
| Temperature | | 25 °C | | Measured pH | | 6.7 | |
| Pressure | | 1 atm | | Measured Alkalir | nity | Not reco | orded |
| Name Style | | Display | Formula | Density | | Not reco | orded |
| Unit Set | | Metric, I | Batch, | Specific Electrica | ıl | Not reco | orded |
| | | Concen | tration | Conductivity | | | |
| Framework | | MSE | | | | | |
| Calculation Type | | Water A | nalysis | | | | |
| Analysis Name | Analysis Name Basic W | | ater Analysis | | | | |
| | | | Comp | osition | | | |
| Neutrals | s (mg/L) | | Cations | (mg/L) | Anions (mg/L) | | (mg/L) |
| CO2 | 150 | | Na+1 | 1060 | CI-1 | | 3896 |
| H2S | 15 | | K+1 | 50 | SO4-2 | | 54 |
| NH3 | 5 | | Ca+2 | 773 | | | |
| | | | Mg+2 | 177 | | | |
| | | | Sr+2 | 0.18 | | | |
| | | | Ba+2 | 0.46 | | | |
| | | | Fe+2 | 62.1 | | | |
| | | | Mn+2 | 2.80 | | | |
| | | | Al+3 | 0.74 | | | |

Setting the Water Analysis

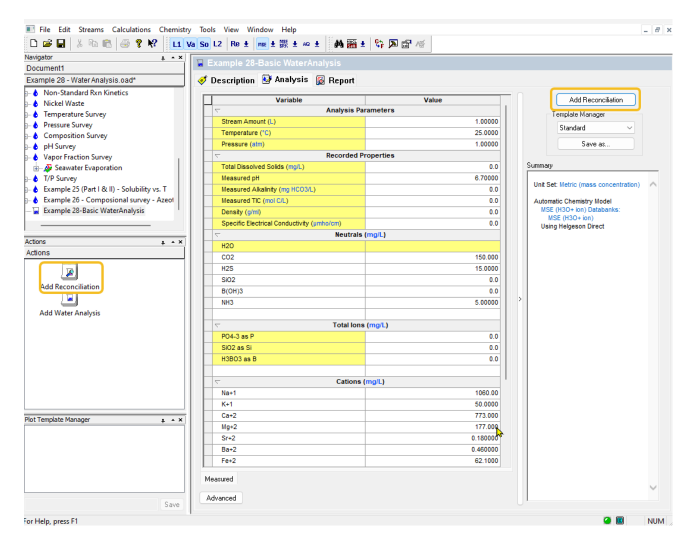
Add a Water Analysis

Click on the new Water Analysis and press **<F2>** to change the name to *Basic Water Analysis*Select the **MSE** thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the **Units Manager** Icon, and select Metric, Batch, Concentration (it may be defined by default)
Under the **Analysis** Tab, enter the Analysis Parameters, Recorded properties, and composition of the water given in the table above.

Go to the **Add Reconciliation** button of the top right corner or select **Add Reconciliation** from the Actions Panel.



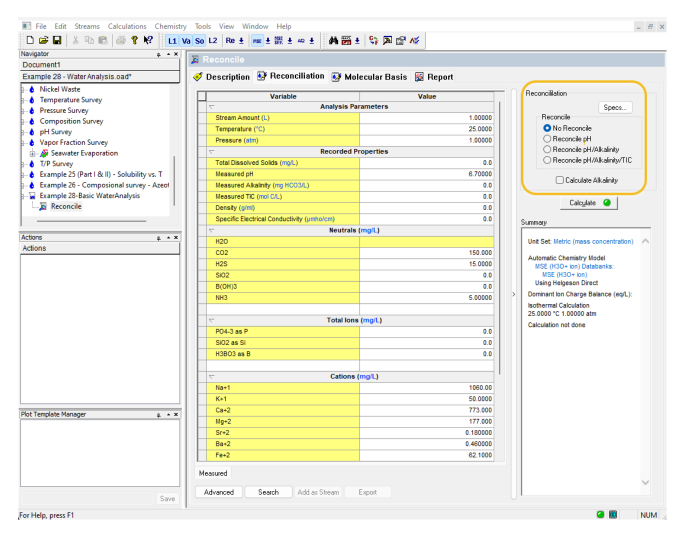
Note: You may notice that after selecting **Add Reconciliation** option, the navigator panel displays a sub-stream called **Reconcile**, and this **Reconcile** sub-stream opens a new tab named **Reconciliation**. The Reconcile sub-stream copies the original inputs entered in the *Water Analysis*. Any changes made in the sub-stream will not change the original Water Analysis.

Additionally, four different types of reconciliation are enabled in the upper right corner of the window: (1) No Reconcile, (2) Reconcile pH, (3) Reconcile pH/Alkalinity, and Reconcile pH/Alkalining/TIC. These types of reconciliation will be described in more detail later in the next section.

The selected default calculation is **No Reconcile**. This option means that the software will compute the water properties based on the current concentration of neutral, cations, and anions species. The calculation will not use the measured pH, or the measured alkalinity entered (if any).

The Calculate Alkalinity box (Calculate Alkalinity) allows you to compute the alkalinity, also based on the concentration entered.

Select the No Reconcile option

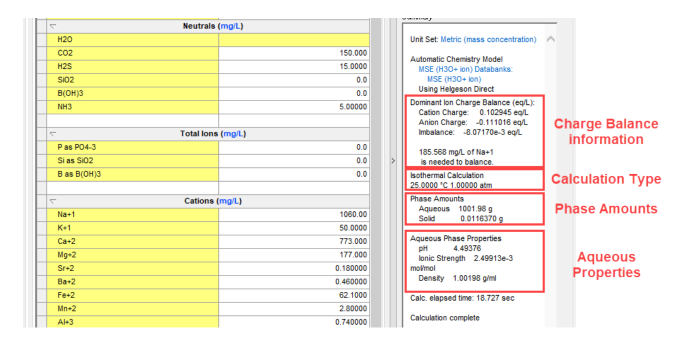


Now, we are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. Create a new file and name it: *Water Analysis Calculations*.

Analyzing the Results

Viewing the Summary Box

Review the Summary Box. Let's analyze it in detail since it contains several pieces of important information, as shown in the image below.



The top section contains the **charge balance information** showing the total equivalent charge (positive and negative), and the charge imbalance. This example has a negative imbalance of -8.072×10⁻³ eq/L, indicating that more cations are needed in solution in order to reach electroneutrality. The software calculates that 185.568 mg/L of Na+ are needed to balance the solution. The software adds this amount of Na+ to the solution. This information can be further confirmed in the **Report Tab** in the Charge Balance table.

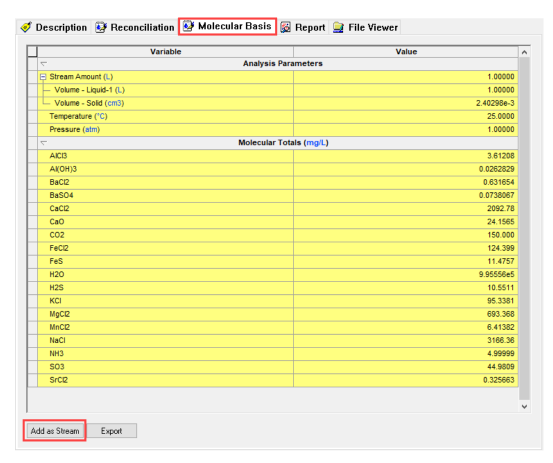
The **calculation type information** shows the calculation type the software used. In this example, the software performed a default isothermal calculation at 25 °C and 1 atm. If we selected other calculation specifications, they would appear here.

The **phase amounts information** shows the distribution of species in the different phases. This analysis contains two phases: aqueous and solid.

The **aqueous properties information** shows the computed pH, ionic strength, and density of the solution. It is important to note here that the measured pH is 6.70 and the computed pH is 4.49. The density of this solution is 1.00198 g/ml. Remember, for this example the software used only the concentration of neutrals, cations, and anions in solution to do a reconciliation.

Viewing the Molecular Basis Tab

Let's review the **Molecular Basis** Tab. The Molecular Basis tab is one of several tabs of the Reconciliation object. The information contained in this tab is the molecular composition of the solution after it has been reconciliated. In other words, the software has converted the **ionic inflows** into a **molecular stream**.



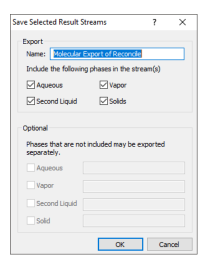
Note: The software generates molecular concentrations based on two priorities. The first priority is to create the least number of molecular inflows. This example contains eighteen inflows (plus H2O not shown). The second priority is to create the least number of total moles. Moles are not shown here, since the units are mg/l. The moles are shown in the Report tab.

You can use this molecular form as a new Stream in OLI Studio, by using the **Add as Stream** button, or it can be exported to a separate software like OLI Flowsheet: ESP, Aspen Plus, UniSim Design, or other simulators.

Converting the Ionic Inflows into a Molecular Stream

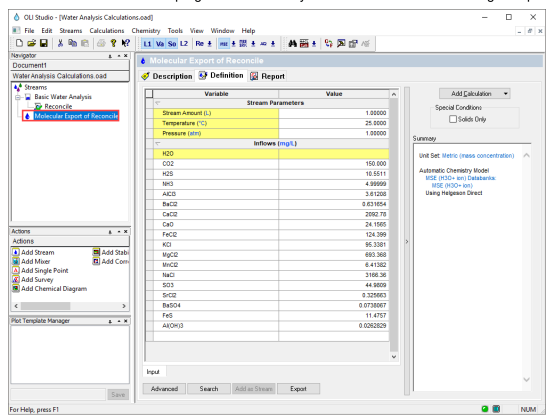
For practice, let's add this Molecular Basis as a Stream into the Navigator Panel.





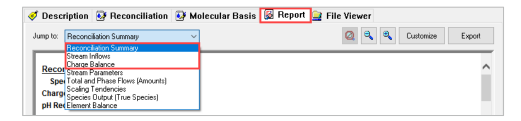
Note: By default, the name of the stream is **Molecular Export of Reconcile**. You can change the name if you prefer. Additionally, you can include or exclude the phases that you want to export into your stream by checking or unchecking the corresponding boxes.

Leave the defaults and click OK. The program automatically adds a new stream in the navigation panel.



Viewing the Report Tab

The **Report** tab is slightly different from the Single Point report described in Section 1. This Report contains three additional tables, all of which are located at the top. These tables are the Reconciliation Summary, Stream Inflows, and Charge Balance.



The **Reconciliation Summary** table confirms the specifications for the reconciliation such as the charge balance method and the reconciliation type (these concepts will be discussed in more detail later in the manual). Additionally, shows the conditions at which the calculation was run, in this case 25°C and 1 atm. And finally, shows a comparison between the measured and calculated properties, in this case it shows the measured vs calculated pH.

Reconciliation Summary Specification Charge Balance Method Dominant Ion pH Reconciliation Type No reconciliation

| | Measured | Calculated |
|-----------------|----------|------------|
| Temperature, °C | 25.0000 | |
| Pressure, atm | 1.00000 | |
| pH | 6.70000 | 4.49376 |
| Density, g/ml | | 1.00198 |
| Water, mg/L | | 9.95555e5 |

The **Stream Inflows** table summarizes all the species that were entered in the water analysis grid.

<u>Stream Inflows</u> Row Filter Applied: Only Non Zero Values

| | Input | Output |
|---------|-----------|-----------|
| Species | mg/L | mg/L |
| H2O | 1.00000e6 | 9.95555e5 |
| Na+1 | 1060.00 | |
| K+1 | 50.0000 | |
| Ca+2 | 773.000 | |
| Mg+2 | 177.000 | |
| Sr+2 | 0.180000 | |
| Ba+2 | 0.460000 | |
| Fe+2 | 62.1000 | |
| CI-1 | 3896.00 | |
| SO4-2 | 54.0000 | |
| CO2 | 150.000 | 150.000 |
| H2S | 15.0000 | 15.0000 |
| NH3 | 5.00000 | 4.99999 |
| Mn+2 | 2.80000 | |
| AI+3 | 0.740000 | |

The **Charge Balance** table contains the concentrations entered and the final balanced values. In this case only the sodium (Na+1) concentration was changed.

Charge Balance

 Cation Charge:
 0.102945 eq/L

 Anion Charge:
 -0.111016 eq/L

 Imbalance:
 -8.07170e-3 eq/L

 Adjusted Species:
 Na+1

| Charged Species | Input | Balanced | Difference |
|-----------------|----------|----------|------------|
| | mg/L | mg/L | % |
| Na+1 | 1060.00 | 1245.57 | 17.51 |
| K+1 | 50.0000 | 50.0000 | |
| Ca+2 | 773.000 | 773.000 | |
| Mg+2 | 177.000 | 177.000 | |
| Sr+2 | 0.180000 | 0.180000 | |
| Ba+2 | 0.460000 | 0.460000 | |
| Fe+2 | 62.1000 | 62.1000 | |
| Mn+2 | 2.80000 | 2.80000 | |
| AI+3 | 0.740000 | 0.740000 | |
| CI-1 | 3896.00 | 3896.00 | |
| SO4-2 | 54.0000 | 54.0000 | |
| HCO3-1 | 0.0 | 0.0 | |
| HS-1 | 0.0 | 0.0 | |
| C2H3O2-1 | 0.0 | 0.0 | |

Exploring Reconciliation Options

When reconciling a Water Analysis, there are three options for reconciliation:

No Reconcile: The software will run an electroneutrality reconciliation only, and then compute the water properties such as pH, density, etc., based on the <u>entered concentration of neutral, cations, and anions species</u>. In the <u>No Reconcile</u> option you may allow the program to pick the species to adjust for **electroneutrality** or you may manually choose the species to perform the adjustment. (<u>See electroneutrality options</u>).

Reconcile pH: The software will run both an <u>electroneutrality</u> and <u>pH reconciliation</u>. This type of reconciliation will match your recorded pH. Additionally, the software will compute the water properties such as, density, electrical conductivity, etc. The **pH** of the solution is automatically **adjusted** by the software by **adding** either **HCI** or **NaOH**, or you may select your preferred acids and bases to adjust the pH.

Reconcile pH/Alkalinity: The software will run an electroneutrality, pH and alkalinity reconciliation. This type of reconciliation will match your recorded pH and alkalinity values. Additionally, the software will compute the water properties such as density, electrical conductivity, etc. The pH of the solution is automatically adjusted by the software by adding either HCl or NaOH or you may select your preferred acids and bases to adjust the pH. The Alkalinity is automatically adjusted by the software, using CO₂ as the alkalinity titrant, H₂SO₄ as the alkalinity pH titrant and 4.5 as the alkalinity end point pH.

Reconcile pH/Alkalinity/TIC: The software will run an electroneutrality, pH, alkalinity and Total Inorganic Carbon (TIC) reconciliation. This type of reconciliation will match your recorded pH, alkalinity and TIC values. Additionally, the software will compute the water properties such as density, electrical conductivity, etc. The TIC is automatically calculated by the software, using CO₂ as the titrant, H₂SO₄ as the alkalinity pH titrant and 4.5 as the end point pH. The (total) Alkalinity is adjusted by the software by adjusting the acetate concentration (organic acids) to match the target Alkalinity. The Alkalinity is adjusted by adding or removing acetic acid. In this calculation, you cannot change the CO₂ or Acetic Acid for the alkalinity adjustment, these are fixed by the software.

Additionally, there is the option to Calculate Alkalinity: Calculate Alkalinity. It is important to note, that this is only an alkalinity calculation based on the concentration entered, it is not an alkalinity reconciliation.

We will explore the different types of reconciliation and introduce the different specification options according to the reconciliation type.

In this section we will create a new file. We will run a basic water analysis, from which the various reconciliation options will be explored.

Example 28: Water Analysis – No Reconcile Option

In this example we will calculate the **pH** and different properties of a Produced Water sample based upon its measured composition at 1 atm and 25 °C.

Starting the Simulation

Use the inputs and parameters from the table below to create the water analysis. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Water Analysis – Reconcile Options | | | | | | | |
|------------------------------------|----------|----------------|-------------|---------------------|---------------|----------|--------|
| Analysis Parameters/Settings | | | | Recorded Properties | | | |
| Stream Amount | | 1 L (Def | ault) | Total Dissolved S | Solids | 36500 n | ng/L |
| Temperature | | 25 °C | | Measured pH | | 7.8 | |
| Pressure | | 1 atm | | Measured Alkalin | nity | 160 mg | HCO3/L |
| Name Style | | Display | Formula | Density | | 1.013 | |
| Unit Set | | Metric, I | Batch, | Specific Electrica | ıl | Not reco | orded |
| | | Concen | tration | Conductivity | | | |
| Framework | | MSE | | | | | |
| Calculation Type | | Water Analysis | | | | | |
| Stream Name | | Water A | nalysis – | | | | |
| | | Reconc | ile Options | | | | |
| | | | Comp | osition | | | |
| Neutrals | s (mg/L) | | Cations | (mg/L) | Anions (mg/L) | | (mg/L) |
| SiO2 | 16 | | Na+1 | 10000 | CI-1 | | 19000 |
| | | | Ca+2 | 500 | SO4-2 | | 2700 |
| | | | Mg+2 | 1200 | HCO3-1 | | 142 |
| | | | Sr+2 | 200 | AsO4-3 | | 12 |
| | | | Ba+2 | 5 | HCOO- | 1 | 20 |
| | | | Fe+2 | 5 | C2H3O2 | 2-1 | 50 |

Calculating the pH

Setting the Water Analysis

Add a Water Analysis

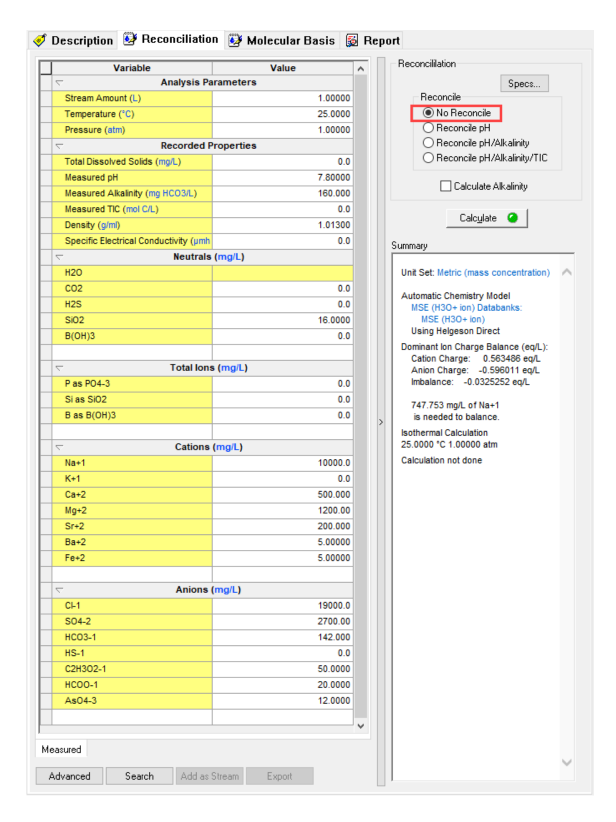
Click on the new Water Analysis and press **<F2>** to change the name to *Water Analysis – Reconcile Options*Select the **MSE** thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the **Units Manager** Icon, and select Metric, Batch, Concentration (it may be defined by default)
Under the **Analysis** Tab, enter the Analysis Parameters, Recorded properties, and Composition of the water given in the table above.

Go to the **Add Reconciliation** button of the top right corner or select **Add Reconciliation** from the Actions Panel, and name it **No Reconcile**.

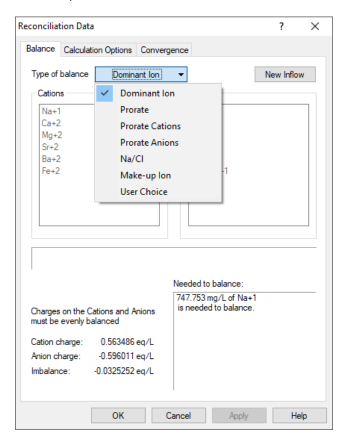
Select the **No Reconcile** option (selected by default)



Note: In the **No reconcile** option, the software runs an electroneutrality reconciliation. The **type of balance** for reconciling electroneutrality is the **Dominant Ion** method. You can select different types of balance for electroneutrality by clicking on the **Specs** button.

Types of Balance for Electroneutrality

Click on the **Specs** button. This will open a new window.



There are 7 different Types of Balance:

Dominant Ion: This is the default method. The largest counter ion is used to adjust the electroneutrality. In our example,747.753 mg/L of Na+1 are added since there is an excess of negative charge (see the above image).

Prorate: This option keeps the relative amount of the counter ions (for this example, the cations) the same and are adjusted up or down equally until reaching electroneutrality.

Prorate Cations: All cations are adjusted up or down equally

Prorate Anions: All anions are adjusted up or down equally

Na+/CI-: Sodium is added when there is an excess of negative charge. Chloride is added when there is an excess of positive charge.

Make-up Ion: This option allows for a single ion species to be adjusted. User selects an ion to increase or decrease.

User Choice: The software determines if an anion or cation is needed to balance the solution and the user choses the specific ion.

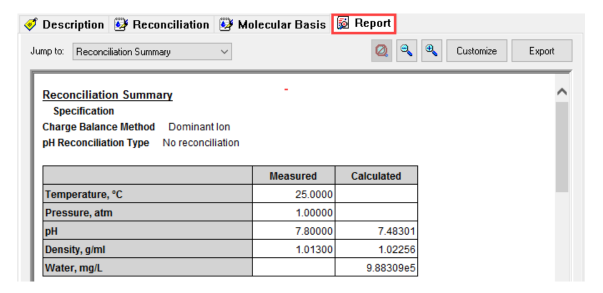
Keep the default option, **Dominant Ion**, as the balance type then press **OK**

Now, we are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named *Water Analysis Calculations*.

Analyzing the Results

Once you run the simulation there are two options to analyze the results: The summary box and the Report

1. Go to the **Report** Tab and look at the **Reconciliation Summary** Table.



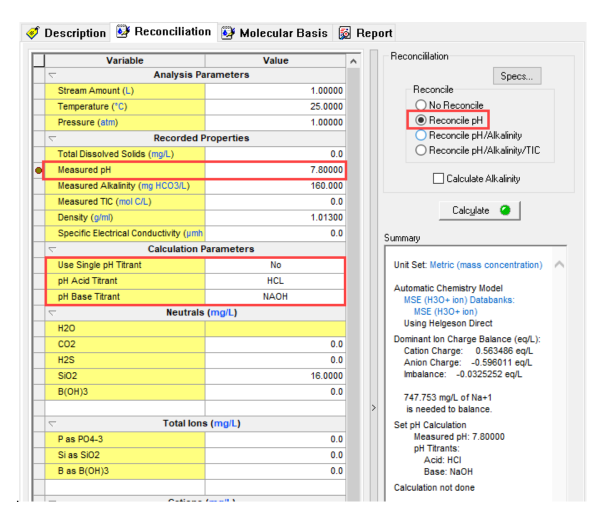
The calculated pH and density are different to the measured values. The software only used the concentration data to calculate these properties.

Example 29: Water Analysis - pH Reconcile Option

In this example we will reconcile the sample by **pH** and also calculate the **alkalinity** of the same Produced Water sample used in the previous Example 28: Water Analysis – No Reconcile Option.

Starting the Simulation

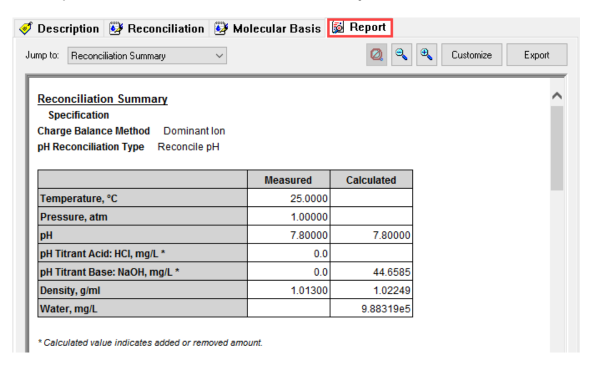
Under the *Water Analysis – Reconcile Options* Stream go to the **Add Reconciliation** button **Click** on the new Reconcile-1 icon and press **<F2>** to change the name to *Reconcile pH*The default units, names and framework have been previously defined
Select the **Reconcile pH** option



Note that by selecting the *Reconcile pH* option, now the measured pH is a fixed value (and brown dot appears in front of this cell). To match the measured pH, the software added a Calculation Parameters section which indicates the acid and basic pH titrants to adjust the pH. By default, HCl and NaOH are selected by the software. This is similar to the set pH single point calculation.

Now, we are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named *Water Analysis Calculations*.

1. Go to the **Report** Tab and look at the **Reconciliation Summary** Table.

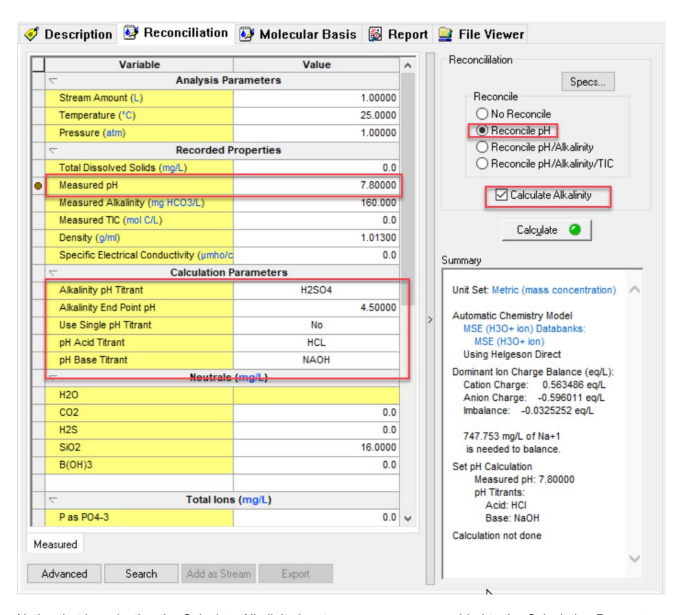


The measured and calculated pH are the same. The software added 44.6585 mg/L of NaOH to match the experimentally measured pH.

Calculating Alkalinity

Now we will instruct the software to calculate Alkalinity.

Go back to the Reconciliation Tab and check the Calculate Alkalinity box.



Notice that by selecting the Calculate Alkalinity box two more rows were added to the Calculation Parameters section: Alkalinity pH Titrant and Alkalinity End Point pH, besides the pH titrants rows. These two new rows indicate that the software will add H₂SO₄ (selected by default) until the pH reduces to 4.5. The software then converts the amount of additional H₂SO₄ into the alkalinity value.

Now, we are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key. It is time to **save** your file (**File >Save as...**) or using the **save** icon in the tool bar.

Analyzing the Results

1. Go to the Report Tab and look at the Reconciliation Summary Table.

Reconciliation Summary

Specification

Charge Balance Method Dominant Ion pH Reconciliation Type Reconcile pH

Alkalinity pH Titrant: H2SO4

| | Measured | Calculated |
|-------------------------------|----------|------------|
| Temperature, °C | 25.0000 | |
| Pressure, atm | 1.00000 | |
| рН | 7.80000 | 7.80000 |
| pH Titrant Acid: HCI, mg/L * | 0.0 | |
| pH Titrant Base: NaOH, mg/L * | 0.0 | 44.6585 |
| Alkalinity, mg HCO3/L | 160.000 | 249.667 |
| Density, g/ml | 1.01300 | 1.02249 |
| Water, mg/L | | 9.88319e5 |

^{*} Calculated value indicates added or removed amount.

The calculated alkalinity value is different to the measured value. This is because the software is not doing an alkalinity reconciliation, rather the software is using only the concentration data to calculate alkalinity. You can see this as single point alkalinity calculation.

Note that in this example, the alkalinity being reported because we checked the **Calculate Alkalinity** box; if the box wasn't checked we would not see the calculated value in the Reconciliation Summary table.

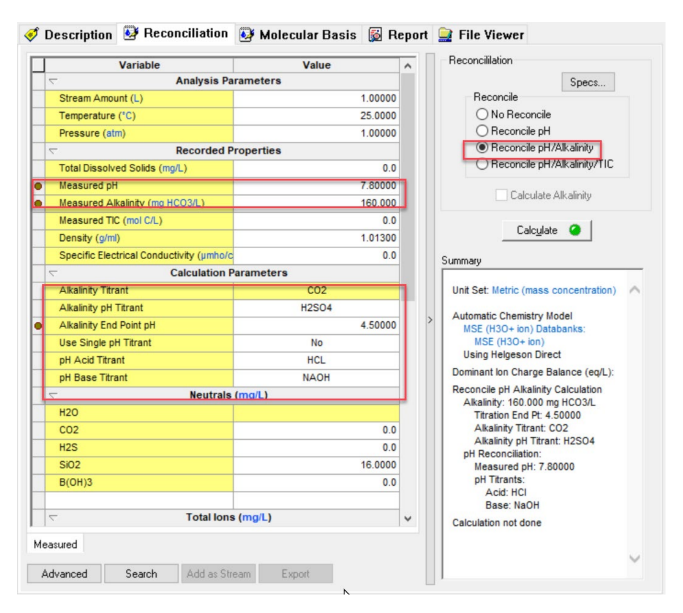
Example 30: Water Analysis – Reconcile pH/Alkalinity Option

In this example we will reconcile the sample by **pH and alkalinity** and also calculate the **pre-scaling tendencies** of the same Produced Water sample used in the previous Example 28: Water Analysis – No Reconcile Option.

Starting the Simulation

Under the *Water Analysis – Reconcile Options* Stream go to the **Add Reconciliation** button **Click** on the new Reconcile-1 icon and press **<F2>** to change the name to *Reconcile pH/Alkalinity*The default units, names and framework have been previously defined

Select the **Reconcile pH/Alkalinity** option



Note that by selecting the *Reconcile pH/Alkalinity* option, now both the measured pH and alkalinity values are fixed values (and brown dot appears in front of this cell).

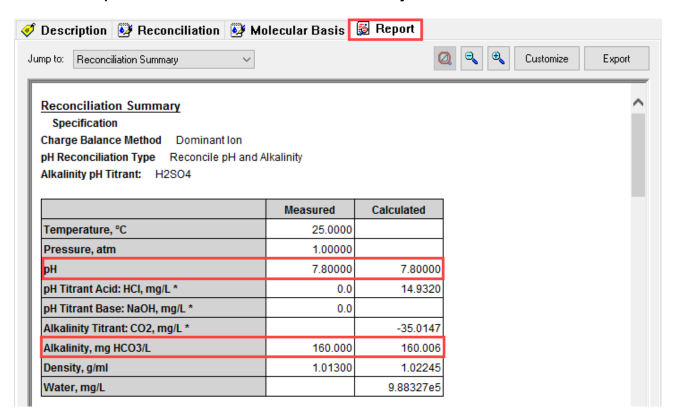
To match the measured alkalinity value, the software added a **Calculation Parameters** section which indicates the alkalinity titrant and the pH titrant used to adjust the alkalinity. CO2 is used by default as the alkalinity titrant. This is similar to an *alkalinity* single point calculation.

To match the measured pH, the software also added a **Calculation Parameters** section which indicates the acid and basic pH titrants to adjust the pH. By default, HCl and NaOH are selected by the software. This is similar to the *set pH* single point calculation. Also notice that the alkalinity box is grayed out. This is because we have asked the software to match the measured alkalinity value. This means that calculated alkalinity value is no longer needed.

Now, we are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named *Water Analysis Calculations*.

Analyzing the Results

1. Go to the **Report** Tab and look at the **Reconciliation Summary** Table.



The measured and calculated pH and alkalinity values are the same. The software added 14.9320 mg/L of HCl to match the experimentally measured pH and removed 35.0147 mg/L of CO2 to match the experimentally measured alkalinity value.

Calculating Pre-scaling Tendencies

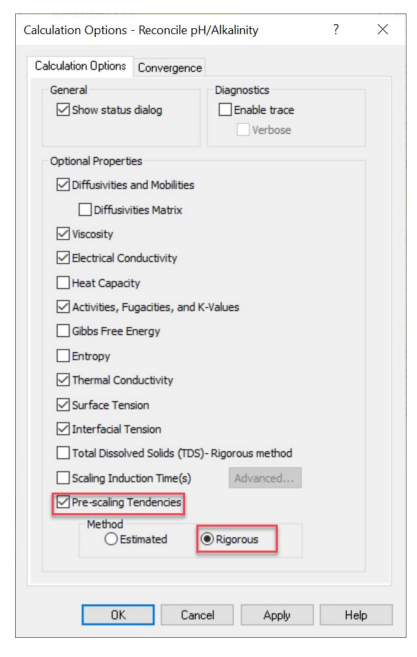
Now we will instruct the software to calculate Pre-scaling tendencies. In OLI Studio: Stream Analyzer, Pre-scaling tendencies are turned OFF by default. We need to enable this option.

Setting the Water Analysis

Go back to the **Reconciliation** tab of the *Reconcile pH/Alkalinity* water analysis

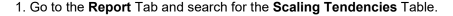
Click on the **Calculation Options** icon . This will open the Calculation Options window.

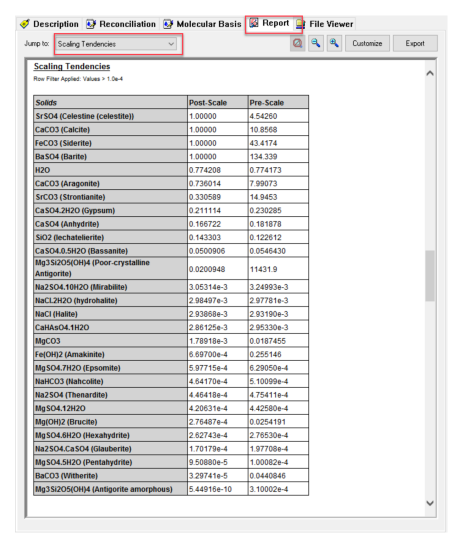
Check the Pre-scaling Tendencies box and select the Rigorous method. Then click OK.



Now, we are ready to perform the calculation. Click on the Calculate button or press the <F9> key

It is time to **save** your file (**File >Save as...) or** using the **save** icon in the tool bar.





This table reports both the Post-Scale and Pre-scale Tendencies. See the section 3.1 Basic Terminology to see a more detailed explanation.

The Scaling Tendencies (reported in the software as **Post-Scale**), are essentially saturation ratios. When the scaling tendency for a given mineral is equal to 1.0 (saturation conditions) it indicates that the mineral is in equilibrium with water, a solid phase has formed. A scaling tendency below 1.0 indicates sub-saturation and that the solid phase will not form. A solid is at risk of forming when the scale tendency value is greater than 1.0 (supersaturation conditions).

The Pre-Scaling Tendencies (reported in the software as **Pre-Scale**), is defined as the scaling tendency before any solids are formed (this can be seen as all the species suspended in solution).

Note: The Post-Scale and the Pre-Scale tendencies are equivalent if the solid phase is turned OFF.

Section 4. Hydrocarbons and Pseudocomponents

So far, we have learned how to enter molecular and ionic inflows in Stream Analyzer. Stream Analyzer also allows you to enter assays or pseudocomponents in order to enter crude oils as streams.

Crude oils are complex groups of organic molecules containing hundreds, perhaps thousands, of pure components in a single oil. Modeling crude oils using pure components is impractical because analyzing for each pure component is cost-prohibitive and the number of species would make calculations overwhelming. A convenient solution to this problem and to modeling the properties of a crude oil is to create pseudocomponents.

Crude oil properties may be defined through a distillation curve, where each boiling point range is a progression of molecular weights, densities, solubilities, viscosities, and other properties associated with that section. It is reasonable for low boiling point molecules to be low molecular weight, low density, low viscosity, and more soluble in water. We can dice boiling point curves using well accepted methods standard to create pseudocomponents that in combination reflect the property of the whole oil.

There are two ways to create a crude oil stream on Stream Analyzer:

The first is to start with a PVT curve and create pseudocomponents using one of the three thermodynamic methods coded into the software.

The second is to enter the pseudocomponent data directly and using the same thermodynamic methods to predict the component properties.

The three thermodynamic methods are: API, Lee Kesler, and Cavett. More details about this method can be found in Section 11. .

A brief introduction of how to enter PVT curves and/or pseudocomponents will be shown in the examples below.

Entering a PVT Analysis

Example 31: Creating an Assay

In this example, we are going to learn how to enter a simple PVT analysis in order to create a Crude Oil Stream.

Starting the Simulation

Add a new Stream

Click on the new Stream and press <F2> to change the name to Crude Oil

Select the MSE-SRK thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

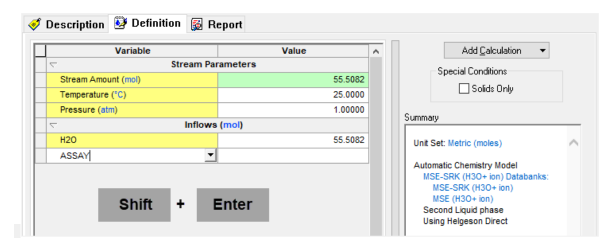
Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the conditions of the Stream T= 25 °C and 1 atm

| Crude Oil Stream | | | | | |
|---------------------------------|-----------------|---------------|---------|--|--|
| Calculation Settings Conditions | | | | | |
| Stream Name | Crude Oil | Stream Amount | Default | | |
| Name Style | Display Formula | Temperature | 25 °C | | |
| Unit Set | Metric, Moles | Pressure | 1 atm | | |
| Framework | MSE-SRK | | | | |

In the inflows grid, in the white cell below H2O, type ASSAY and then press **Shift + Enter>.** You can assign another name to your Assay; however, you are limited to 5 characters for the name of the Assay.

Note: The <Shift + Enter> action instructs the software to create an Assay rather than look for a species or pure component from the database.

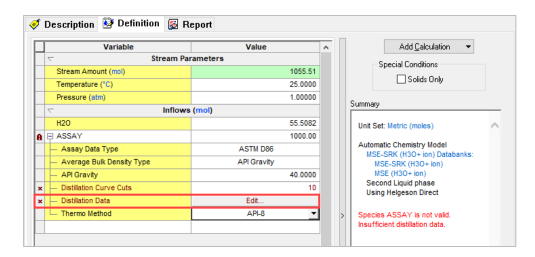


Note: After typing the name of your Assay, immediately press Shift + Enter command keys together.

Use the information provided in the table below to complete the grid.

| Assay Information | | | |
|---------------------------|----------------------|-------------------|-----------------|
| Calculation Settings | | Distillation Data | |
| Assay amount | 1000 moles | Volume % | Temperature, ⁰C |
| Assay Data Type | ASTM D86 | 1 | 20 |
| Average Bulk Density Type | API Gravity | 5 | 30 |
| API Gravity | 40 | 10 | 50 |
| Distillation Curve Cuts | 10 | 20 | 60 |
| Distillation Data | See next two columns | 40 | 80 |
| Thermo Method | API-8 | 60 | 120 |
| | | 80 | 150 |
| | | 90 | 180 |
| | | 95 | 200 |
| | | 99 | 220 |
| | | 100 | 240 |

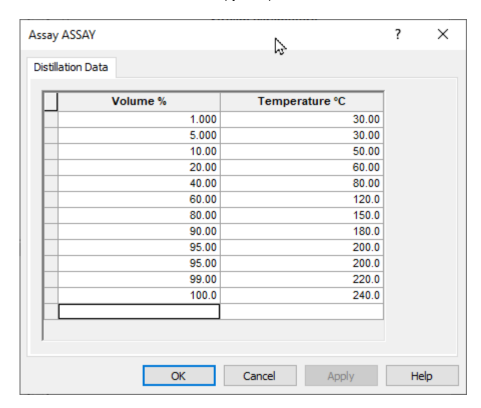
Your screen should look like the image below:



Note: You can change the Assay Data Type, the Average Bulk Density Type, and the Thermo Method options by clicking on the white cells. This will enable a drop-down arrow that will show you the different options.

Click on the **Edit** button next to the Distillation Data. This will open a new window. Complete the Distillation Data using the information given in the table above. Then click **OK**.

Note: Ctrl+C and then Ctrl+V can be used to copy and paste the data in the Distillation Data table.



The Distillation Curve Cuts will eventually become individual pseudocomponents, each with its own critical parameters and thermodynamic reference data.

At this point we are ready to perform a calculation. **Add** a **Single Point** calculation, and then select an **Isothermal** calculation.

Change the SinglePoint name to Assay using the <F2> key

Click on the Calculate button or press the <F9> key to run the simulation

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. Save the calculations of this section under the name *Hydrocarbons and Pseudocomponents*.

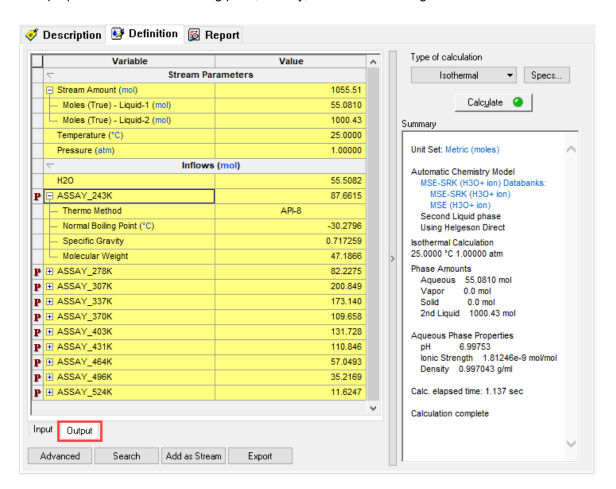
Analyzing the Results

We will review the results first in the Output Tab and then in the Report Tab.

Click on the Output Tab

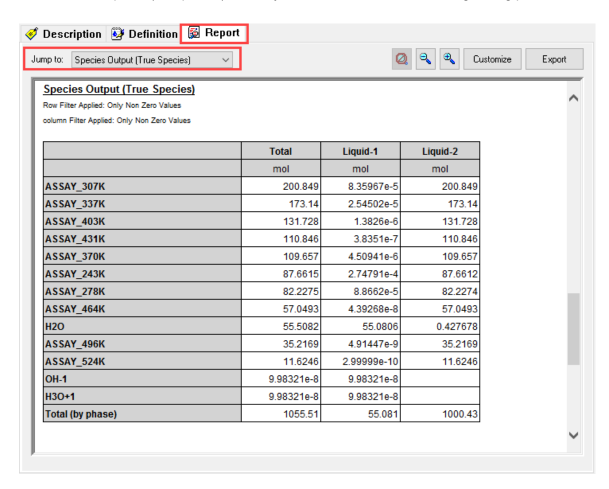
The Output Tab shows a grid with 10 different pseudocomponents, identified with the letter P, corresponding to each distillation cut of the crude. Notice that the name of each cut is the combination of the oil name (ASSAY for this example) plus its boiling point in K.

The '+' sign adjacent to each Assay expands a sub-table that contains the critical properties of each cut. These properties are: Normal boiling point, density, and molecular weight.



Click on the Report Tab and scroll down to the Species Output table.

This table shows how the pseudocomponents are distributed between the Liquid-1 and Liquid-2 phases. Notice how the Liquid-1 (or aqueous) solubility decreases with each increasing boiling point.



Entering Pseudocomponents to a Stream

There are times when the user does not have (or does not want to use) distillation data. Rather, the user had the individual properties for a single pseudocomponent. OLI allows the user to enter the individual pseudocomponents. In the following example, we will illustrate how.

Example 32: Creating Pseudocomponents

In this example, we will enter two properties - boiling point and density. The software will use one of four thermodynamic models (API-8, API-5, Cavett, or Kessler-Lee) to calculate critical properties of pseudocomponents.

Starting the Simulation

Add a new Stream

Click on the new Stream and press <F2> to change the name to Crude Oil - Pseudocomponents

Select the MSE-SRK thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Moles

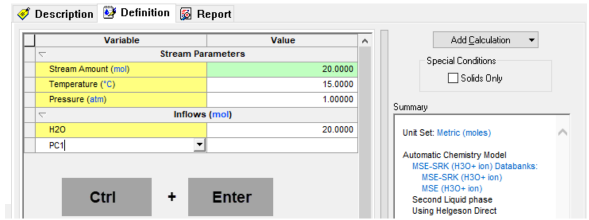
Enter the conditions of the Stream T= 15 °C and 1 atm

Change H2O from the default value to 20 moles

| Crude Oil Stream | | | |
|----------------------|------------------------------|---------------|------------|
| Calculation Settings | | Conditions | |
| Stream Name | Crude Oil - Pseudocomponents | Stream Amount | Calculated |
| Name Style | Display Formula | Temperature | 15 ℃ |
| Unit Set | Metric, Moles | Pressure | 1 atm |
| Framework | MSE-SRK | H2O | 20 moles |

In the inflows grid, in the white cell below H2O, type PC1 and then press <CrtI + Enter>

Note: The <Crtl + Enter> action instructs the software to create a pseudocomponent rather than look for a species or pure component from the database.

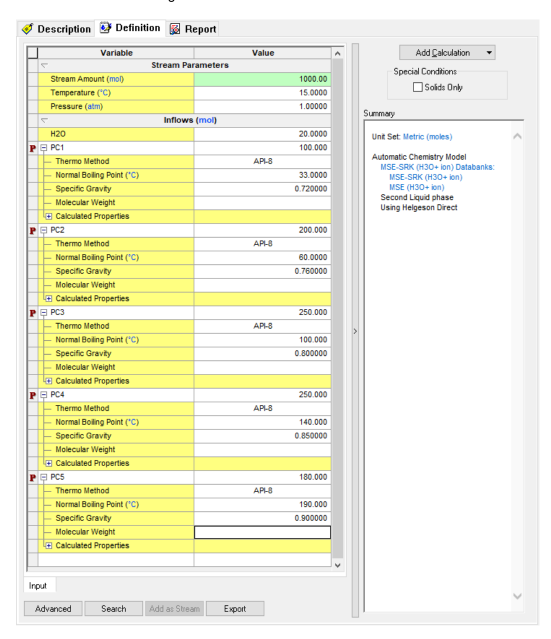


Note: After typing the name of your pseudocomponent, immediately press Ctrl + Enter command keys together.

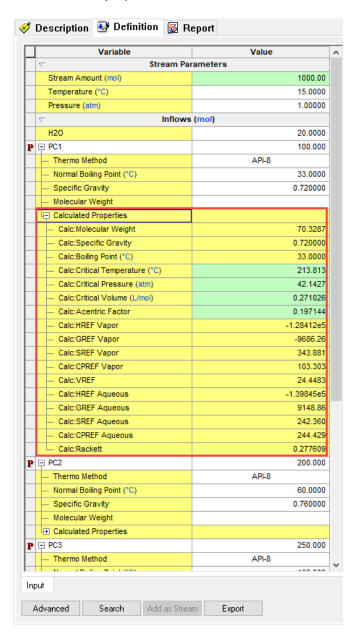
Repeat step 8 four more times with the following names: PC2, PC3, PC4, and PC5. Use the information provided in the table below to complete the grid. The molecular weight will be calculated by the software. In general, only two of the three parameters need be entered.

| Pseudocomponents Information | | | | | |
|------------------------------|-------|-------|-------|-------|-------|
| Pseudocomponent name | PC1 | PC2 | PC3 | PC4 | PC5 |
| Inflow (moles) | 100 | 200 | 250 | 250 | 180 |
| Thermo Method | API-8 | API-8 | API-8 | API-8 | API-8 |
| Boiling Point (°C) | 33 | 60 | 100 | 140 | 190 |
| Specific Gravity | 0.72 | 0.76 | 0.8 | 0.85 | 0.9 |

Your screen should look like the image below:



Notice that at the bottom of each pseudocomponent there is a "+" and a field called "Calculated Properties." Stream Analyzer immediately predicts all the calculated properties for each pseudocomponent. Click the "+" sign next to PC1 to expand and see these properties.



The values in the green boxes can be edited to match data from other programs.

At this point we are ready to perform a calculation. **Add** a **Single Point** calculation, and then select an **Isothermal** calculation.

Change the **SinglePoint** name to **Pseudocomponents** using the **<F2>** key

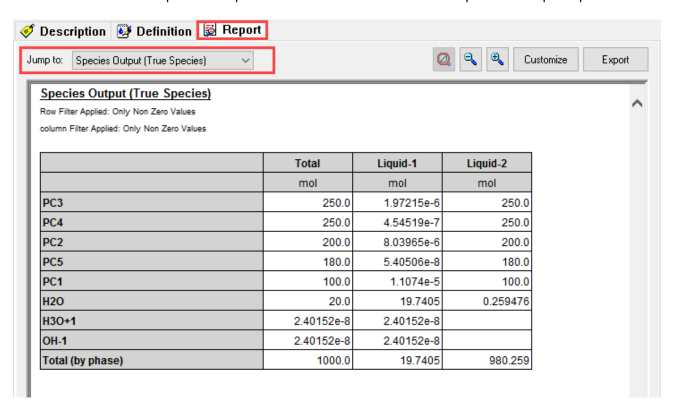
Click on the Calculate button or press the <F9> key to run the simulation

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. Save the calculations of this section under the name *Hydrocarbons and Pseudocomponents*.

Analyzing the Results

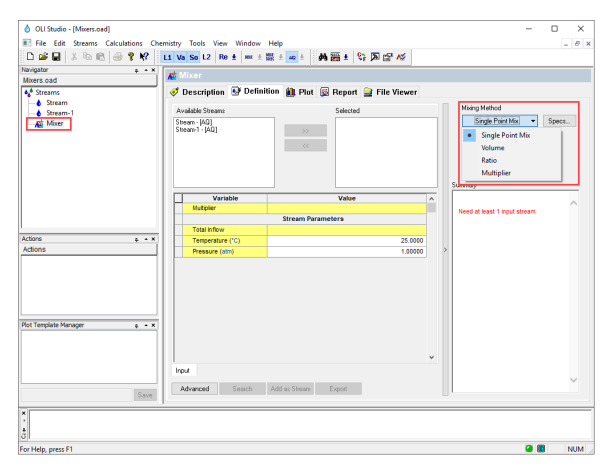
Click on the Report Tab and scroll down to the Species Output table.

This table shows how the pseudocomponents are distributed between the Liquid-1 and Liquid-2 phases.



Section 5. Mixers

Mixers are a useful tool to mix one or more streams. This tool allows you to mix Molecular Streams or Reconciled Streams.



In this section, you will learn how to set up a **Mixer** and differentiate the different mixing options to get the most out of your simulation results.

Mixing Options (Definitions)

The Mixer Object has four different mixing options: Single point mix, Volume, Ratio, and Multiplier. A quick summary of what each mixing option allows you to do is summarized here.

Single Point Mix

This option allows you to multiply up or down a stream inflow. When selecting the streams to mix, this option will enable the multiplier option for both streams. You can leave the default values that are set by default to 1.0 or change the multiplier values to any other number.

Volume

In this option, one stream stays at a constant volume, while the other stream's volume changes within a range specified by the user, as is illustrated in the table below:

| Stream 1 | Stream 2 |
|----------|----------|
| 1 L | 1 L |
| 1 L | 2 L |
| | |
| 1 L | n L |

Ratio

In the ratio mixing type, both streams change at the same time. For example, stream will be 100% and Stream will be 0%, then Stream 1 will be 90% and Stream 2 will be 10%, and so on, until it reaches Stream 1 0% and Stream 2 100%.

| Stream 1 | Stream 2 |
|----------|----------|
| 100 % | 0 % |
| 90 % | 10% |
| | |
| 0 % | 100 % |

Multiplier

In this mixing option, one stream stays constant, while the other stream changes using a multiplier. The range of the multiplier values is given by the user.

| Stream 1 | Stream 2 |
|----------|--------------|
| 1 | Stream 2 × 0 |
| 1 | Stream 2 × 1 |
| | |
| 1 | Stream 2 × n |

A Basic Mixer Calculation

A brief introduction to the Mixer tool will be shown in the example below. As we go through the example, basic definitions, functionalities, and reporting for the *Mixer tool* will be introduced.

Example 33: Calculating the Heat of Mixing

You will mix 1 L of a 10 wt% Ca(OH)₂ solution with 1 L of a 10% HCl solution, and then you will ask the software to calculate the **heat of mixing** and **the final temperature** of the mixture.

Starting the Simulation

In this example, we need to create two different streams: a 10 wt% Ca(OH)₂ solution and a 10% HCl solution.

Use the inputs and parameters from the table below to create each stream. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Basic Mixing Calculation | | | |
|--------------------------|---------------------------|------------------------|---------------------------|
| 1 st Stream | | 2 nd Stream | |
| Stream Name | Ca(OH)2 | Stream Name | HCI |
| Name Style | Display Formula | Name Style | Display Formula |
| Unit Set | Metric, Batch, Mass Frac. | Unit Set | Metric, Batch, Mass Frac. |
| Framework | MSE | Framework | MSE |
| Stream Amount | 1 kg | Stream Amount | 1 kg |
| Temperature | 25 °C | Temperature | 25 ℃ |
| Pressure | 1 atm | Pressure | 1 atm |
| H2O | Calculated | H2O | Calculated |
| Ca(OH)2 | 10 mass% | HCI | 10 mass% |

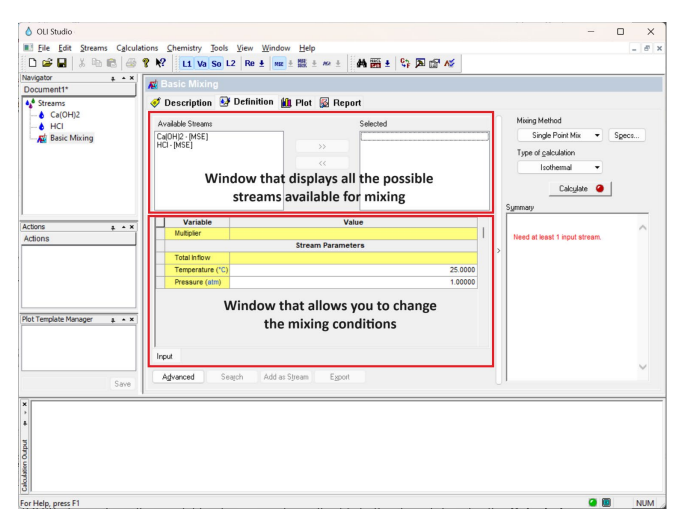
After creating the streams, **Add a Mixer.** The Mixer object can be accessed from the Menu Bar by selecting Calculations > **Add Mixer** or by selecting the **Add Mixer** in the Actions Pane.

Press <F2> to rename the mixer to Basic Mixing

Notice that the display layout for the Mixer tool is slightly different. There are two different windows (see image below).

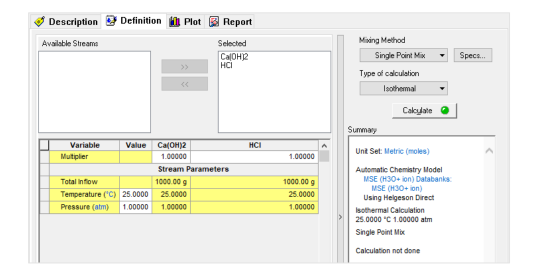
The first window displays all possible streams available for the mix calculation. There is no limit to the number of streams to be mixed.

The second window allows you to change the mixing conditions, e.g., temperature, and pressure of the mixing.



Select the Ca(OH)2 stream from the available streams and use the >> button to put it under the **Selected** window

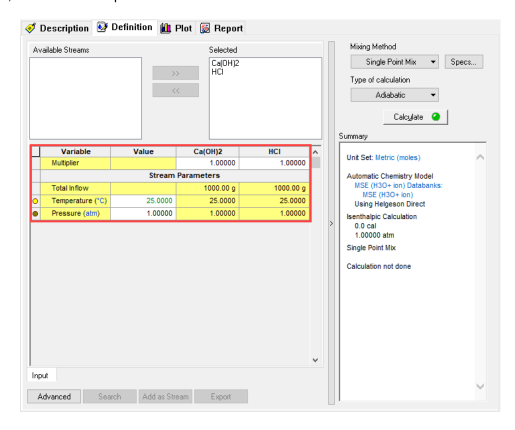
Select the HCl stream from the available streams and use the >> button to put it under the Selected window



Also notice that the default calculation is the **Single Point Mix** Method at **Isothermal** conditions. In this example, we want to calculate the heat of mixing and the final temperature of the mixture. Thus, an adiabatic calculation is required to see this effect.

Click on the **Type of Calculation** button and select the **Adiabatic** calculation option

Note: Once the streams for the mixing have been selected, notice that the multipliers of both streamsCa(OH)2 and HCl can be edited. This specific calculation will be carried out at a constant pressure of 1 atm, and the final temperature of the mixture will be calculated.



Now, we are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file **(File >Save as...) or** use the **save** icon in the tool bar. You can save it and name it as *Mixer Calculations*.

Analyzing the Results

Review the Summary Box. After mixing these two streams, the temperature increased from 25 °C to approximately ~48.65 °C at a constant pressure of 1 atm. This indicates that the enthalpy (or heat) of mixing is exothermic.

A Titration Experiment

In this example, you will explore the volume mixing option to recreate a titration experiment. As we go through the example, different functionalities, reporting, and plotting for the *Mixer tool* will be introduced.

Example 34: HF Titration with CaCl₂

In this example, we will first create two new streams, a 0.1 m HF solution, and a 0.1 m CaCl₂ solution. What is the pH of these individual streams?

Second, we will mix these two streams in equal amounts. What is the final pH of the mixture?

And finally, the 0.1 m HF solution will be titrated with a 0.1 m CaCl₂ solution. What will the titration curve look like?

Starting the Simulation

For this example, we need to create two different streams: a 0.1 m HF solution and a 0.1 m CaCl₂ solution. Note that we have switched thermodynamic frameworks to the **AQ** framework. Please make sure that this framework is selected for both streams in this example.

Use the inputs and parameters from the table below to create each stream. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Titration Calculation | | | |
|------------------------|----------------------|------------------------|----------------------|
| 1 st Stream | | 2 nd Stream | |
| Stream Name | 0.1 HF | Stream Name | 0.1 CaCl2 |
| Name Style | Display Formula | Name Style | Display Formula |
| Unit Set | Metric, Batch, Moles | Unit Set | Metric, Batch, Moles |
| Framework | AQ | Framework | AQ |
| Stream Amount | Calculated | Stream Amount | Calculated |
| Temperature | 30 °C | Temperature | 30 °C |
| Pressure | 1 atm | Pressure | 1 atm |
| H2O | 55.5082 moles | H2O | 55.5082 moles |
| HF | 0.1 moles | CaCl2 | 0.1 moles |

Calculating the pH of the individual streams

Add a Single Point - Isothermal calculation for each stream

Change the name to pH 0.1 m HF and pH 0.1 m CaCl2, respectively, using the <F2> key

Calculate the isothermal calculation for each stream

Check the pH values of the individual streams in the Summary Box

Results - pH of the individual streams

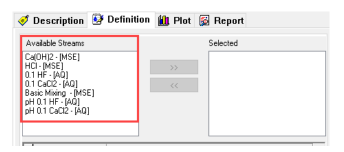
| pH 0.1 m HF | pH 0.1 m CaCl2 |
|-------------|----------------|
| 2.13 | 6.82 |

Calculating the pH of the Mixture

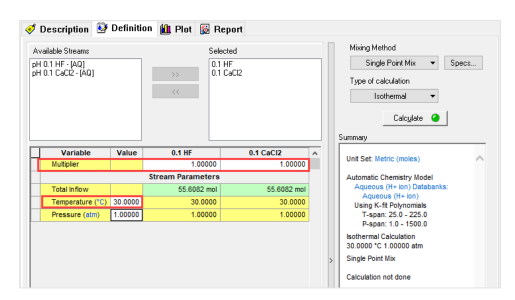
After creating the streams, **Add a Mixer**. The Mixer object can be accessed from the Menu Bar by selecting Calculations > **Add Mixer** or by selecting the **Add Mixer** in the Actions Pane.

Press <F2> to rename the mixer to *Titration*

Note: Notice that the Available Streams window is showing all the available streams for the mixing calculation. Also notice that this window is showing both *Streams* and *Single Point calculations*. Additionally, the available streams' thermodynamic framework appears within brackets, e.g., AQ, MSE, or MSE-SRK. This is important because in order to carry out mixing calculations, the selected streams need to use the same thermodynamic framework.



Select the 0.1 HF – [AQ] stream. Use the >> button to move it to the Selected window
Select the 0.1 CaCl2 stream. Use the >> button to move it to the Selected window
Select Single Point Mix as the Mixing Method and Isothermal as the Type of Calculation
Leave the default values for the multipliers (1.0) and change the temperature at which the mixture takes place to 30 °C.



Now, we are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key At this point is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save under the same file that we created before named as *Mixer Calculations*.

Check the pH values of the individual streams in the Summary Box

Results - pH of the individual streams

The resultant pH is 1.44. How can this be? An acid stream (the HF stream had a pH of approximately 2.13) and a nominally basic stream (the CaCl₂ stream had a pH of 6.82) are mixed, and the pH is outside the value of either stream.

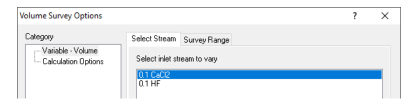
We will continue using Stream Analyzer to further examine the chemistry in more detail. We will titrate the HF stream with CaCl₂.

Titrating HF with CaCl₂

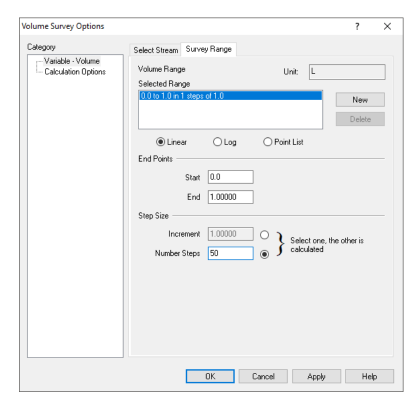
Change the Single Point Mix as the Mixing Method to the Volume option.

Note: You will receive a warning message: Need 1 stream selected as the variable stream (Specs...)

Click on the **Specs** button. This will open a new window indicating you to select the Stream to vary. In the **Select Stream** tab, **Select** the 0.1 m CaCl2 stream



Select the **Survey Range** tab. Change the Volume Range from 0 L to 1 L. Change the **Number of Steps** to 50. Then click **OK**.



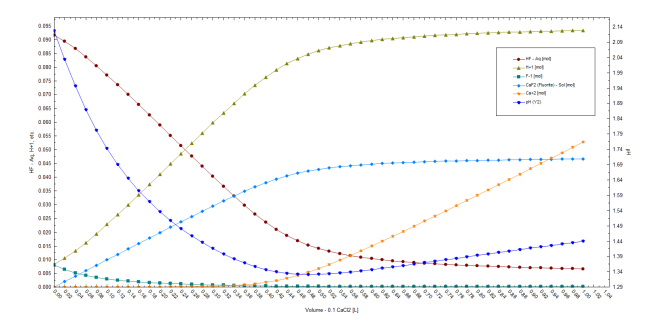
Now, we are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar. You can save under the same file that we created before named as *Mixer Calculations*.

Titration Results and Plotting

Click on the **Plot** tab. **Note:** You will see an empty plot, since we need to define the variables that we want to report.

Click on the Variables button and expand the Aqueous section. Select and put the following species in the Y1 axis using the >> button: HF-Aq, H+1, F-1, Ca+2. Expand the Solid section and select CaF2(s).

Next, expand the **Additional Parameters** Section and Select **pH**. Put it in the **Y2 Axis** using the **>>** button. Then click **OK**.



As you can see, adding $CaCl_2$ to this solution lowers the pH to values as low as 1.33. The pH does begin to slowly increase after 0.48 L of $CaCl_2$ have been added.

Why the unusual pH behavior? You can see that a small amount of Ca^{+2} in solution, causes the formation of the solid CaF_2 . This effectively removes F^- from solution according to the following equation:

$$Ca^{+2} + 2F^- \rightarrow CaF_{2(s)}$$

As we add more $CaCl_2$ in solution, more CaF_2 is formed. As a result, the molecular HF decreases because it dissociates in order to maintain the equilibrium by producing more F^- ions. This shifts the following equilibrium to the right:

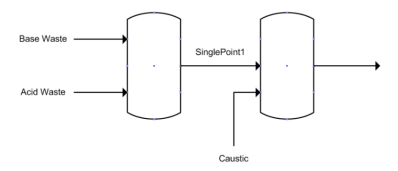
$$HF \rightarrow H^+ + F^-$$

This causes an increase of H^+ in solution and, as a result, a decrease in the pH. At some point, at around 0.48 L of 0.1 m CaCl₂, the solid reaches a steady value, indicating that the solid has reached its saturation (maximum formation) value.

Cascading Mixers

Example 35: Cascading Mixers

In this example, we will mix two streams, one basic and one acidic, and mix them together under adiabatic conditions. The output of this mixer will then be used as the input to a second mixer where a new caustic stream is added. The schematic in the figure below illustrates the layout.

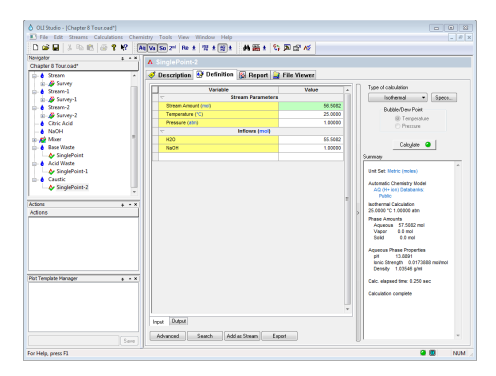


For this example, we need to create three streams. You have already done this in other examples, so we will only give you the input data (for all streams, we will use the AQ thermodynamic framework)

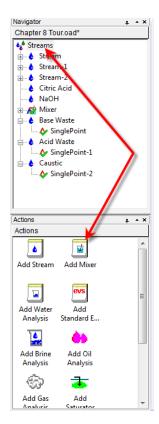
Cascading Mixer Stream Data

| Parameter | Value | Units |
|------------------|-------------------|-------------|
| Stream Name | Base Waste | |
| Temperature | 25.0 | °C |
| Pressure | 1.0 | Atmospheres |
| Stream Amount | 55.5082 (Default) | Mole |
| H ₂ O | 55.5082 | Mole |
| CO2 | 0.1 | Mole |
| NH3 | 0.01 | Mole |
| SO2 | 0.01 | Mole |
| Stream Name | Acid Waste | |
| Temperature | 25.0 | °C |
| Pressure | 1.0 | Atmospheres |
| H ₂ O | 55.5082 | Mole |
| HCI | 0.1 | Mole |
| H2SO4 | 1.0 | Mole |
| Stream Name | Caustic | |
| Temperature | 25.0 | °C |
| Pressure | 1.0 | Atmospheres |
| H ₂ O | 55.5082 | Mole |
| NaOH | 1.0 | Mole |

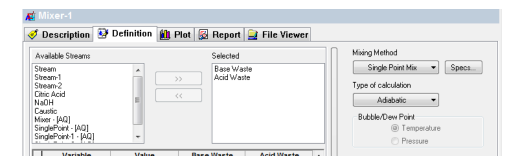
To test that our streams are representing their respective names, perform a single point isothermal flash on each stream. When you are done you should have a window that looks like the image below.



Now we are ready to begin. Click on the "Streams" at the top of the tree-view in the left-hand window, and then select Add Mixer from the actions panel.

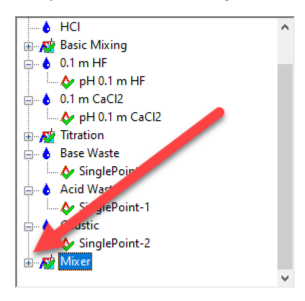


We can now add the "Base Waste" and "Acid Waste" stream as we have done in previous examples. The only difference here is that we are selecting an **Adiabatic** calculation instead of the default isothermal calculation.

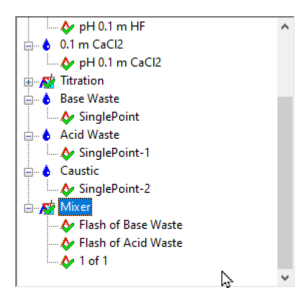


Let's calculate this mixer. Click the Calculate button. If everything is set up correctly the resultant mixed pH should be approximately 0.3.

In the stream tree-view panel, locate your mixer and click the "+" sign.

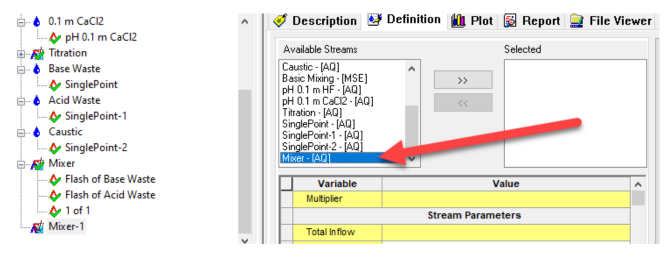


Click the "+" sign to expand the tree.



You can see that the program has performed an equilibrium calculation on each of our input streams as well as calculated the mixer. The results of the mixer are stored in the object "1 of 1".

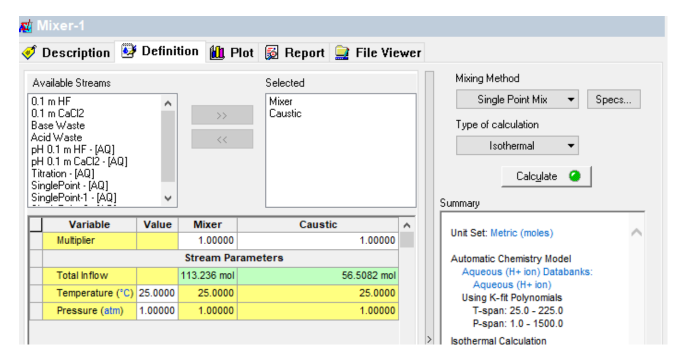
We will now add a second mixer from the actions panel.



When we add a subsequent mixer, we will see all our previous objects still displayed. It is possible to reuse a stream that we already used (this is not possible in OLI's other simulators such as OLI Flowsheet: ESP except by using a virtual stream.) We want to connect the output from our first mixer (Mixer) to the inlet of the <u>Caustic</u> stream. Click the object

Mixer - [AQ].

Then add the **Caustic** stream. Select the type of calculation as Isothermal.



We are now ready to calculate the second mixer. Press the calculate button.

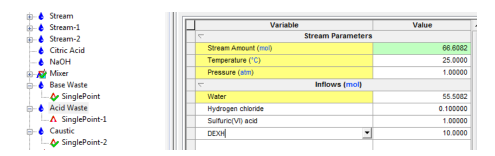
The resultant pH should be approximately 1.0. We want to increase this value. Change the <u>Multiplier</u> value for Caustic Stream from 1.0 to 2.4

| Variable | Value | Mixer | Caustic |
|------------------|----------|-------------|-------------|
| Multiplier | | 1.00000 | 2.40000 |
| | Stream P | arameters | |
| Total Inflow | | 113.236 mol | 135.620 mol |
| Temperature (°C) | 25.0000 | 25.0000 | 25.0000 |
| Pressure (atm) | 1.00000 | 1.00000 | 1.00000 |

Click the calculate button again. The new pH should be approximately 12.1.

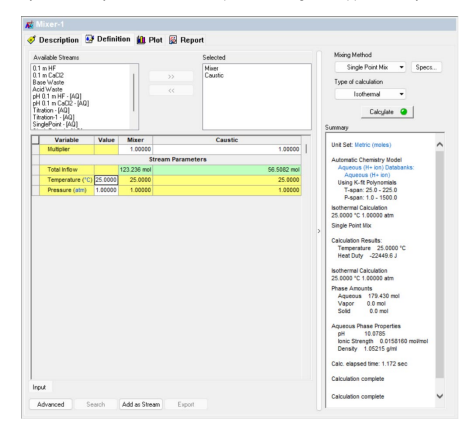
The real value of the cascading mixer is the quick ability to re-calculate the objects if we want to change something. Let's add some diethanolamine to the series of calculations. Click the **Acid Waste** stream (not the single point calculation below it) and add the component DEXH ¹⁰ to the grid with a value of 10.0 moles.

 $^{^{10}}$ This is the OLI Tag name for diethanolamine, which is easier to type if you know the name.



Return to the last mixer you created (Mixer-1 in our example) and then reduce the *Caustic Multiplier back to* 1.0

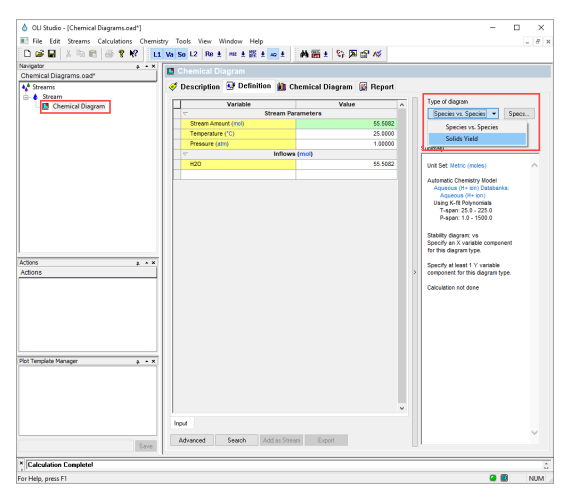
To recalculate all the objects at once, press the control+F9 keys and all objects will be calculated. Now if you look at the summary of Mixer-1 you will see that the pH has changed to approximately 10.



Section 6. Chemical Stability Diagrams

"Predicting the stability of chemical compounds as a function of solution chemistry is crucial towards understanding the electrochemical characteristics of materials in real-world applications. There are several commonly considered factors that affect the stability of a chemical compound, such as metal ion concentration, mixtures of ion concentrations, pH, buffering agents, complexation agents, and temperature. Chemical stability diagrams graphically describe the relative stabilities of chemical compounds, ions, and complexes of a single element as a function of bulk solution chemistry (pH and metal ion concentration) and also describe how solution chemistry changes upon the thermodynamically driven dissolution of a species into solution as the system progresses towards equilibrium" 11.

Stream Analyzer has the capability of building chemical stability diagrams by using the *Chemical Diagram tool* that allows to study the precipitation of species as function of ion concentration and other parameters such as pH. In this section we will explore with different examples how to set up cases to use the Chemical Diagram tool.



¹¹ Santucci, R.J., McMahon, M.E. and Scully, J.R., 2018. Utilization of chemical stability diagrams for improved understanding of electrochemical systems: evolution of solution chemistry towards equilibrium. npj Materials Degradation, 2(1), p.1.

Building a Chemical Diagram

In this section, first we will go through the steps of building a basic chemical diagram, and then we will explore how the concentration of species and other variables such as temperature and pressure have a significant influence on the stability of chemical species.

Example 36: La(C2H₃O₂)₃ – K₃PO₄ Chemical Diagram

For this example, we are going to reproduce the work found in the following paper: Andelman, T., Tan, M. C., & Riman, R. (2010). Thermochemical engineering of hydrothermal crystallization processes. Materials Research Innovations, 14(1), 9-15. https://doi.org/10.1179/143307510X12599329343123.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| La(C2H ₃ O ₂) ₃ – K ₃ PO ₄ Chemical Diagram | | | |
|---|----------------------|-----------------------------------|---------------|
| Calculation Settings | | Stream Composition and Conditions | |
| Stream Name | Chemical Diagrams | Stream Amount | Calculated |
| Name Style | Display Formula | Temperature | 25 °C |
| Unit Set | Metric, Batch, Moles | Pressure | 1 atm |
| Framework | AQ | H2O | 55.5082 moles |
| Type of Calculation | Chemical Diagram | La[C2H3O2]3 ^{Note} | 0.1 moles |
| | | K3PO4 | 0.1 moles |
| | | HNO3 | 0.0 moles |
| | | NaOH | 0.0 moles |

Note: Notice the use of square brackets instead of the traditional parenthesis

Add a new **Stream**

Click on the new Stream and press <F2> to change the name to Chemical Diagrams

Select the AQ thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

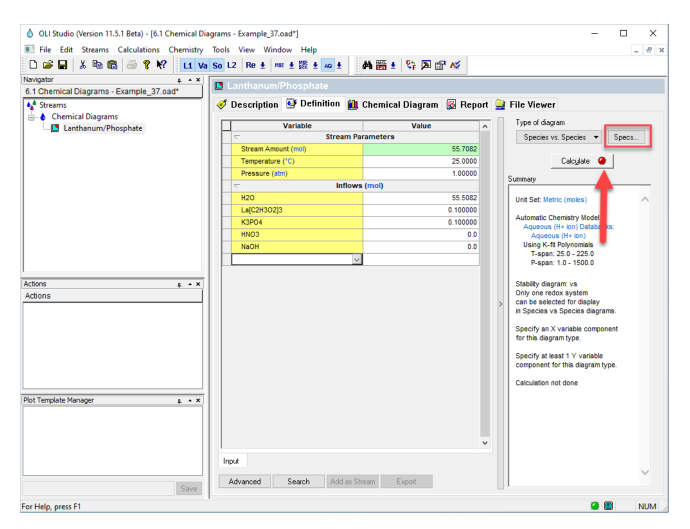
Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the composition of the stream given in the table above. Notice that the concentrations of La(C2H3O2)3 and K3PO4 are held at a 1:1 molal ratio.

Go to the **Add Calculation** button and select **Chemical Diagram** calculation or by selecting the **Chemical Diagram** icon in the Actions Pane

Select Species vs Species as Type of Diagram - Default

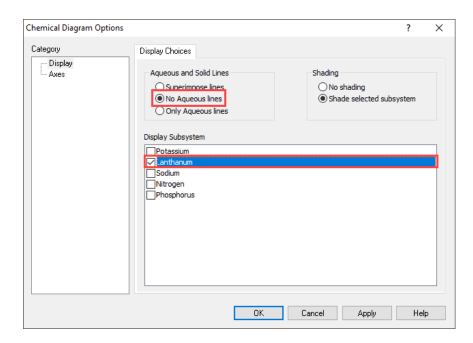
Change the Chemical Diagram name to Lanthanum/Phosphate using the <F2> key



Notice that the calculation is not ready

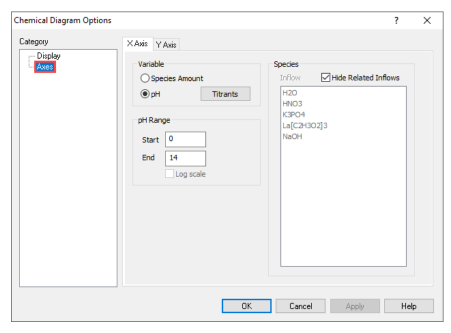
Click on the **Specs** button. This will open the **Chemical Diagram Options** window

In the **Display Choices** tab, select the **No Aqueous lines** button and check the box for **Lanthanum** as the subsystems to display



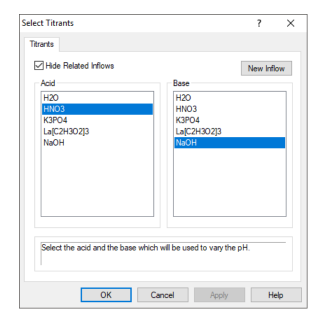
Click on the **Axes** category. The dialog changes to display the **X Axis** tab.

In the **Variable** section select the **pH** option. Leave the default **pH Range**: 0-14. This also enables the **Titrants** button.



Click on the **Titrants** button. Select **HNO3** as the **acid titrant** and **NaOH** as the **base titrant**. Then click **OK**.

Note: These titrants were added in the stream definition, and that's why they appear as possible titrants.

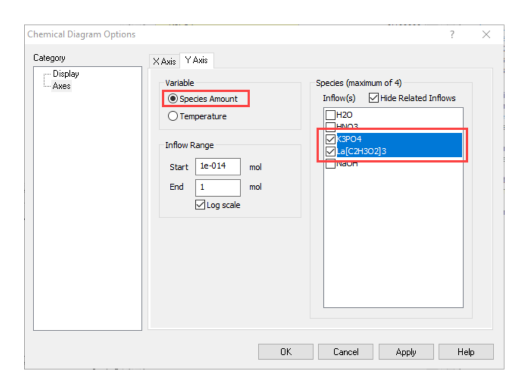


Select the **Y Axis** tab. Select the **Species Amount** button as the variable to change Select La[C2H3O2]3 and K3PO4 species

For the Inflow Range keep the default range: from 1E-14 to 1.0 moles on a log scale.

Note: This means we will have initially 1.0E-14 moles of $La(C_2H_3O_2)_3$ and 1.0E-14 moles of K_3PO_4 increasing equally until we have 1.0 moles of each.

Click **OK**

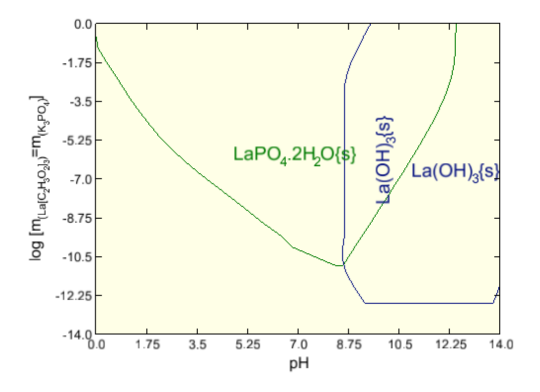


We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file using **(File >Save as...)** or using the **save** icon in the tool bar. You can save it and name it *Chemical Diagrams*.

Analyzing the Results

Click on the **Chemical Diagram** tab (Chemical Diagram). This tab displays the Species vs pH diagram for the Lanthanum species.

The X axis of the diagram is pH, varied by adding HCl or NaOH, and the Y axis variable is the concentrations of $La(C_2H_3O_2)_3$ and K_3PO_4 , held at a 1:1 molar ratio. From this figure, we can see that at room temperature there is a large pH range over which lanthanum phosphate will be the thermodynamically stable product. As pH increases, and more OH ions are present, $La(OH)_3$ begins to form. The pH range in which $La(OH)_3$ forms is the right bounded region in the diagram. The area below the lines is where only aqueous species exist.



Example 37: Adding 50% Lanthanum Acetate

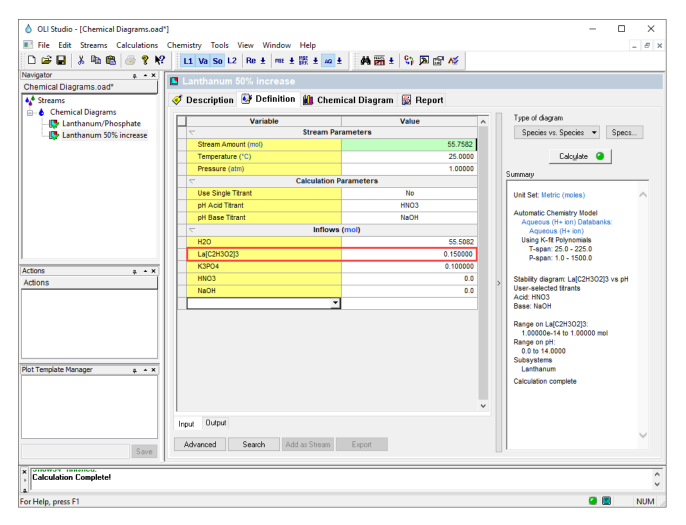
In this second example we will add 50% more to the lanthanum acetate species and see the effect of this increase in concentration on the chemical diagram.

Starting the Simulation

Copy the *Lanthanum/Phosphate* chemical diagram calculated in the Example 36: La(C2H3O2)₃ – K3PO4 Chemical Diagram, and **paste** it under the *Chemical Diagrams* stream.

Change the name to Lanthanum 50% increase using the <F2> key

Change the Lanthanum Acetate amount to 0.15 moles

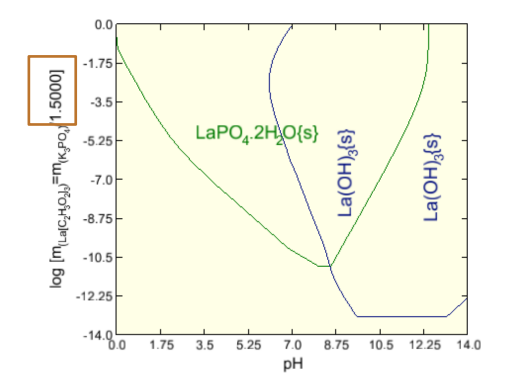


We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file using **(File >Save as...)** or using the **save** icon in the tool bar.

Analyzing the Results

Click on the **Chemical Diagram** tab (**Chemical Diagram**). It is important to note that the Y axis label has changed to reflect the increased amount of the lanthanum acetate.

From this figure, we can see that by increasing amount of Lanthanum acetate, it favors the stability of La(OH)₃ over a wider range of pH.



Example 38: Changing temperature and pressure

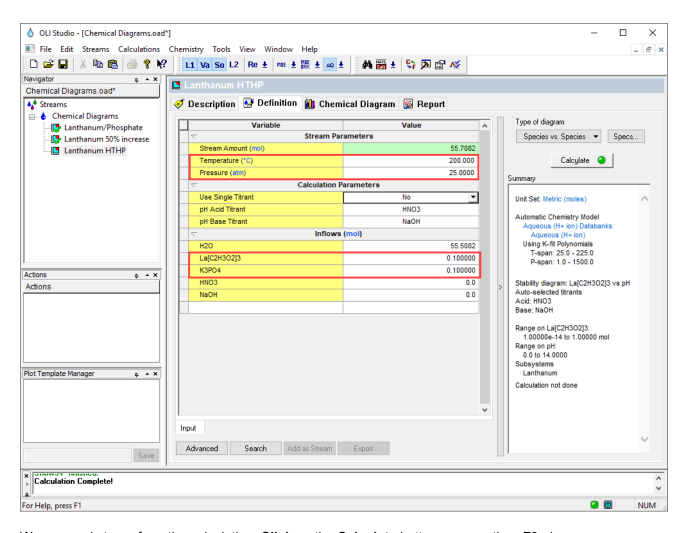
In this third example we will use the original amount of the lanthanum acetate species and see the effect of this increase in temperature and pressure on the chemical diagram.

Starting the Simulation

Copy the *Lanthanum/Phosphate* chemical diagram calculated in the Example 36: La(C2H3O2)₃ – K3PO4 Chemical Diagram, and **paste** it under the *Chemical Diagrams* stream.

Change the name to Lanthanum HTHP using the <F2> key

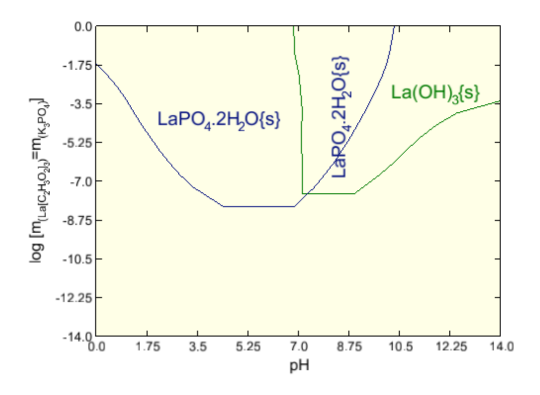
Change the temperature to 200 °C and the pressure to 25 atmospheres



We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file using **(File >Save as...)** or using the **save** icon in the tool bar.

Analyzing the Results

Click on the **Chemical Diagram** tab (Chemical Diagram). This figure shows the stability diagram for $La(C_2H_3O_2)_3$ and K_3PO_4 over a range of pH values at 200 °C and 25 atm. Under these conditions, there is a decrease in the pH range over which $LaPO_4$ is the stable product. This is because as temperature increases, the insoluble rare earth hydroxides $[RE(OH)_3]$ become more stable. The aqueous area of the single-phase increases at elevated temperatures due to the increased solubility of the different species. Again, at high pH values, only $La(OH)_3$ will form.



Chapter III – OLI Studio: Corrosion Analyzer

Corrosion Analyzer is a module within OLI Studio. A separate license enables this module. Corrosion Analyzer is first-principles corrosion prediction tool. As in Stream Analyzer, for all calculations we will create one or more objects, referred to as a **Streams**, which are used to define a particular chemistry, temperature, and pressure.

There are two different types of calculations that can be carried out in Corrosion Analyzer: Corrosion Rates and Stability Diagrams (Potential vs pH and Potential vs Species Diagrams). A brief definition of each type of calculation is given below.



Stability Diagram is used to predict the stability of metals, metal ions, oxides, etc. as a function of T, P, and solution composition. With this tool you can draw conclusions about the ranges of immunity to corrosion, possible passivation, and dissolution of metals in the presence of species that promote or inhibit corrosion.

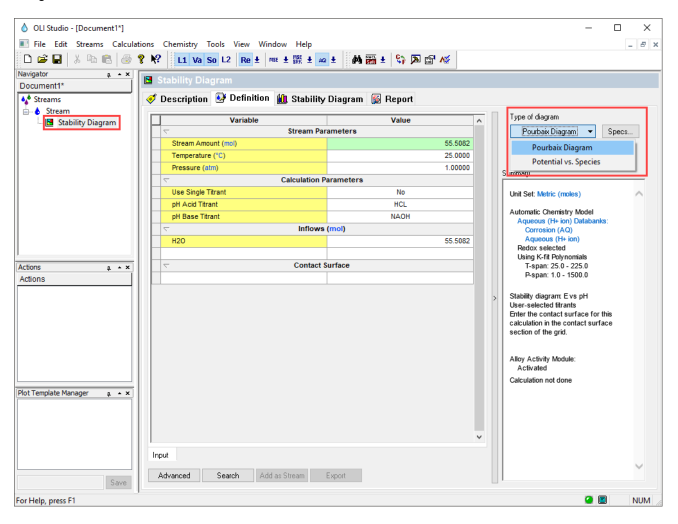


Corrosion Rates is used to predict the general corrosion rate, the propensity for localized corrosion (pitting or crevice corrosion) to occur, polarization curves, and heat treatment effects of metallic materials at one specific equilibrium state. The effects of temperature, pressure, pH, concentration of species, and velocity on corrosion are also included.

In this chapter several examples will be provided to cover these calculation types.

Section 7. Stability Diagrams

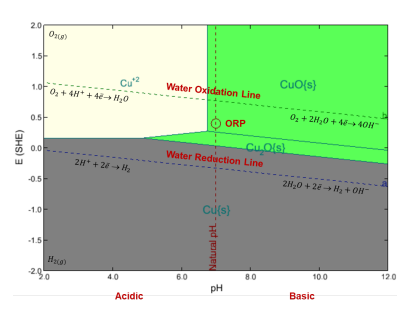
In this section you are going to learn how to set up a Stability Diagram and how to interpret the results when using this tool.



How to Interpret a Stability Diagram

A Stability Diagram, also known as a Pourbaix Diagram or simply a Potential vs pH diagram, maps out the possible stable species in an aqueous environment at different pH and potential combinations. This diagram, however, does not provide information of reaction rates or kinetic effects.

A schematic representation of the Pourbaix diagram for copper in water at 25 °C and 1 atm is shown in the image below.



The gray color represents the immune to corrosion region. It is the stability field of the elemental metal. Cu is the base metal in the plot on the left.

The green color represents the range of possible passivation. It is the stability field of a sparingly soluble compound (usually an oxide,hydroxide, or salt). This compound will form a layer on the surface of the metal, which may protect the metal from corrosion. Having determined that a layer is formed, it is necessary to verify whether it is protective or not because this depends on the crystalline structure of the sparingly soluble compound. In the above plot, the green area contains CuO and Cu₂O as the solid phases produced by oxidation of Cu.

The light-yellow color represents the corrosion region. It is the stability field(s) of dissolved (ionic or neutral) metal species in which neither the metal nor passivating solids are stable. In the plot above, Cu²⁺ is the ionic species that is the most stable.

The diagonal dashed lines, represented by the letters a and b, are the water reduction and water oxidation lines, respectively.

Below the water reduction line (a): Water decomposes to form H_2 gas. Above the water oxidation line (b): Water decomposes to form O_2 gas. The region within these lines (a and b) is called the stability region of water.

The lines that divide different species in the Pourbaix diagram show the equilibrium conditions. These lines represent the equilibrium for chemical and electrochemical reactions.

Finally, as a reference, the values of the Natural pH and the Oxidation Reduction Potential (ORP) are superimposed on the diagram. The natural pH line of the water sample is the water's computed pH before it is adjusted using with acid or base to create the diagram. The ORP is represented by the red circle and indicates the initial electrochemical potential of the water phase. This is the value before the potential is adjusted using a potentiostat, for example.

How to determine if corrosion in the absence of oxygen is possible?

In the absence of oxygen, the most common reduction reaction is the reduction of the proton to elemental hydrogen (as shown by line a in the plot below). Depending on the pH, the reactions that can occur are the following:

Acidic:
$$2H^+ + 2\bar{e} \rightarrow H_2$$

Basic:
$$2H_2O + 2\bar{e} \rightarrow H_2 + OH^-$$

For a corrosion process to proceed, the line **a** must lie above a line that corresponds to an equilibrium between the metal and metal-containing ions.

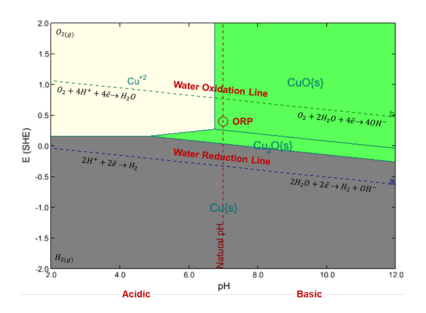
In oxygen-containing solutions, O_2 can be reduced to H_2O (as shown by line \boldsymbol{b}). Depending on the pH, the reactions that can occur are the following:

Acidic:
$$O_2 + 4H^+ + 4\bar{e} \rightarrow H_2O$$

Basic:
$$O_2 + 2H_2O + 4\bar{e} \rightarrow 4OH^-$$

For a corrosion process to occur, the line **b** must lie above a line that corresponds to an equilibrium between the metal and metal-containing ions.

Passivation is likely if **b** lies above a line that corresponds to an equilibrium between the metal and a sparingly soluble compound.



Stability diagram for Copper in Water at 25°C and 1 atm

Building a Pourbaix Diagram

Example 39: Pourbaix Diagram of Fe-H₂O System

After completing this example, you will learn how to set up a Pourbaix Diagram using the **Stability Diagram** tool.

Starting the Simulation



To start the software, double-click the OLI Studio icon on the desktop, which will take you to the OLI Studio interface where you can start creating your calculations.

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Stability Diagram Calculation | | | |
|-------------------------------|----------------------|-----------------------------------|---------------|
| Calculation Settings | | Stream Composition and Conditions | |
| Stream Name | Fe-H2O System | Stream Amount | Calculated |
| Name Style | Display Formula | Temperature | 25 °C |
| Unit Set | Metric, Batch, Moles | Pressure | 1 atm |
| Framework | AQ | pH Range | 2-12 |
| Calculation Type | Stability Diagram | H2O | 55.5082 moles |
| Contact Surface | Fe | | |

Add a new Stream

Click on the new Stream and press <F2> to change the name to Fe-H2O System

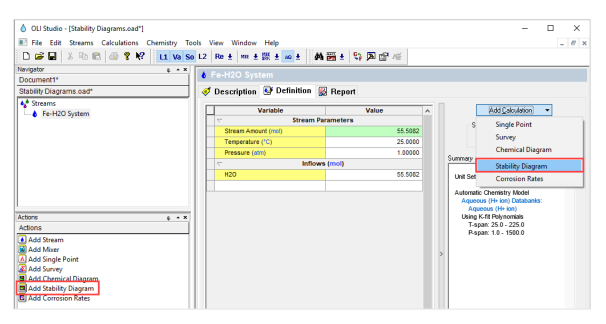
Select the AQ thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the composition, temperature, and pressure of the stream given in the table above

Go to the **Add Calculation** button and select **Stability Diagram** calculation or by selecting the **Chemical Diagram** icon in the Actions Pane.



Select Pourbaix Diagram as Type of Survey - Default

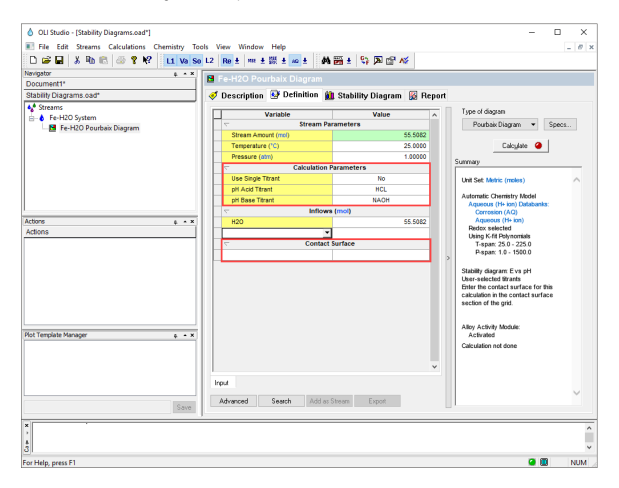
Click on the new Stability Diagram and press <F2> to change the name to Fe-H2O Pourbaix Diagram

Note: Notice that two new sections appear in the **Definition** tab: **Calculation Parameters** and **Contact Surface** grids. Additionally, the Redox button is turned ON.

In the **Calculation Parameters** grid is where the pH titrants need to be defined. These default pH titrants selected by default are HCl and NaOH. You have the option to change the default titrants to your preferred titrants.

In the **Contact Surface** grid is where the metal or alloy of interest needs to be defined. E.g. Fe, Cu, carbon steel, etc.

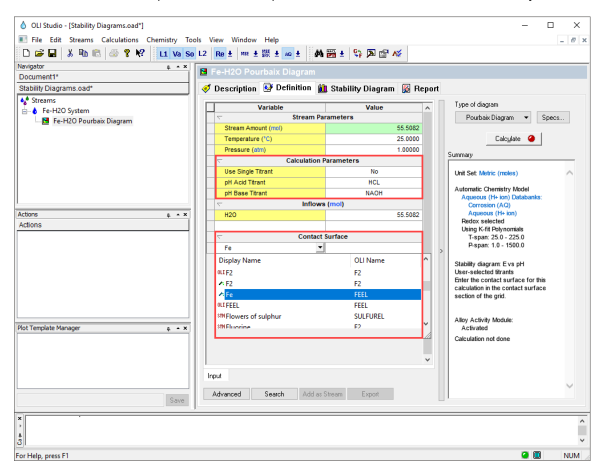
The **Redox** button (Re) is turned ON for this calculation since reduction and oxidation reactions are needed to create the Pourbaix Diagram. This option will enable the different oxidation states of the selected metal.



Leave the default titrants for this calculation: HCl and NaOH

Under the Contact Surface grid type Fe

Note: You also have the option to use the drop-down arrow to search for the metal or alloy of interest.

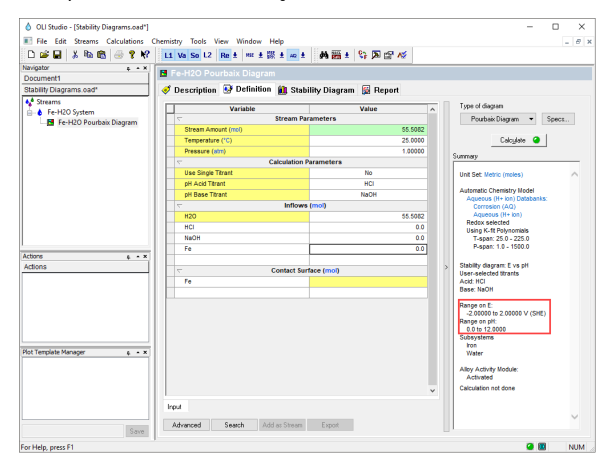


At this point all the basic inputs to create a Pourbaix Diagram have been defined. Notice that in the summary box the following default plotting settings have been defined:

• Potential range: -2 to 2 V (SHE) and

pH range: 0-14.

You can modify these default values before running the calculation.



Click on the Specs button. This will open the Stability Diagram Options Window

Under the **Display** Category you will see the **Display Choices** tab. Under this tab make sure to select the following options:

No Aqueous lines

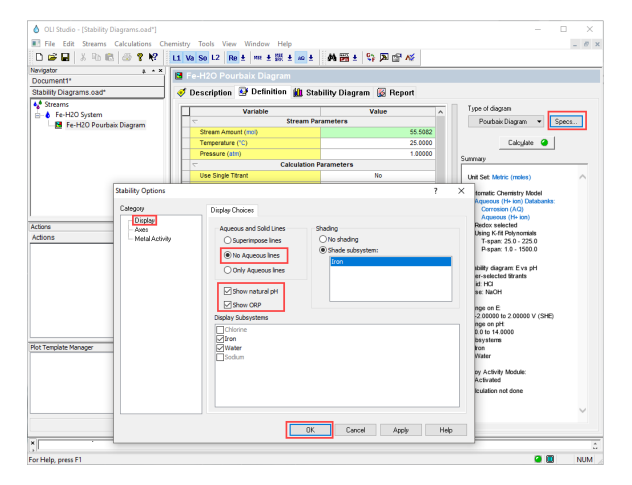
Show natural pH

Show ORP

Under Display Subsystems check the Iron and Water boxes

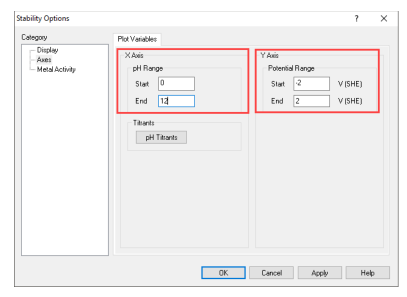
Under Shading select the option Shade Subsystem and select Iron.

Click OK to close the window



Click on the Axes Category. This will show you the Plot Variables tab.

Note: In the **Axes** Category you can modify the axes of the Pourbaix Diagram and change the default titrants.



Change the X Axis or pH range to 2-12

Leave the default values for the Y Axis of Potential Range -2 to 2 V (SHE)

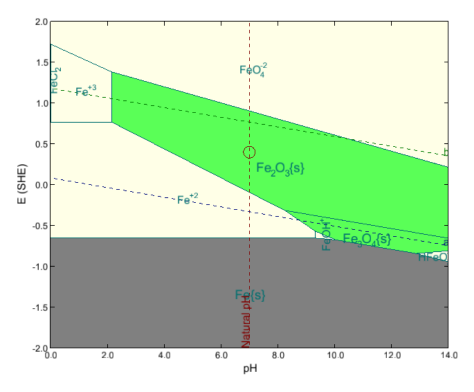
Click **OK** to close the window

We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save it and name it as *Stability Diagrams*.

Analyzing the Results

Click on the **Chemical Diagram** tab (Stability Diagram). This tab displays the Pourbaix Diagram for the Fe-H₂O system at 25 °C and 1 atm.



The obtained diagram is useful for assessing the corrosion behavior of iron in pure water.

Elemental iron (gray section) is stable from -2 to -0.6 V SHE. At a potential greater than -0.6 V SHE, Fe oxidizes to Fe⁺². If the pH is above 10, the boundary is at a lower potential, for example, -0.7 V SHE at 12 pH.

The software computes that Fe $_3$ O₄ (magnetite) is stable between a pH range of ~8.3 and 12. This is also known as the *Passive Film* that protects the iron surface from active corrosion. At higher potentials, the software computes Fe $_2$ O₃ (hematite). Hematite is the dehydrated form of Fe(OH) $_3$ and Fe $_3$ O₄, two forms of rust. These phases provide less corrosion protection to the iron surface than the magnetite. These two phases are represented in the plot by the green color.

The white sections of the plot represent the dominant iron-containing species dissolved in solution. The species with the largest range is Fe⁺². The software computes stability for this species from -0.6 V to 0.77 V SHE, at low pH, and it is no longer stable at pH 10. Active corrosion of the iron surface is predicted if the pH is between 0 and 10. The actual corrosion rate cannot be predicted from this plot, nor can the pH of the water film (diffusion layer) in contact with the iron surface. Therefore, the pH range of 0 to 10 is a guideline and not a direct prediction.

The natural pH and the oxidation reduction potential (ORP) of water, which is represented by the red circle, are showing that when placing Fe in pure water at 25 °C and 1 atm. The point at which the natural pH and the ORP intersect (7 pH, 0.4 V SHE) indicates that Fe₂O₃ (hematite) is the most stable species that forms.

Additionally, notice that the oxidation of iron can be coupled with the reduction of water because water reduction line (line a) lies above the lines that represent the oxidation of iron. Therefore, corrosion of iron can occur with the evolution of hydrogen and formation of soluble iron-containing ions, e.g., Fe²⁺.

Selective Oxidation and Reduction Chemistry

Occasionally, you may want to remove a specific oxidation state for an element when building a Stability Diagram. There are a variety of reasons to perform such an activity, one being that a specific oxidation state may be kinetically unavailable for the reaction. Another case is that perhaps you need to compare and contrast two systems. We will explore how to do this in the following example.

Example 40: Turning off the Fe⁺⁶ oxidation state in the Stability Diagram

In this example you will create a Stability diagram for iron in water at ambient conditions (see previous example for details on how to create this stream), learn how to change the default titrants, and turn off/on different specific oxidation states of the metal of interest.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Stability Diagram Calculation | | | |
|-------------------------------|----------------------|-----------------------------------|----------------|
| Calculation Settings | | Stream Composition and Conditions | |
| Stream Name | Fe-H2O System | Stream Amount | Calculated |
| Name Style | Display Formula | Temperature | 25 ℃ |
| Unit Set | Metric, Batch, Moles | Pressure | 1 atm |
| Framework | AQ | pH Range | 0-14 (Default) |
| Calculation Type | Stability Diagram | H2O | 55.5082 moles |
| Contact Surface | Fe | H2SO4 | 0 moles |
| | | NaOH | 0 moles |

Under the *Fe-H2O System* add a new Stability Diagram calculation. Go to the **Add Calculation** button and select **Stability Diagram** calculation or by selecting the **Add Stability Diagram** icon in the Actions Pane.

Select the AQ thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the composition, temperature and pressure of the stream given in the table above. Add H2SO4 and

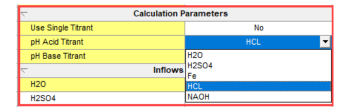
NaOH as inflows. Note: H2SO4 and NaOH will be used as titrants to adjust pH

Select Pourbaix Diagram as Type of Survey - Default

Click on the new *Stability Diagram* and press **<F2>** to change the name to *Fe-H2O Selective Oxidation* Under the **Contact Surface** grid type **Fe**

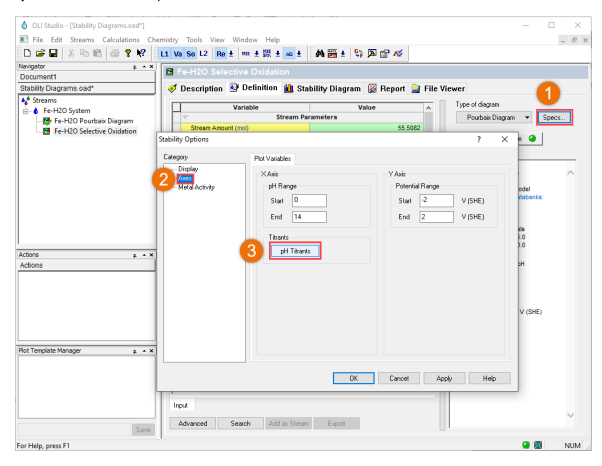
Changing the Default Titrant

In the **Calculation Parameters** grid, **click** on HCl (which is default titrant). A drop-down arrow appears, **click** on it and **select** H2SO4. NaOH has been selected by default.

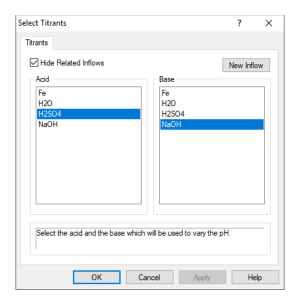


Note: You can also change the default titrant using the Specs button

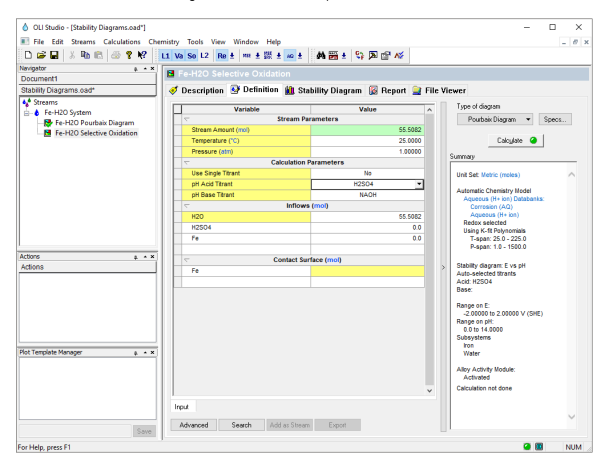
Click on the Specs button Select the Axes option under the Category window Click on pH Titrants. This will open a new window



Select H2SO4 as the Acid titrant, and NaOH as the Base titrant. Then click OK to exit both windows.



The screen should look like the image below after all the inputs and definitions have been entered:

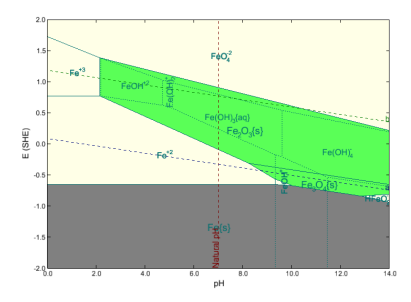


We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Stability Diagrams*.

Analyzing the Results

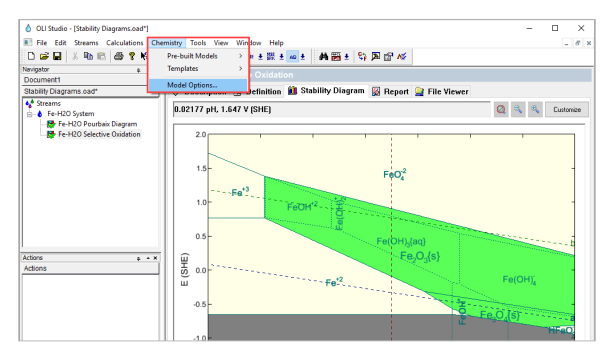
Click on the **Chemical Diagram** tab (Stability Diagram). This tab displays the Pourbaix Diagram for the Fe-H₂O system at 25 °C and 1 atm.



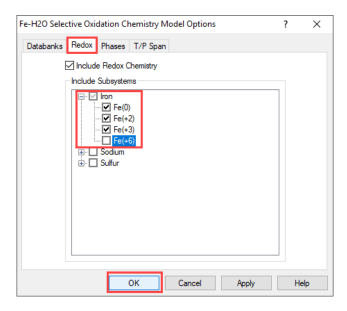
Marcell Pourbaix did not consider the FeO_4^{2-} ion in his work. This is iron in the +6 oxidation state for which there was little thermodynamic data available in the 1960s. To reproduce his work, we need to remove the redox subsystem that pertains to Fe(+6).

Selective Redox, removing an undesired oxidation state

In the Menu bar go to **Chemistry > Model Options**. This will open a new window



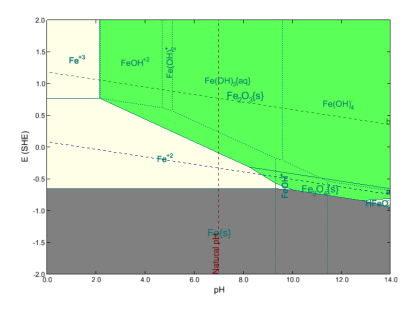
Select the **Redox** tab, expand the Iron Subsystems by clicking on the '+' icon, and **uncheck** the **Fe(+6)** oxidation state. Note that if you have an inflow component with this oxidation state it will remain in the calculation. Click the **OK** button to save your changes and close the window.



We are ready to recalculate to consider the changes. Press the **<F9>** key or go to the **Definition** tab and click on the **Calculate** button.

Analyzing the Results

Click on the **Chemical Diagram** tab (Stability Diagram). In the figure below, you can see that the region dominated by the FeO₄²⁻ ion is not present.



Modeling the Effects of Hydrogen Sulfide on Corrosion

Example 41: The Effect of Hydrogen Sulfide on the Corrosion of Iron

The H₂S (S⁻²) forms a relatively insoluble precipitate with Fe⁺². This precipitate has the potential to cover the metal surface and enhance the pH region where iron is protected from the water. We will model this using the Stability Diagram tool.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Stability Diagram Calculation | | | | |
|-------------------------------|----------------------|-----------------------------------|----------------|--|
| Calculat | tion Settings | Stream Composition and Conditions | | |
| Stream Name | Fe-H2O System | Stream Amount | Calculated | |
| Name Style | Display Formula | Temperature | 25 °C | |
| Unit Set | Metric, Batch, Moles | Pressure | 1 atm | |
| Framework | AQ | pH Range | 0-14 (Default) | |
| Calculation Type | Stability Diagram | H2O | 55.5082 moles | |
| Contact Surface | Fe | H2S | 1e-4 moles | |

Under the *Fe-H2O System* add a new Stability Diagram calculation. Go to the **Add Calculation** button and select **Stability Diagram** calculation or by selecting the **Add Stability Diagram** icon in the Actions Pane.

Select the AQ thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

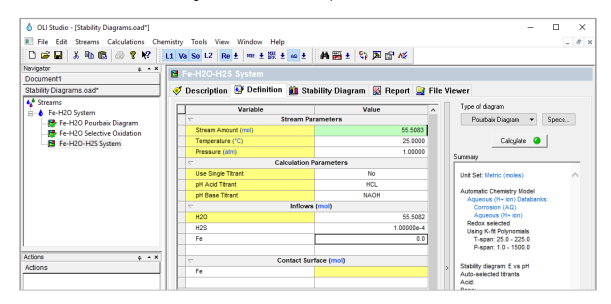
Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the composition, temperature and pressure of the stream given in the table above. **Add 1e-4 moles of H2S** as an inflow.

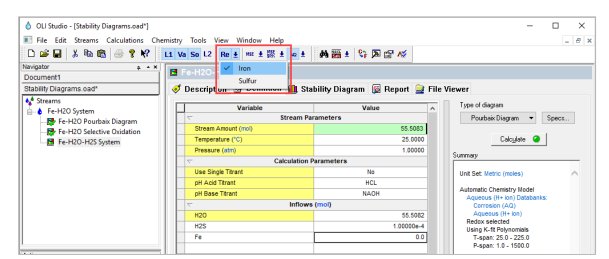
Select **Pourbaix Diagram** as Type of Survey – Default

Click on the new *Stability Diagram* and press **<F2>** to change the name to *Fe-H2O-H2S System* Under the **Contact Surface** grid type **Fe**

The screen should look like the image below after all the inputs and definitions have been entered:



Note: By default, only the transition metals are turned ON. The different oxidation states of elements that are not transition metals are not turned on automatically. You need to turn them ON manually.



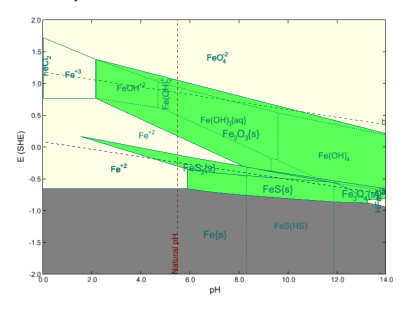
Go to the **Redox** button (**Re**), and **click** on the drop-down arrow **Select Sulfur**. This will turn the oxidation states of sulfur ON

We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Stability Diagrams*.

Analyzing the Results

Click on the **Chemical Diagram** tab (Stability Diagram). This tab displays the Pourbaix Diagram for the Fe-H₂O-H₂S system at 25 °C and 1 atm.



Inspection of the diagram reveals a profound effect of H_2S on the corrosion of iron. New stability fields of FeS and FeS₂ are observed. Elemental iron is found to be in equilibrium with FeS over for pH values ranging from ca. 6.0 to 12.5.

Since the Fe/FeS equilibrium line lies below the H⁺ reduction line (**a**), a process consisting of the reduction of H⁺ to H⁰ and oxidation of Fe to FeS is likely to occur in de-aerated environments. FeS forms a passive film and offers some protection against corrosion.

In fact, the protection due to the formation of FeS is possible over a much wider pH range than that due to the formation of Fe_3O_4 (magnetite) in the absence of H_2S .

This has important implications for corrosion in refinery installations, where H₂S is frequently present.

Modeling Corrosion at High Temperature

Example 42: High Temperature Iron in Water

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Stability Diagram Calculation | | | | |
|-------------------------------|----------------------|-----------------------------------|----------------|--|
| Calcula | tion Settings | Stream Composition and Conditions | | |
| Stream Name | Fe-H2O System | Stream Amount | Calculated | |
| Name Style | Display Formula | Temperature | 300 °C | |
| Unit Set | Metric, Batch, Moles | Pressure | 150 atm | |
| Framework | AQ | pH Range | 0-14 (Default) | |
| Calculation Type | Stability Diagram | H2O | 55.5082 moles | |
| Contact Surface | Fe | | | |

Under the *Fe-H2O System* add a new Stability Diagram calculation. Go to the **Add Calculation** button and select **Stability Diagram** calculation or by selecting the **Add Stability Diagram** icon in the Actions Pane. Select the **AQ** thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

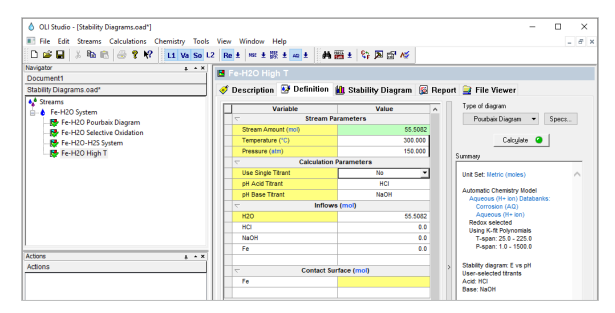
Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the composition, temperature and pressure of the stream given in the table above

Select Pourbaix Diagram as Type of Survey - Default

Click on the new *Stability Diagram* and press **<F2>** to change the name to *Fe-H2O High T* Under the **Contact Surface** grid type **Fe**

The screen should look like the image below after all the inputs and definitions have been entered:

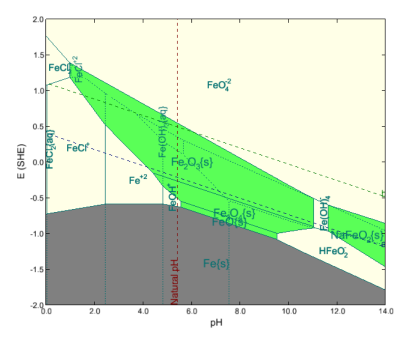


We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Stability Diagrams*.

Analyzing the Results

Click on the **Chemical Diagram** tab ($^{\text{LL}}$ Stability Diagram). This tab displays the Pourbaix Diagram for the Fe-H₂O system at 300 $^{\circ}$ C and 150 atm.



We can see from the diagram above that passivation is only possible at moderate pHs.

Example 43: Neutralization of Refinery Streams with Alkanolamines

In this example you will create a stability diagram for alkaline neutralization in an oil refinery.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Stability Diagram Calculation | | | | |
|-------------------------------|----------------------|-----------------------------------|------------------------|--|
| Calculat | tion Settings | Stream Composition and Conditions | | |
| Stream Name | Fe-H2O System | Stream Amount | Calculated | |
| Name Style | Display Formula | Temperature | 50 °C | |
| Unit Set | Metric, Batch, Moles | Pressure | 1 atm | |
| Framework | AQ | pH Range | 0-14 (Default) | |
| Calculation Type | Stability Diagram | H2O | 55.5082 moles | |
| Contact Surface | Fe | HCI | 0 moles (Acid Titrant) | |
| | | DEA 12 | 0 moles (Base Titrant) | |
| | | C8H18 | 2E-07 moles | |
| | | C7H16 | 8E-07 moles | |
| | | C3H8 | 1.2E-04 moles | |
| | | C4H10 | 2E-05 moles | |
| | | C5H12 | 7E-6 moles | |
| | | C6H14 | 2E-6 moles | |
| | | H2S | 0.01 moles | |

Under the *Fe-H2O System* add a new Stability Diagram calculation. Go to the **Add Calculation** button and select **Stability Diagram** calculation or by selecting the **Add Stability Diagram** icon in the Actions Pane Select the **AQ** thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the composition, temperature and pressure of the stream given in the table above

Select Pourbaix Diagram as Type of Survey - Default

Click on the new Stability Diagram and press <F2> to change the name to Fe-H2O-Alkanoamines

Under the Contact Surface grid type Fe

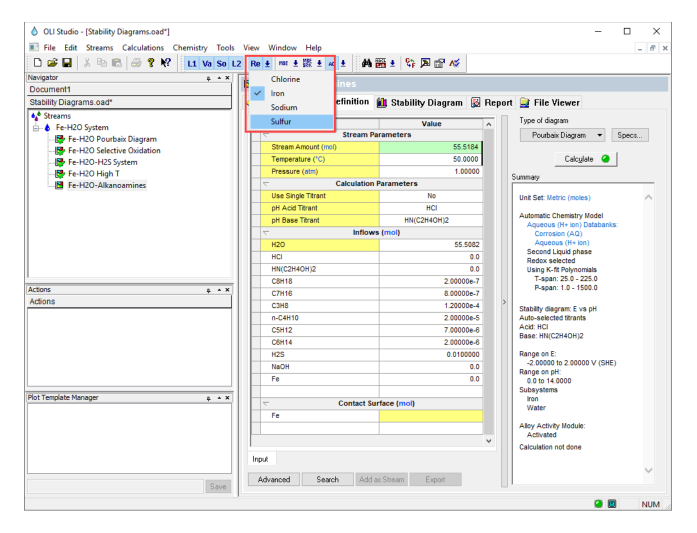
Change the Base Titrant to DEA

Turn Redox ON for Sulfur. Click on the drop-down arrow next to the Re button and select Sulfur

Note: You can also enable Redox reactions via the menu bar: *Chemistry > Model Options > Redox* tab. Make sure that Sulfur is checked.

¹² The OLI Flowsheet: ESP name for this species is DEXH, which can be used as an input to make your life easier. The formula name is: HN(C2H4OH)2

The screen should look like the image below after all the inputs and definitions have been entered:

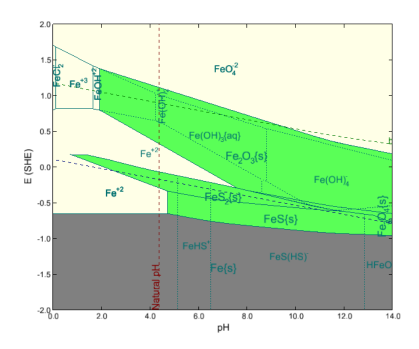


We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Stability Diagrams*.

Analyzing the Results

Click on the **Chemical Diagram** tab ($^{\textcircled{ll}}$ Stability Diagram). This tab displays the Pourbaix Diagram for the Fe-H₂O system at 50 $^{\circ}$ C and 1 atm.



Modeling the Effects of Complexation on Corrosion

In this section we will simulate the reaction of Copper with Ammonia and Gold metal with Cyanide. This section attempts to answer the question of how strong complexing agents affect the passivation of these metals.

Example 44: Copper and Ammonia

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Stability Diagram Calculation | | | | |
|-------------------------------|----------------------|-----------------------------------|----------------|--|
| Calculat | tion Settings | Stream Composition and Conditions | | |
| Stream Name | Cu-H2O System | Stream Amount | Calculated | |
| Name Style | Display Formula | Temperature | 25 °C | |
| Unit Set | Metric, Batch, Moles | Pressure | 1 atm | |
| Framework | AQ | pH Range | 0-14 (Default) | |
| Calculation Type | Stability Diagram | H2O | 55.5082 moles | |
| Contact Surface | Cu | NH3 | 0 moles | |

Add a new Stream

Click on the new Stream and press **<F2>** to change the name to *Cu-H2O System* Select the **AQ** thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

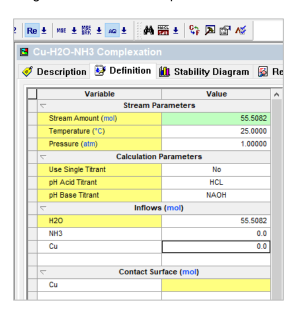
Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the composition, temperature and pressure of the stream given in the table above

Under the *Cu-H2O System* Stream add a new Stability Diagram calculation. Go to the **Add Calculation** button and select **Stability Diagram** calculation or by selecting the **Chemical Diagram** icon in the Actions Pane Select **Pourbaix Diagram** as Type of Survey – Default

Click on the new *Stability Diagram* and press <F2> to change the name to *Cu-H2O-NH3* Under the **Contact Surface** grid type **Cu**

The screen should look like the image below after all the inputs and definitions have been entered:

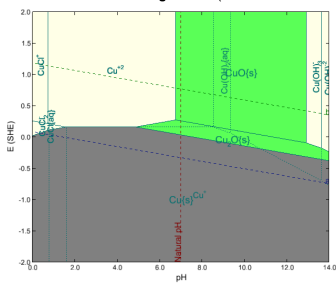


We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Stability Diagrams*.

Analyzing the Results

Click on the Chemical Diagram tab (Stability Diagram). This tab displays the Pourbaix Diagram for the Cu-



 This tab displays the Pourbaix Diagram for the Cu-H₂O system at 25 °C and 1 atm.

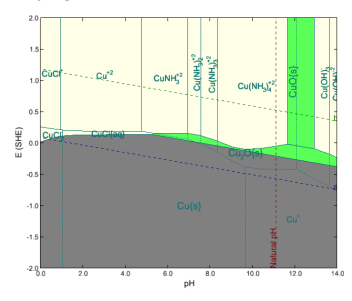
In the absence of oxygen (looking at only the \boldsymbol{a} line), we can see that the copper equilibrium line lies above the hydrogen \boldsymbol{a} line. This means there is insufficient oxidizing power in the water to corrode copper metal in pure water.

What happens if Ammonia (NH_3) is added to the solution? The next step is to understand the effect of NH_3 on the stability of copper.

Effect of NH3 on the Stability of Copper

Go to the **Definition** tab, and change the amount of NH3 from 0 to 0.1 moles **Click** on the **Calculate** button or press the **<F9>** key

Analyzing the Results



Click on the **Chemical Diagram** tab (
Stability Diagram). This tab displays the
Pourbaix Diagram for the Cu-H₂O-NH₃ system at
25 °C and 1 atm in a 0.1 m of NH₃ solution.

Notice that a large area of corrosive liquid has appeared in the stability field for the copper oxides. This means that it is thermodynamically possible for the ammonia to break down the passivation layer of copper oxide in the presence of oxygen. Notice that

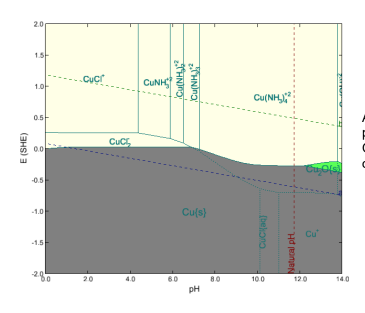
in the absence of oxygen (the \boldsymbol{a} line only), copper is still stable.

Now repeat the exercise with 1.5 moles of NH₃

Go to the **Definition** tab, and change the amount of NH₃ from 0 to 1.5 moles **Click** on the **Calculate** button or press the **<F9>** key

Analyzing the Results

Click on the **Chemical Diagram** tab (Stability Diagram). This tab displays the Pourbaix Diagram for the Cu-H₂O-NH₃ system at 25 °C and 1 atm in a 1.5 m of NH₃ solution.



At this concentration of ammonia, most, if not all the passivating copper oxide, has been reacted away. Only at very high pH values are there any stable oxides.

Example 45: Gold in the presence of Cyanides

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Stability Diagram Calculation | | | | |
|--|----------------------|---------------|----------------|--|
| Calculation Settings Stream Composition and Conditions | | | | |
| Stream Name | Au-H2O System | Stream Amount | Calculated | |
| Name Style | Display Formula | Temperature | 25 °C | |
| Unit Set | Metric, Batch, Moles | Pressure | 1 atm | |
| Framework | AQ | pH Range | 0-14 (Default) | |
| Calculation Type | Stability Diagram | H2O | 55.5082 moles | |
| Contact Surface | Au | NaCN | 0 moles | |

Add a new Stream

Click on the new Stream and press **<F2>** to change the name to *Au-H2O System* Select the **AQ** thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

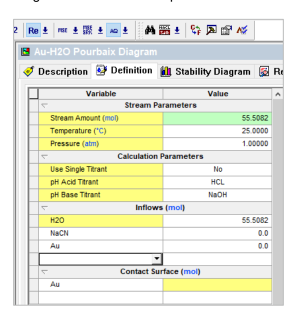
Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the composition, temperature and pressure of the stream given in the table above

Under the *Au-H2O System* Stream add a new Stability Diagram calculation. Go to the **Add Calculation** button and select **Stability Diagram** calculation or by selecting the **Chemical Diagram** icon in the Actions Pane Select **Pourbaix Diagram** as Type of Survey – Default

Click on the new Stability Diagram and press <F2> to change the name to Au-H2O Under the Contact Surface grid type Au

The screen should look like the image below after all the inputs and definitions have been entered:

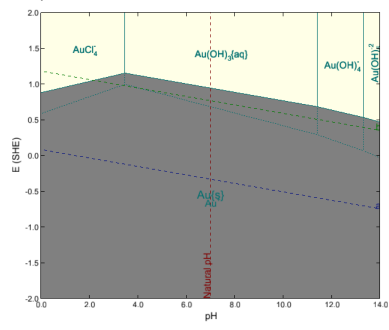


We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file (**File >Save as...**) **or** use the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Stability Diagrams*.

Analyzing the Results

Click on the **Chemical Diagram** tab ($^{\text{LL}}$ Stability Diagram). This tab displays the Pourbaix Diagram for the Au-H₂O system at 25 $^{\circ}$ C and 1 atm.



You can see that without oxygen, gold metal is immune to corrosion. The hydrogen line **a** is below the gold equilibrium line. In the presence of oxygen, gold is still immune to corrosion except at very low pH.

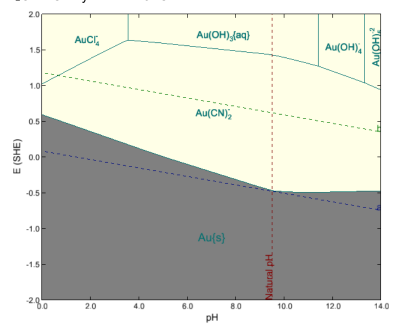
One of the most commonly used leaching processes for **gold extraction** is the **cyanidation** process. This process is a hydrometallurgical technique for extracting gold from low-grade ores by converting the gold to a water-soluble coordination complex. Basically, this approach adds cyanide salts to water. We are going to simulate this using the software.

Effect of Cyanide on the Stability of Gold

Go to the **Definition** tab, and change the amount of NaCN from 0 to 1e-4 moles **Click** on the **Calculate** button or press the **<F9>** key

Analyzing the Results

Click on the **Chemical Diagram** tab (Stability Diagram). This tab displays the Pourbaix Diagram for the Au-H₂O-NaCN system at 25 °C and 1 atm.



In the presence of oxygen, gold completely corrodes with cyanide. This is primarily due to the formation of the gold complex: $Au(CN)_2^-$. This complex is stable over all regions where water is also stable. This means that now gold can be processed in water in a variety of conditions.

Modeling the Effect of Oxidizing Inhibitors on Corrosion

In this chapter we will look at the effect of modeling corrosion inhibitors. We will do this by superimposing two stability diagrams over one another. If one solid field overlaps the corrosion range of the other system, then passivation is likely.

Example 46: Iron in the presence of chromates

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Stability Diagram Calculation | | | | |
|-------------------------------|----------------------|-----------------------------------|----------------|--|
| Calculation Settings | | Stream Composition and Conditions | | |
| Stream Name | Inhibitors | Stream Amount | Calculated | |
| Name Style | Display Formula | Temperature | 25 °C | |
| Unit Set | Metric, Batch, Moles | Pressure | 1 atm | |
| Framework | AQ | pH Range | 0-14 (Default) | |
| Calculation Type | Stability Diagram | H2O | 55.5082 moles | |
| Contact Surface | Fe | | | |
| Contact Surface | Cr | | | |

Add a new Stream

Click on the new Stream and press <F2> to change the name to Inhibitors

Select the AQ thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the composition, temperature and pressure of the stream given in the table above

Under the *Inhibitors* Stream add a new Stability Diagram calculation. Go to the **Add Calculation** button and select **Stability Diagram** calculation or by selecting the **Chemical Diagram** icon in the Actions Pane

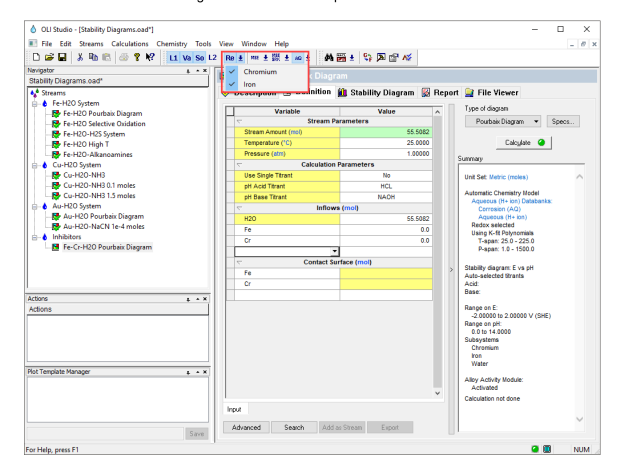
Select Pourbaix Diagram as Type of Survey – Default

Click on the new Stability Diagram and press <F2> to change the name to Fe-Cr-H2O Pourbaix Diagram Under the Contact Surface grid type Fe and then Cr

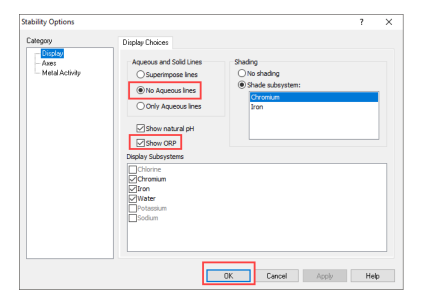
Note: Make sure that both **Fe** and **Cr** are enabled for Redox reactions. Use the arrow next to the **Re** button to expand the list.

You can also enable Redox reactions via the menu bar: **Chemistry** > **Model Options** > **Redox** tab. Make sure that Fe and Cr boxes are checked.

The screen should look like the image below after all the inputs and definitions have been entered:



Click the **Specs** button and under the Display Choices tab check the following options: No Aqueous lines
Show ORP



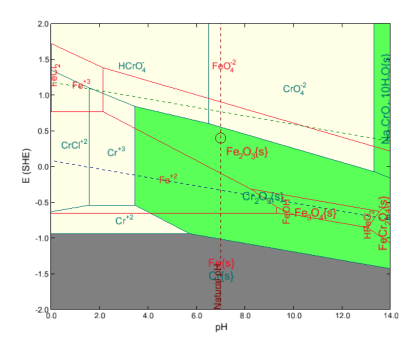
Click **OK** to close the Stability Options window

We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file (**File >Save as...**) or use the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Stability Diagrams*.

Analyzing the Results

Click on the **Chemical Diagram** tab (Stability Diagram). This tab displays the Pourbaix Diagram for the Fe-Cr-H₂O system at 25 °C and 1 atm.



As you can see in this diagram, the shaded chrome passivating solid (Cr(OH)_{3ppt}) overlays the corrosive region of the iron system. This means that there is potential for passivating the metal in that region.

The $Cr(OH)_3$ field overlaps with the corrosion range of Fe in most of the pH range. This causes inhibition because of the coupling of the following: oxidation of Fe to Fe²⁺, reduction of chromates to $Cr(OH)_3$, and the deposition of a protective layer of $Cr(OH)_3$.

Example 47: Iron in the presence of arsenates

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Stability Diagram Calculation | | | | |
|-------------------------------|----------------------|-----------------------------------|----------------|--|
| Calcula | tion Settings | Stream Composition and Conditions | | |
| Stream Name | Inhibitors | Stream Amount | Calculated | |
| Name Style | Display Formula | Temperature | 25 °C | |
| Unit Set | Metric, Batch, Moles | Pressure | 1 atm | |
| Framework | AQ | pH Range | 0-14 (Default) | |
| Calculation Type | Stability Diagram | H2O | 55.5082 moles | |
| Contact Surface | Fe | | | |
| Contact Surface | As | | | |

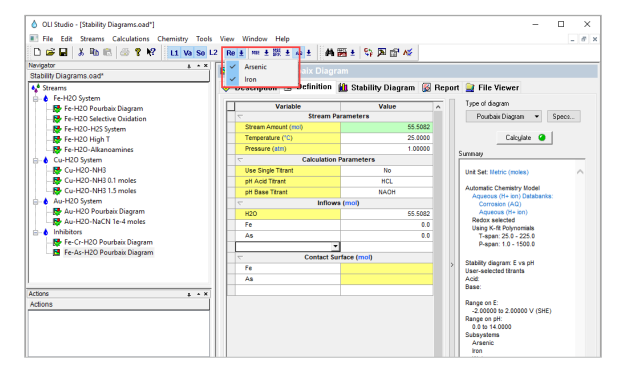
Under the *Inhibitors* Stream add a new Stability Diagram calculation. Go to the **Add Calculation** button and select **Stability Diagram** calculation or by selecting the **Chemical Diagram** icon in the Actions Pane Select **Pourbaix Diagram** as Type of Survey – Default

Click on the new Stability Diagram and press <F2> to change the name to Fe-As-H2O Pourbaix Diagram Under the Contact Surface grid type Fe and then As

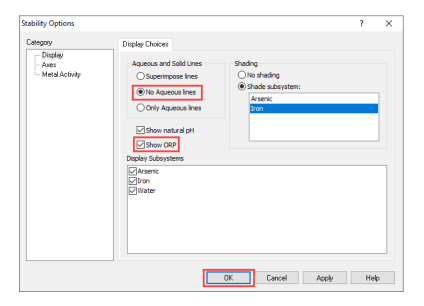
Note: Make sure that both **Fe** and **As** are enabled for Redox reactions. Use the arrow next to the **Re** button to expand the list.

You can also enable Redox reactions via the menu bar: *Chemistry > Model Options > Redox* tab. Make sure that Fe and As boxes are checked.

The screen should look like the image below after all the inputs and definitions have been entered:



Click the **Specs** button and under the Display Choices tab check the following options: No aqueous lines Show ORP

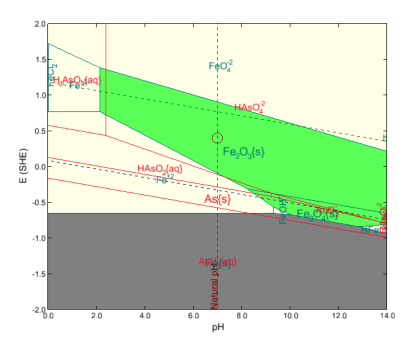


Click **OK** to close the Stability Options window

We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file (**File >Save as...**) **or** use the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Stability Diagrams*.

Analyzing the Results

Click on the **Chemical Diagram** tab (Stability Diagram). This tab displays the Pourbaix Diagram for the Fe-As-H₂O system at 25 °C and 1 atm.



The elemental arsenic field overlaps with the corrosion range of Fe in most of the pH range provided that the conditions are reducing (absence of oxygen). This promotes inhibition because of the coupling of the following: oxidation of Fe to Fe⁺² and the reduction of arsenates to elemental As. This promotes the deposition of a protective layer of As. This can only work in reducing environments; otherwise, the protective layer of As will oxidize and dissolve.

Implications of Stability Diagrams on Cathodic Protection

Cathodic protection works by shifting the potential of the metal into its immunity range. Stability diagrams can help you to answer the following questions:

What is the potential range that ensures that the metal stays in the immunity range?

What is the effect of environmental variables on the immunity domain?

We will explore the Fe-H₂O system at different temperatures and pressures to answer these questions.

Example 48: Effect of Temperature and Pressure on the Pourbaix Diagram of Fe-H2O System

Starting the Simulation

Iron at 30 °C and 1 atm

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Stability Diagram Calculation | | | | |
|-------------------------------|----------------------|-----------------------------------|----------------|--|
| Calculat | tion Settings | Stream Composition and Conditions | | |
| Stream Name | Cathodic Protection | Stream Amount | Calculated | |
| Name Style | Display Formula | Temperature | 30 °C | |
| Unit Set | Metric, Batch, Moles | Pressure | 1 atm | |
| Framework | AQ | pH Range | 0-14 (Default) | |
| Calculation Type | Stability Diagram | H2O | 55.5082 moles | |
| Contact Surface | Fe | | | |

Add a new Stream

Click on the new Stream and press <F2> to change the name to Cathodic Protection

Select the AQ thermodynamic Framework

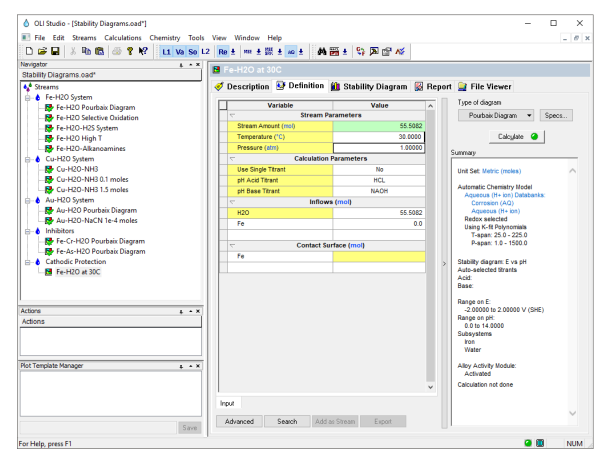
Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the composition, temperature and pressure of the stream given in the table above

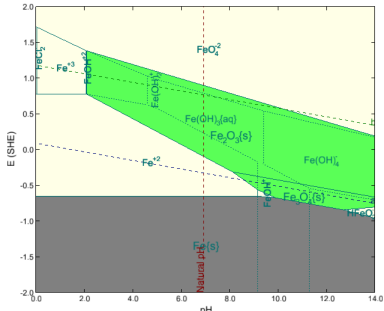
Under the *Cathodic Protection* Stream add a new Stability Diagram calculation. Go to the **Add Calculation** button and select **Stability Diagram** calculation or by selecting the **Chemical Diagram** icon in the Actions Pane Select **Pourbaix Diagram** as Type of Survey – Default

Click on the new Stability Diagram and press <F2> to change the name to Fe-H2O at 30C and 1 atm Under the Contact Surface grid type Fe The screen should look like the image below after all the inputs and definitions have been entered:



Analyzing the Results

Click on the **Chemical Diagram** tab (Stability Diagram). This tab displays the Pourbaix Diagram for the Fe-H₂O system at 30 °C and 1 atm.



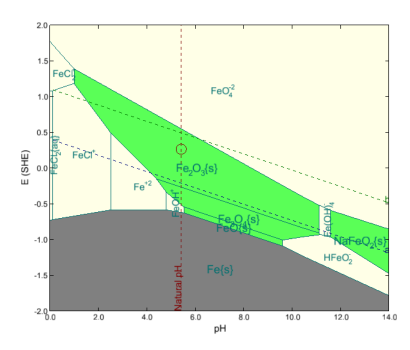
In the whole pH range, cathodic protection will require shifting the potential to moderately negative values.

Iron at 300 °C and 100 atm

Go to the **Definition** tab, and change the Temperature and Pressure to 300°C and 100 atm **Click** on the **Calculate** button or press the **<F9>** key

Analyzing the Results

Click on the **Chemical Diagram** tab (Stability Diagram). This tab displays the Pourbaix Diagram for the Fe-H₂O system at 300 °C and 100 atm.

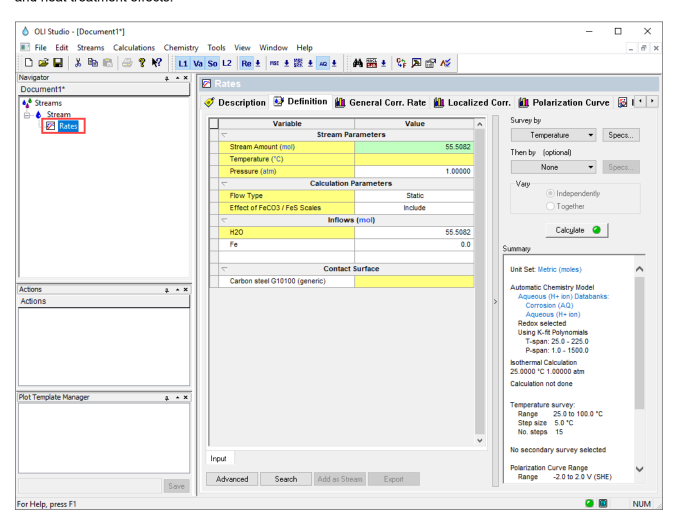


The immunity range in acidic and neutral solutions is weakly affected by temperature. However, the immunity range in alkaline solutions is shifted to much lower potentials which makes cathodic protection much more difficult.

Section 8. Introduction to Rates of Corrosion

In this section you are going to learn how to set up a Corrosion Rate calculation, how to study the effect of different variables on the corrosion rates of different alloys, such as temperature, pH, flow velocity, etc., and how to interpret the results when using this tool.

We will also explore how to interpret the results for the propensity to localized corrosion, polarization curves, and heat treatment effects.



A Basic General Corrosion Rate Calculation

A brief introduction to the corrosion rate tool will be shown in the example below. As we go through the example, the basic definitions, functionalities, and reporting for the *Corrosion rate* tool will be introduced. Note: As of version 11.5 and earlier, only the AQ thermodynamic framework supports the corrosion rate calculation.

Example 49: Corrosion rate of an oxygenated 0.1 m NaCl solution

Starting the Simulation

To start the software, double-click the OLI Studio icon on the desktop, which will take you to the OLI Studio interface where you can start creating your calculations.

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Corrosion Rate Calculation | | | | |
|----------------------------------|-------------------------------|-----------------------------------|---------------|--|
| Calcu | lation Settings | Stream Composition and Conditions | | |
| Stream Name | CR of carbon steel | Stream Amount | Calculated | |
| Name Style | Display Formula | Temperature | 25 °C | |
| Unit Set | Unit Set Metric, Batch, Moles | | 1 atm | |
| Framework | AQ | H2O | 55.5082 moles | |
| Calculation Type Corrosion Rates | | 02 | 2.5e-4 moles | |
| Contact Surface | Carbon Steel G10100 (generic) | NaCl | 0.1 moles | |

Add a new Stream

Click on the new Stream and press <F2> to change the name to CR of carbon steel

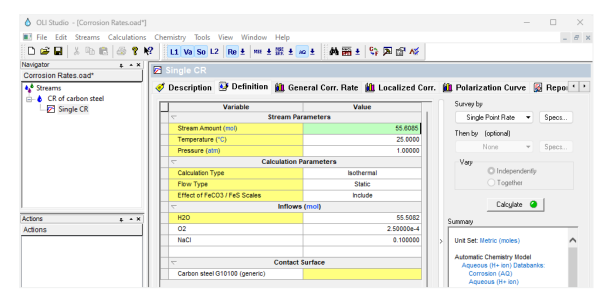
Select the AQ thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

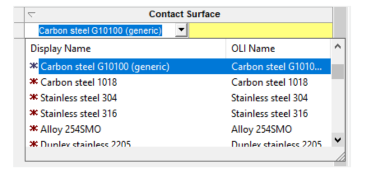
Click on the Units Manager Icon, and select Metric, Batch, Moles

Enter the composition, temperature and pressure of the stream given in the table above

Go to the **Add Calculation** button and select **Corrosion Rates** calculation or by selecting the **Add Corrosion Rates** icon in the Actions Pane. Your screen should look like the image below.

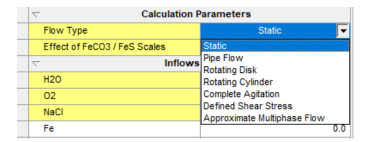


Note: By default, the software has selected **Carbon steel G10100 (generic)** as the default contact surface to perform the corrosion rates on. You can click on the white space of the **Carbon steel G10100 (generic)** name, and it will show a drop-down arrow. You can then click on the dropdown arrow, and you will be able to see and select any alloy present in the **Alloy** database.



Additionally, the **Calculation Parameters** grid shows two options: (1) Flow Type and (2) Effect of FeCO3 /FeS scales on corrosion rates.

By default, the software selects **Static** as **Flow Type**. There are 7 options for Flow Type, and a brief description of the first five options will be given below.



Static

The solution is not flowing in this calculation.

Pipe Flow

The fluid is flowing through a pipe. The pipe diameter and flow velocity must be defined. The default pipe diameter is 0.1 meters and the default flow velocity is 2 m/s.

Rotating disk

This reproduces a type of experiment that is used quite frequently in the laboratory. A disk is rotated to bring fluid to the surface of the electrode in a predictable manner. The diameter of the disk is specified as well as the revolutions per minute (RPM). The default diameter is 0.01 meters, and the default RPM is 5000 RPM.

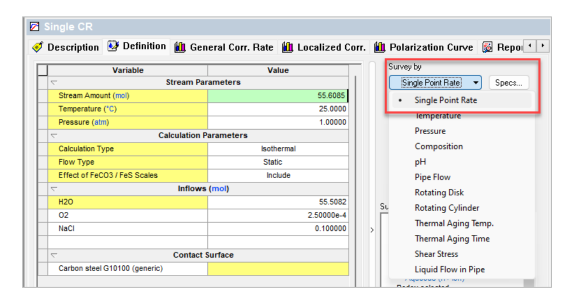
Rotating Cylinder

This reproduces a type of experiment that is used quite frequently in the laboratory. A cylindrical rotor is rotated to bring fluid to the surface of the electrode in a predictable manner. The diameter of the rotor is specified as well as the revolutions per minute (RPM). The default diameter is 0.01 meters, and the default RPM is 5000 RPM.

Complete Agitation

In this calculation, the liquid phase is completely agitated, and no mass transfer limitations apply.

Leave the default alloy **Carbon steel G10100 (generic)** as the contact surface **Go** to the **Survey by** option and select **Single Point Rate**. This option will allow you to calculate the corrosion rate at the temperature and pressure specified.



Note: You can also study the effect of other variables such as temperature, pressure, (chemical species) composition, pH, pipe flow, etc. on the corrosion rate. The effect of these variables can be set up as a survey calculation. In the upcoming corrosion rate examples, survey calculations to study corrosion rates will be shown in more detail.

A brief description of the most common survey types for studying corrosion rates is given below.

pH Survey – This calculation is like the pH survey available in OLI Studio: Stream Analyzer. The specification requires a titrant acid and base to change the pH.

Temperature Survey – The default range is from 25-100 °C. Any range may be used by changing the Range option. The user should consider that some points in the survey may not converge due to phase changes (e.g., boiling off of aqueous liquids).

Composition Survey – The composition of a chemical compound, for example NaCl, can be varied to study its effects on corrosion rates. The range of the chemical compound defaults from 0 to 1 mole with an increment of 0.1 moles. This range can be changed via the Range option.

Care should be taken when adding salts that can form hydrates (e.g., CaCl₂.6H₂O). When these hydrated salts begin to precipitate from solution, large amounts of water may be complexed with the crystal. The solution may dehydrate, and non-convergence may be the result.

Pressure Survey – The pressure of the system can also be varied. The default range can be changed via the Range option. Care should be taken when working at very low pressures since the solution may inadvertently boil off the liquid and non-convergence may result.

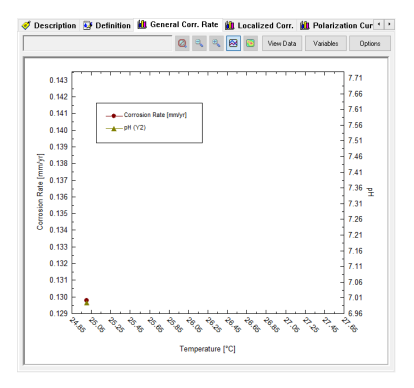
Flow Velocity Survey - In systems that are flowing, the flowrate of the stream can be varied.

Now, we are ready to perform the calculation. Click on the Calculate button or press the <F9> key

It is time to save your file (File >Save as...) or use the save icon in the tool bar. Create a new file and name it: Corrosion Rates.

Analyzing the Results

Click on the **General Corr. Rate** tab (General Corr. Rate). This tab displays a plot showing the results of corrosion rate and pH.



Click on the View Data button to see the results in tabulated form.

| | | Temperature | Corrosion Rate | pН |
|---|---|-------------|----------------|---------|
| I | | °C | mm/yr | |
| ľ | 1 | 25.0000 | 0.129801 | 6.99516 |

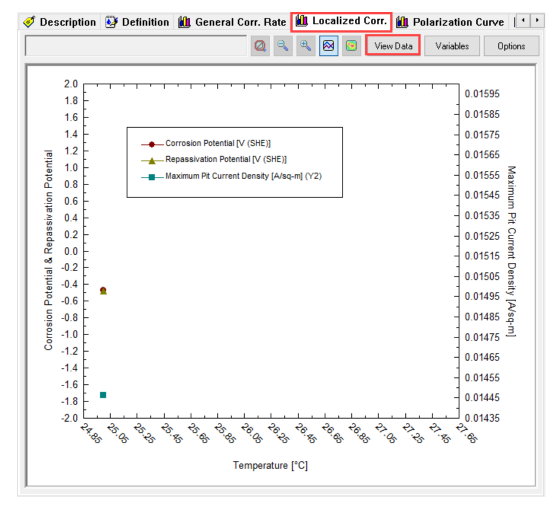
The predicted corrosion rate of carbon steel at 25 °C and 1 atm is ~0.13 mm/year when exposed to an oxygenated 0.1 m NaCl solution. It is important to highlight that this predicted corrosion rate is the corrosion rate once the system has reached a steady state.

Click on the **Localized Corr.** Rate tab (Localized Corr.). This tab displays a plot showing the results of three different calculated data:

Corrosion Potential (red dot)

Repassivation Potential (yellow triangle)

Maximum Pit Current Density (blue square)

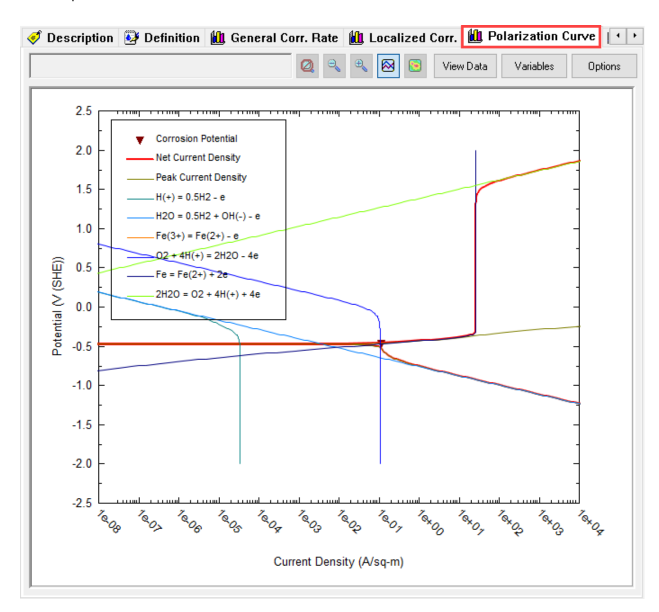


Click on the View Data button to see the results in tabulated form.

| | | Temperature | Corrosion Potential | Repassivation Potential | Maximum Pit Current Density |
|---|---|-------------|---------------------|-------------------------|-----------------------------|
| | | °C | V (SHE) | V (SHE) | A/sq-m |
| • | 1 | 25.0000 | -0.472418 | -0.481902 | 0.0144636 |

Notice that in this case, the corrosion potential has a higher value than the repassivation potential. This indicates that carbon steel at 25 °C and 1 atm when exposed to an oxygenated 0.1 m NaCl solution will likely suffer localized corrosion (pitting or crevice corrosion). The maximum pit current density gives the worst-case pitting rate for these conditions.

Now, click on the **Polarization Curve** tab (Polarization Curve). This tab displays a plot showing the calculated polarization curve of carbon steel.



The net polarization curve is given by the red curve. This red curve is obtained by adding up all the currents of the half reactions at a given potential. The corrosion potential is calculated by applying the mixed potential theory, and it is represented by this red triangle. Once the corrosion potential is obtained, the corrosion current density is also computed and, subsequently, so is the corrosion rate.

Exploring corrosion rate calculation options

Example 50: Corrosion in a Water-Filled Carbon Steel Tank

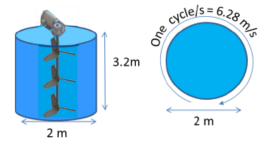
In the following example, you are going to study the corrosion rate of a carbon steel tank with a volume of 10 m³ (3.2 cm high and 200 cm in diameter) that is filled to the top with water. The tank's wall thickness is 1.27 cm.

These are the operation conditions:

Sometimes the water in the tank remains static (mixer off), closed to the atmosphere Some other times the tank is fully mixed (complete agitation)

The tank when operating has a variable speed mixer with a rotation speed between 0 and 12,000 rpm At other times, the tank is open to the atmosphere. The effects of O₂, CO₂, and flow will be studied.

We will evaluate each instance to compute the corrosion risk.



Starting the Simulation

First operation condition: Static flow, tank closed to the atmosphere at ambient conditions

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc., will require further adjustments, and will be described as necessary.

| Corrosion rate calculation of tank as a function of flow | | | | |
|--|-------------------------------|---------------|------------|--|
| Calculation Settings Stream Composition and Conditions | | | | |
| Stream Name | CR vs flow | Stream Amount | 10000 L | |
| Calculation Type | Corrosion Rates | Temperature | 25 ℃ | |
| Name Style | Display Name | Pressure | 1 atm | |
| Unit Set | Metric, Batch, Concentration | Water | Calculated | |
| Framework | AQ | | | |
| Contact surface | Carbon Steel G10100 (generic) | | | |

Add a new Stream

Click on the new Stream and press **<F2>** to change the name to *CR vs flow* Select the **AQ** thermodynamic Framework

Click on the Names Manager Icon, and select the *Display Name* option, and click **OK**Click on the Units Manager Icon, and select the *Metric, Batch, Conc.* option, and click **OK**



In the Definition tab, notice that the units of the Stream Amount changed from moles (mol) to liters (L).

Enter the composition, temperature, and pressure of the stream given in the table above Go to the **Add Calculation** button and select **Corrosion Rates** calculation or select the **Add Corrosion Rates** icon in the Actions Pane.

Change the *Rates* name to *Static Flow, 25C* using the <F2> key.

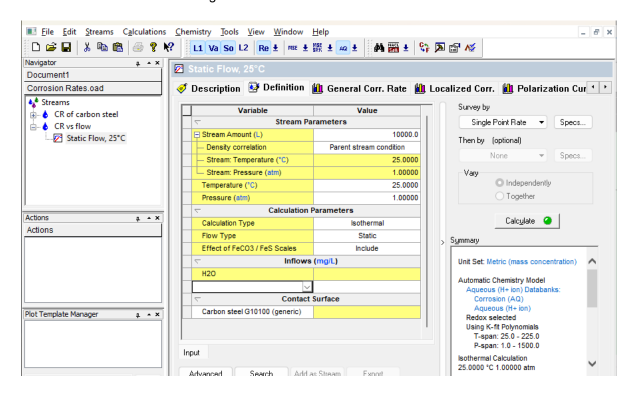
Leave the default Calculation Type as Isothermal

Leave the default Flow Type as Static

Leave the default alloy Carbon steel G10100 (generic) as the contact surface

Go to the **Survey by** option and select **Single Point Rate**. This option will allow you to calculate the corrosion rate at the temperature and pressure specified.

Your screen should look like the image below.

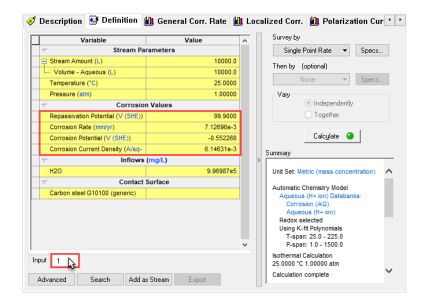


We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key.

It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Corrosion Rates*.

Analyzing the Results

Click on the output tab labeled **1** (next to the input tab), and check the results located in the **Corrosion Values** tab.



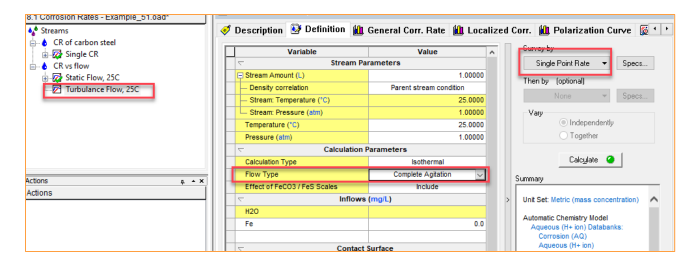
The corrosion rate is 7.13e-3 mm/yr. This is a negligible rate, since the thickness of the tank is \sim 12.7 mm, then corroding half the wall thickness would take about 900 years.

Second operation condition: Turbulent flow, tank closed to the atmosphere at ambient conditions

Under the stream 'CR vs Flow' add new corrosion rate calculation. Go to the Add Calculation button and select Corrosion Rates calculation or select the Add Corrosion Rates icon in the Actions Pane Change the Rates name to Turbulent Flow, 25C using the <F2> key. Leave the default Calculation Type as Isothermal Change the default Flow Type to Complete Agitation

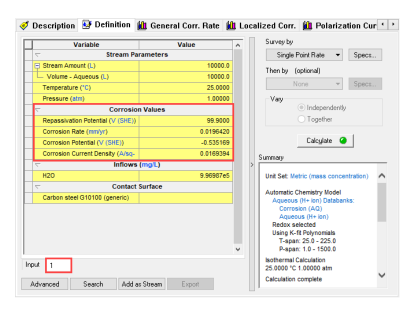
Leave the default alloy Carbon steel G10100 (generic) as the contact surface Go to the Survey by option and select Single Point Rate.

Your screen should look like the image below.



Analyzing the Results

Click on the output tab labeled 1 (next to the input tab), and check the results located in the **Corrosion Values** tab.



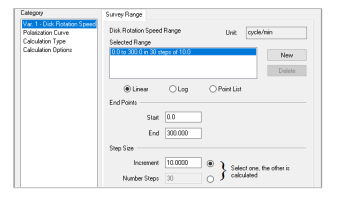
The corrosion rate increased 0.0196 mm/year, still a relatively low value.

Third operation condition: Varying flow, tank closed to the atmosphere at ambient conditions

Under the stream 'CR vs Flow' add new corrosion rate calculation. Go to the Add Calculation button and select Corrosion Rates calculation or by selecting the Add Corrosion Rates icon in the Actions Pane Change the Rates name to Varying Flow, 25C using the <F2> key Go to the Survey by button and select Rotating Disk Under the Calculation Parameters grid. change the Disk Diameter to 200 cm

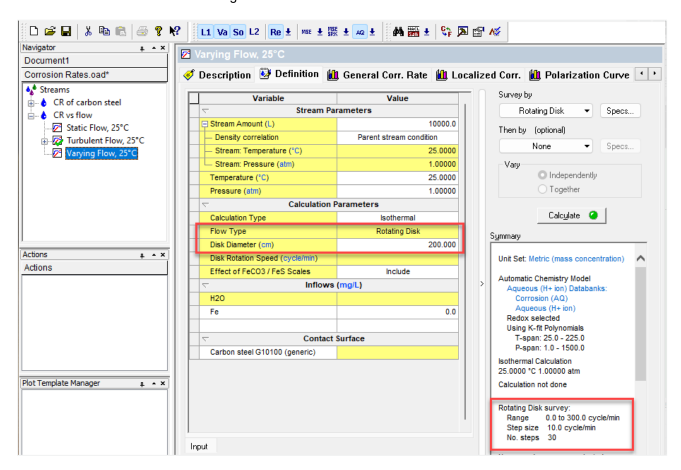
Note: At this point, the vertical dimensions of the tank are unimportant. We will assume that the tank can be modeled like a rotating cylinder. The propeller rotates at 1200 rpm, although it is not expected that the wall velocity will approach this value, and so a lower value will be used (we still want it to be high enough to see the effects of shear).

The next step is to set the speed of the mixer. Go to the **Specs** button next to the **Rotating Disk** option Change the Disk Rotation Speed Range from 0 to 300 cycles/min with an increment of 10 cycles/min. Then click **OK**.



Leave the default alloy Carbon steel G10100 (generic) as the contact surface

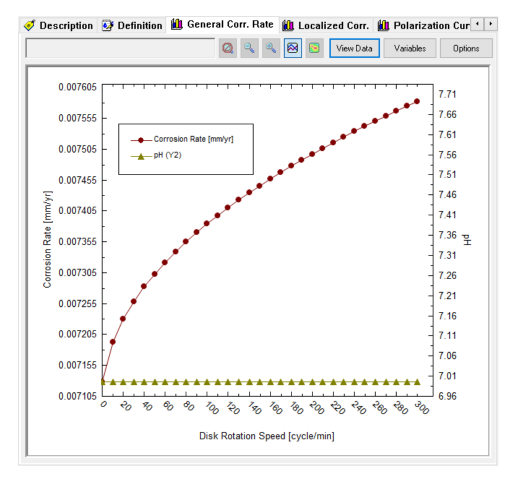
Your screen should look like the image below.



We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file **(File >Save as...) or** use the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Corrosion Rates*.

Click on the **General Corr. Rate** tab (General Corr. Rate). This tab displays a plot showing the results of corrosion rate and pH.



The corrosion rate is computed to increase as the bulk liquid velocity increases from 0 to 300 rpm near the wall surface. The reason is straightforward; the higher velocity reduces the static water film thickness on the metal surface. This *diffusion layer* film limits the mass transfer of corrosion products and bulk chemicals to and from the surface. As the liquid velocity (and therefore shear force) increases, the film thickness decreases, and the concentration gradient increases. This increases the flux of materials to and from the surface, which increases corrosion rates.

Fourth operation condition: Static flow, tank open to the atmosphere (aerated with O₂ only) at ambient conditions

Open steel tanks are in contact with oxygen in the air. To simulate this, follow the steps below.

Under the stream 'CR vs Flow' add new corrosion rate calculation. Go to the **Add Calculation** button and select **Corrosion Rates** calculation or press the **Add Corrosion Rates** icon in the Actions Pane Change the **Rates** name to **Static Flow**, **O2** using the **<F2>** key

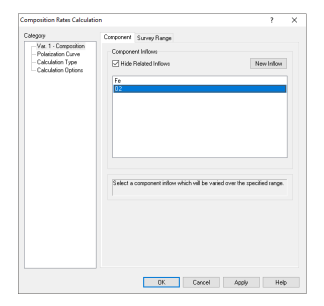
Add Oxygen (O2) to the inflow grid

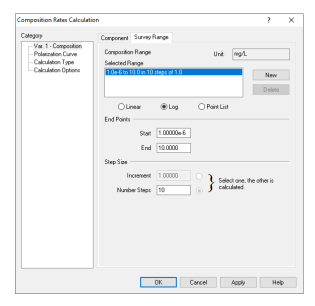
Leave the default Flow Type as **Static**

Go to the Survey by button and select Composition

The next step is specifying the concentration range of oxygen. Go to the **Specs** button next to the **Composition** option. This will open a new window.

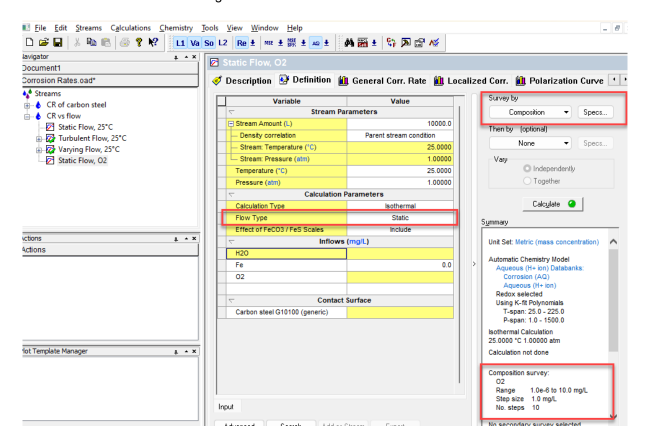
Under **component** select (click on) O2. Then select the **Survey Range** tab and enter the Survey Range as follows: Log Scale, Start=1e-6, End=10, Steps=10. Then click **OK**.





Leave the default alloy Carbon steel G10100 (generic) as the contact surface

Your screen should look like the image below.



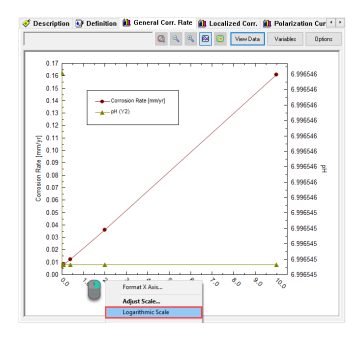
We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file **(File >Save as...) or** use the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Corrosion Rates*.

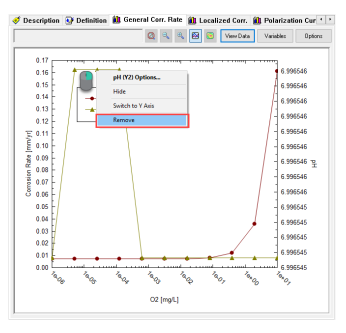
Analyzing the Results

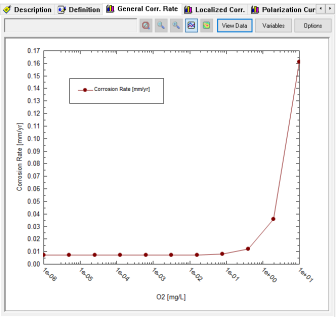
Click on the **General Corr. Rate** tab (General Corr. Rate). This tab displays a plot showing the results of corrosion rate and pH.

Right-mouse click on the X-axis and change it to Logarithmic Scale



Right-mouse click on the pH curve and select Remove





The resulting plot shows the impact of oxygen on the corrosion rate. Corrosion increases by a factor of 20 once the concentration increases beyond 100 μ g/l. Aerated water contains approximately 8 mg/l O₂. Corrosion on the tank wall is 0.16 mm/yr. At this rate, 50% of the 12.3 mm wall will be dissolved in 40 years.

Fifth operation condition: Static flow, tank open to the atmosphere (aerated with O₂ and CO₂) at ambient conditions

The atmosphere contains \sim 400 ppmV CO₂. At this concentration, 0.6 mg/l CO₂ is dissolved in water as molecular CO₂; this CO₂ hydrolyzes water to form the following reactants:

$$CO_2 + H_2O = H^+ + HCO_3^-$$

The resulting pH is about 5.6 at ambient conditions.

The impact of CO₂ on corrosion is two-fold, as two separate reactions occur at the metal surface:

$$H^+ + e^- \leftrightarrow \frac{1}{2}H_2$$

$$HCO_3^- + e^- = \frac{1}{2}H_2 + CO_3^{-2}$$

To test the CO₂ impact, you will recalculate the corrosion rate using two CO₂ concentrations: 0 and 0.6 ppm.

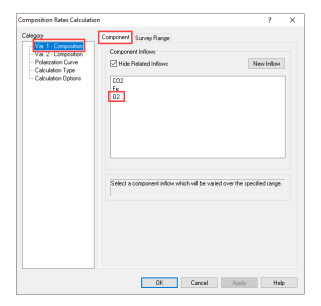
Under the stream 'CR vs Flow' add new corrosion rate calculation. Go to the Add Calculation button and select Corrosion Rates calculation or select the Add Corrosion Rates icon in the Actions Pane Change the Rates name to Static Flow, O2 and CO2 using the <F2> key Add Oxygen (O2) and carbon dioxide (CO2) to the inflow grid Leave the default Flow Type as Static

Go to the Survey by button and select Survey by Composition and then by Composition (as shown below)

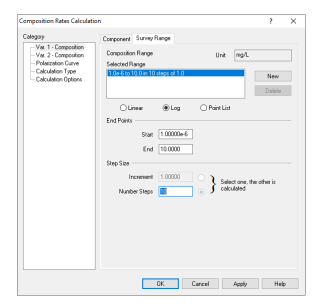


The next step is the concentration range of oxygen and carbon dioxide. Go to the **Specs** button next to the **Composition** option. This will open a new window.

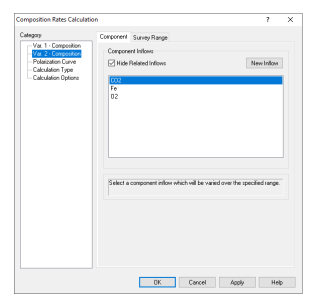
Under the **Category** section, make sure to select **Var. 1 -Composition**. Go to the **Component** tab, and select O2



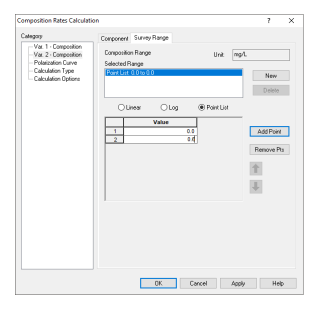
Then select the **Survey Range** tab and enter the Survey Range as follows: Log Scale, Start=1e-6, End=10, Steps=10. Then click **OK**.





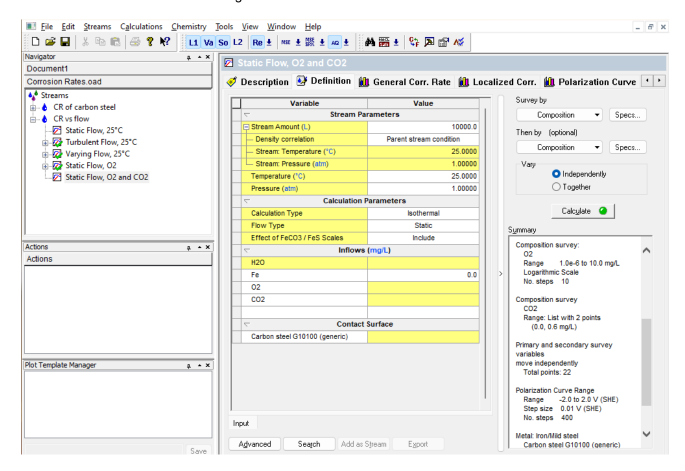


Then select the **Survey Range** tab. Select the Point List option. Create two points with values of 0 and 0.6 mg/L. Then click **OK**.



Leave the default alloy Carbon steel G10100 (generic) as the contact surface

Your screen should look like the image below.



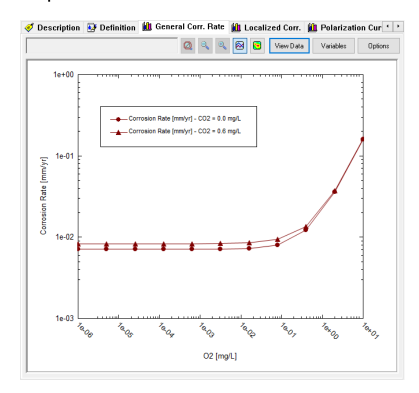
We are ready to perform the calculation. Click on the Calculate button or press the <F9> key

It is time to **save** your file **(File >Save as...) or** use the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Corrosion Rates*.

Analyzing the Results

Click on the **General Corr. Rate** tab (General Corr. Rate). This tab displays a plot showing the results of corrosion rate and pH.

Right-mouse click on the X-axis and change it to Logarithmic Scale Right-mouse click on the Y-axis and change it to Logarithmic Scale Right-mouse click on the pH curve and select Remove Series

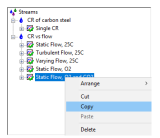


The effects of CO_2 are small, particularly in relation to the oxygen effects. The Corrosion rates at low O_2 content are about 0.0071 mm/yr with no CO_2 and 0.0083 mm/yr with 0.6 mg/l CO_2 ; a small difference between two small rates. At 10 mg/l O_2 the corrosion rate is 0.16 mm/yr, and it is the same with and without CO_2 . Thus, CO_2 has no effect on corrosion at high O_2 concentrations meaning that the surface is corroded by oxygen; small concentrations of CO_2 does not change this rate.

Sixth operation condition: 300 cycles/min flow, tank open to the atmosphere (aerated with O₂ and CO₂) at ambient conditions

Lastly, you will look at the effects of shear rates on the tank in contact with CO2 and O2.

Copy (right-mouse click or Ctrl+C) 'Static Flow, O2 and CO2' corrosion calculation, and **paste** (right-mouse click or Ctrl+V) on the stream 'CR vs Flow'





Change the Static Flow, O2 and CO2-1 name to 300 rpm flow, O2 and CO2 using the <F2> key

Add Oxygen (O2) and carbon dioxide (CO2) to the inflow grid

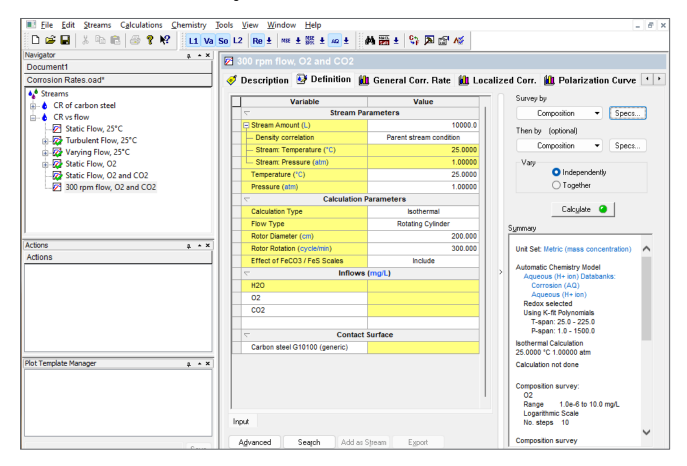
Change the Flow Type from Static to Rotating Cylinder

Set the Rotor Diameter to 200 cm

Set the Rotor Rotation to 300 cycles/min

Leave the default alloy Carbon steel G10100 (generic) as the contact surface

Your screen should look like the image below.

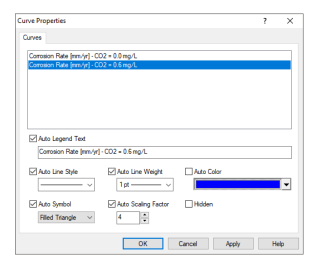


We are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Corrosion Rates*.

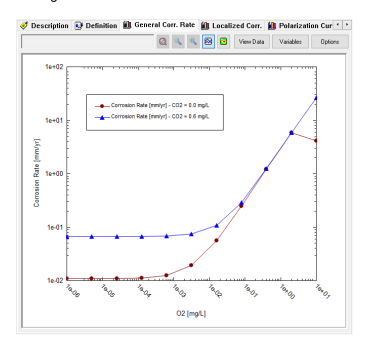
Analyzing the Results

Click on the **General Corr. Rate** tab (General Corr. Rate). This tab displays a plot showing the results of corrosion rate and pH.

Right-mouse click on the X-axis and change it to Logarithmic Scale
Right-mouse click on the Y-axis and change it to Logarithmic Scale
Right-mouse click on the pH curve and select Remove Series
Double click on the "CO2 = 0.6 mg/L" curve. This will open a new window. Change the color to blue.



The plot should look like the image below.



The 0.6 mg/L CO_2 curve shifted to higher values at low O_2 concentrations compared to no CO_2 presence. Now, when comparing the corrosion rate for the 0.6 mg/L CO_2 concentration at 300 rpm vs static conditions, the rates are 0.067 mm/year and 0.008 mm/year, respectively. The effect of shear at high O_2 concentrations (right side of the plot) is also pronounced. Corrosion is still dominated by O_2 attack, but the rate is now over 10 mm/year, about 100x greater than the static conditions.

Gas condensate corrosion

Example 51: Condensed overhead gas and mitigation strategies

An alkanolamine gas sweetening plant has corrosion problems in the condensed overhead gas.

Diethanolamine is used to neutralize an acid gas containing carbon dioxide (CO_2) and hydrogen sulfide (H_2S) . The diethanolamine is regenerated and the acid gases are driven off in a stripper. The off gas from this stripper is saturated with water vapor. As these gases cool, they will condense. This condensate can be very corrosive. The plant's service life can be shortened considerably due to these condensed acid gases.

In this example, you will calculate the gas dew point temperature, remove the condensed aqueous phase, and perform a Corrosion Rate calculation with the condensed water. Lastly, you will consider mitigation strategies for the pipes.

You are introducing *fluid velocity* and *liquid condensation* into the calculation. The software uses a diffusion layer model to compute mass transfer to and from corroding surfaces. Higher rates produce thinner layers, resulting in faster mass transfer rates, and thus higher corrosion rates. The liquid condensation point is straightforward; it calculates the temperature (or pressure) where the first liquid drop forms.

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc., will require further adjustments, and will be described as necessary.

| Corrosion Rate Calculation | | | |
|----------------------------|-------------------------------|-----------------------------------|--------------------|
| Calculation Settings | | Stream Composition and Conditions | |
| Stream Name | Gas condensate | Stream Amount | 1e5 moles |
| Name Style | Display Formula | Temperature | 38 °C |
| Unit Set | Metric, Batch, Mole Fraction | Pressure | 1.2 atm |
| Framework | AQ | H2O | Calculated (mole%) |
| Calculation Type | Corrosion Rates | CO2 | 77.4 |
| Survey | Single Point Rate | N2 | 0.02 |
| Flow Type | Pipe Flow | H2S | 16.6 |
| Pipe Diameter | 10 cm | CH4 | 0.5 |
| Pipe Flow Velocity | 2 m/s | C2H6 | 0.03 |
| Contact Surface | Carbon Steel G10100 (generic) | C3H8 | 0.03 |

Add a new Stream

Click on the new Stream and press <F2> to change the name to Gas Condensate

Select the AQ thermodynamic Framework

Click on the Names Manager Icon and select the Formula option

Click on the Units Manager Icon, and select Metric, Batch, Mole Fraction

Enter the composition, temperature, and pressure of the stream given in the table above

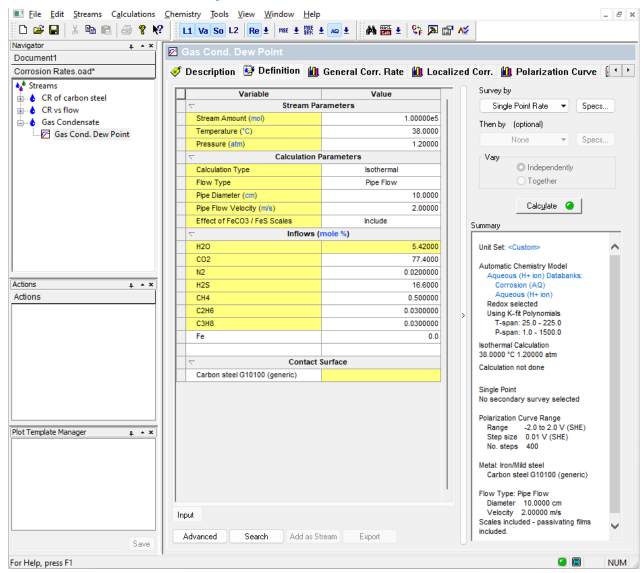
Go to the **Add Calculation** button and select **Corrosion Rates** calculation or select the **Add Corrosion Rates** icon in the Actions Pane.

Go to the **Survey by** option and select **Single Point Rate**. Change the name to **Gas Cond. Dew Point**. **Change** the Flow Type from Static to **Pipe Flow**.

Specify the Pipe Diameter = 10 cm and the Pipe Flow Velocity = 2 m/s.

Leave the default alloy Carbon steel G10100 (generic) as the contact surface

Your screen should look like the image below.

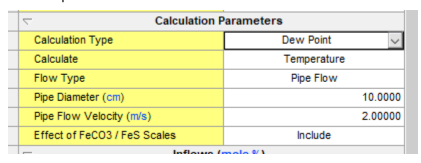


Now that the gas condensate stream is created, the next task is to isolate the condensed water at the dew point temperature.

Click on the Specs button. This will open a new window

Under the Category section, select Calculation Type

Change the Type of Calculation from Isothermal to **Dew Point.** Make sure that radio button for temperature is selected since we are going to calculate the Dew Point temperature for this example. Then click **OK**. Notice that the input grid has been updated:



You could have also used the drop-down arrow next to the "Calculation Type" box to directly pick a dew point (or any other required) calculation type.

Now, we are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Corrosion Rates*.

Analyzing the Results

Click on the General Corr. Rate tab (General Corr. Rate).

Click on the **View Data** button to see the results in tabulated form.

| | Temperature | Corrosion Rate | pН |
|---|-------------|----------------|---------|
| | °C | mm/yr | |
| 1 | 37.5851 | 0.704502 | 3.92719 |

Notice that the calculated dew temperature is 37.6°C, corrosion rate is 0.7 mm/year, and the dew point pH is 3.9. To further interpret the results of this calculation, we are going to study the polarization curve.

Click on the **Polarization Curve** tab (Polarization Curve). Before interpreting this plot, we will format it for easy viewing.

Click on the **Options** button. This will open a new window.

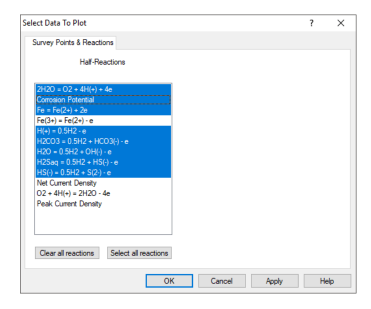


Under the **Category** section select **X** axis. Change the **Minimum** to 1e-6 and **Maximum** to 1e6. Now, select **Y** axis. Change the **Minimum** to -1.5 and **Maximum** to 1.5. Then **click OK**.

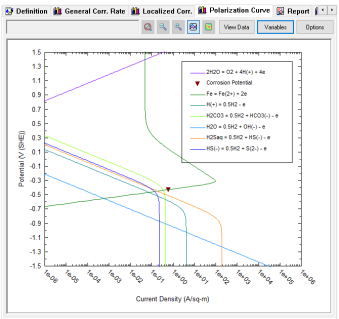
Click on the **Variables** button. This will open a new window.



Remove the following variables by unchecking them. See image below. Then Click OK.



Your polarization curve should look like the image below.



You can click and drag the legend of the polarization curve to your desired position.

This is a measured potential vs. current density plot. The anodic portion (green curve) is the carbon steel. There are cathodic reactions. The sum of the cathodic currents adds up to the anode curve at a single point: The corrosion potential (represented by the inverted red triangle ▼) with a value of -0.43 V vs SHE. At the corrosion

potential, the anodic and cathodic currents are equal, and this value is also known as the i_{corr} . For this example, i_{corr} =0.61 A/m².

Notice that the cathodic reaction that contributes the most to corrosion is H_2CO_3 (light green line). The next contributors are H_2S (orange line) and HS^{-1} (blue line) equally. After that is H^+ (turquoise line). Water reduction (light blue line), contributes the least to corrosion.

Mitigation Strategies

There are several mitigating solutions to this corrosion problem.

One is adding insulation to prevent temperature drops. The dew point is very close to the overhead gas temperature so this may not be a suitable option. Adding heat to keep the temperature above the dew point is usually considered along with insulation.

Changing the chemistry to change the partial oxidation and reduction processes is also an option.

Changing alloys could mitigate the corrosion problems as well.

We will explore the change of chemistry and the change of alloys for this example.

Adjusting the solution chemistry

The condensate pH is approximately 3.9. We can try to add a base to increase the pH. In this section, we will add Diethanolamine to raise the pH to 7.5.

Single Point- Set pH Calculation

Click on the 'Gas Condensate' stream and **go** to the **Add Calculation** button and select **Single Point** calculation or select the **Add Single Point** icon in the Actions Pane.

Change the name to **DEA** using the **<F2>** key.

Add DEA in the inflows grid. (It should have changed to the formula name: HN(C2H4OH)2

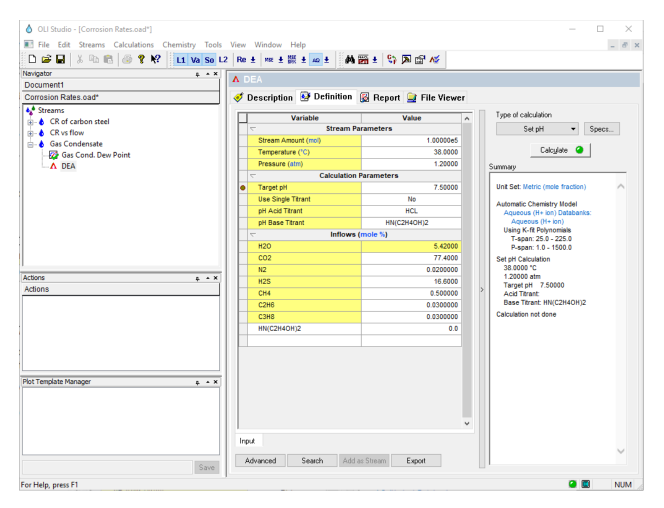
Go to the Type of Calculation option and select Set pH

Change the Target pH to 7.5

Click on the drop-down arrow for the pH Base Titrant and select DEA (or HN(C2H4OH)2)

We have set up the calculation to adjust the amount of DEA to match the target value of pH=7.5.

Your screen should look like the image below.

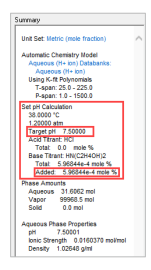


Now, we are ready to perform the calculation. Click on the Calculate button or press the <F9> key

It is time to **save** your file **(File >Save as...) or** use the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Corrosion Rates*.

Analyzing the Results

Check the summary box to analyze the results.



The pH is set to 7.5 and the inflow of DEA to the total gas is approximately 5.97e-4 mole%.

The DEA concentration in the condensing water and the amount remaining in the vapor phase can both be viewed in the Report tab.

Click on the Report Tab and go to the Total and Phase Flows and Species Output tables

Species Output (True Species) Row Filter Applied: Only Non Zero Values

column Filter Applied: Only Non Zero Values

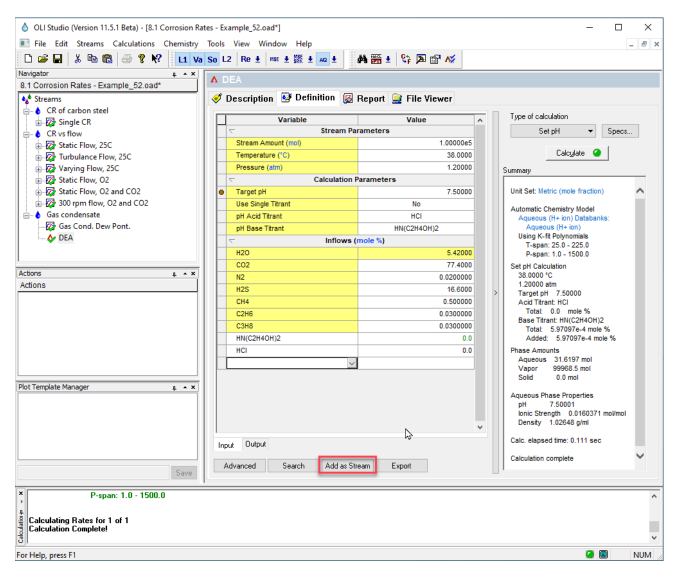
| Total and Phase Flows (Amour | nts) |
|---|------|
| column Filter Applied: Only Non Zero Values | |

| | Total | Aqueous | Vapor |
|-------------|-----------|----------|-----------|
| | mol | mol | mol |
| Mole (True) | 1.00000e5 | 31.6062 | 99968.5 |
| Mole (App) | 1.00001e5 | 32.0477 | 99968.5 |
| | g | g | g |
| Mass | 4.08064e6 | 642.188 | 4.08000e6 |
| | L | L | L |
| Volume | 2.11456e6 | 0.625619 | 2.11456e6 |

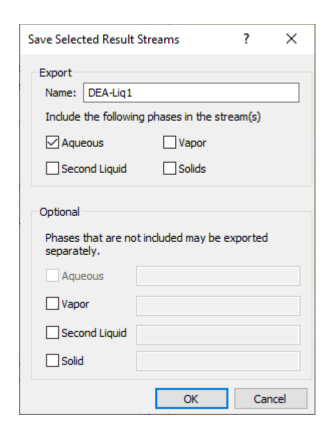
| | Total Aqueous | | Vapor |
|------------------|---------------|------------|------------|
| | mole % | mole % | mole % |
| CO2 | 77.3994 | 0.0416672 | 77.4239 |
| H2S | 16.5999 | 0.0271211 | 16.6052 |
| H2O | 5.41962 | 96.6827 | 5.39076 |
| CH4 | 0.499999 | 1.37153e-5 | 0.500157 |
| C2H6 | 0.03 | 9.08534e-7 | 0.0300094 |
| С3Н8 | 0.03 | 7.06434e-7 | 0.0300094 |
| N2 | 0.02 | 2.56549e-7 | 0.0200063 |
| NH2(C2H4OH)2(+1) | 5.02996e-4 | 1.59145 | |
| HCO3-1 | 3.689e-4 | 1.16718 | |
| C5H10NO4-1 | 6.87352e-5 | 0.217474 | |
| HS-1 | 5.7612e-5 | 0.182281 | |
| HN(C2H4OH)2 | 2.51153e-5 | 0.0778779 | 5.01249e-7 |
| CO3-2 | 3.86988e-6 | 0.0122441 | |
| S-2 | 3.93925e-9 | 1.24636e-5 | |
| OH-1 | 6.55471e-10 | 2.07387e-6 | |
| H+1 | 2.47871e-11 | 7.8425e-8 | |
| Total (by phase) | 100.0 | 100.0 | 100.0 |

Recalculating the corrosion rate with DEA

We first need to capture the condensate from the DEA pH calculation. We want only the liquid portion to perform our corrosion rate calculation. This would represent the first drop of condensate from the gas stream.



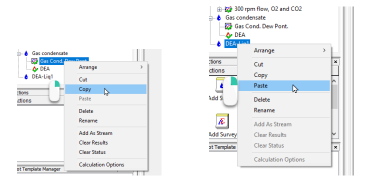
Click the Add as Stream button



Uncheck the following boxes: **Second Liquid, Vapor**, and **Solid.** This removes these phases from the saved stream. Click the **OK** button

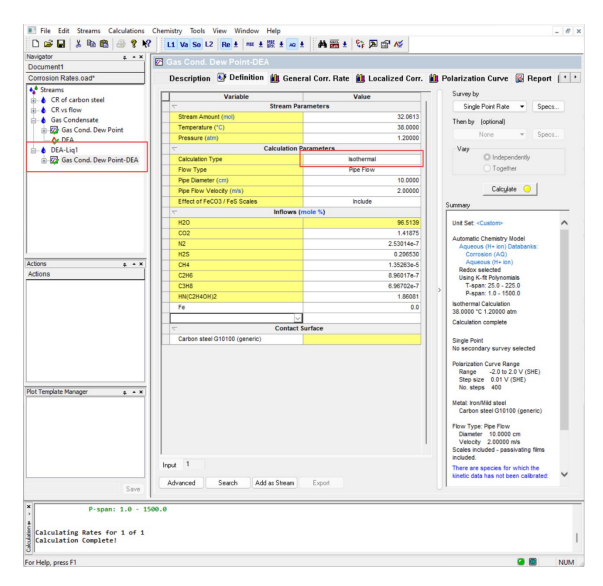
We now can recalculate the corrosion rate with DEA in the stream. This represents some DEA entrained in the vapor phase as the gas leaves the regenerator.

Copy 'Gas Cond. Dew Point' corrosion calculation from the <u>Gas condensate</u> stream, and <u>paste</u> it on the stream '<u>DEA-Liq1</u>' so that it appears under <u>'DEA-Liq1.'</u>



Change the name to *Gas Cond. Dew Point DEA* using the <F2> key The added DEA is already in the inflows gird. Leave the default alloy **Carbon steel G10100 (generic)** as the contact surface Change the calculation type to **Isothermal**

Your screen should look like the image below.



Now, we are ready to perform the calculation. Click on the Calculate button or press the <F9> key

It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Corrosion Rates*.

Analyzing the Results

Click on the General Corr. Rate tab (General Corr. Rate).

Click on the View Data button to see the results in tabulated form.

| | Temperature | Corrosion Rate | pН |
|---|-------------|----------------|---------|
| | °C | mm/yr | |
| 1 | 38.0000 | 0.480601 | 7.50001 |

Notice that the corrosion rate has decreased from 0.7 mm/year to 0.48 mm year. The pH is 7.6, and it remained like the target value of 7.5. Based on these results, neutralizing the pH had a partial effect on corrosion reduction.

Changing the Alloy

Since treating the acid gas with a base is probably not a good idea for metal hydroxides, perhaps we can change the alloy. We will add a new corrosion rates calculation and test different alloys.

13%Cr Stainless Steel

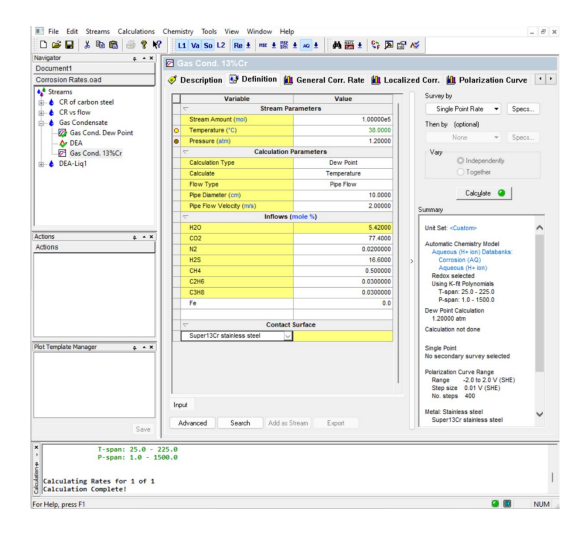
Copy 'Gas Cond. Dew Point' corrosion calculation, and paste it under the stream 'Gas Condensate'

Change the name to *Gas Cond. 13%Cr* using the <F2> key.

Notice that in the inflows, copying and pasting does not bring the DEA and its composition to this corrosion calculation. The reason why is because the DEA was added at the corrosion rate calculation level, and not at the stream level.

Change the contact surface to Super13%Cr stainless steel

Your screen should look like the image below.



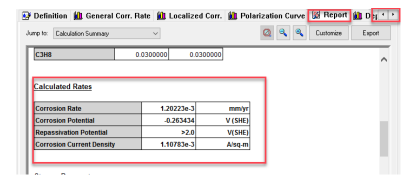
Now, we are ready to perform the calculation. Click on the Calculate button or press the <F9> key

It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Corrosion Rates*.

Analyzing the Results

Click on the **Report** tab to see the results. (You may need to use the ◀▶ buttons to move the tabs and find the Report tab).

Go to the Calculated Rates table



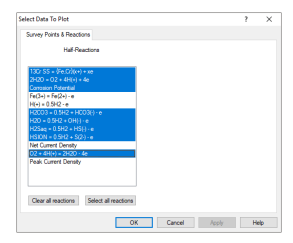
The corrosion rate of Super 13% Cr is several orders of magnitude lower when compared to the corrosion rate for Carbon Steel, i.e., at 0.0012 mm/year for 13%Cr vs 0.7 mm/year for Carbon Steel. This is consistent with the use of Super 13% Cr to protect against CO₂ corrosion.

Click on the **Polarization Curve** tab (Polarization Curve).

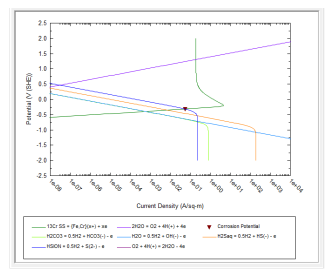
Click on the Variables button. This will open a new window.



Remove the following variables by unchecking them as See image below. Then Click OK.



The polarization curve (after some layout modifications) should look like the image below.



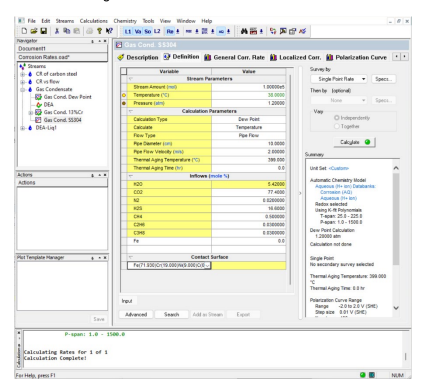
The layout of the plot was changed for easy reading. To change layout, you can rightclick on any part of the plot and select the option *Allow Layout Changes* to modify the appearance of your plot.

The corrosion potential of Super 13%Cr is -0.25 V vs SHE and the corresponding corrosion current i_{corr} =0.001A/m². The i_{corr} for Carbon Steel was 0.61 A/m², this means that 13%Cr is 10x more resistant to corrosion under the same conditions.

Stainless Steel 304

Copy 'Gas Cond. Dew Point' corrosion calculation, and paste it under the stream 'Gas Condensate' Change the name to Gas Cond. SS304 using the <F2> key Change the contact surface to Stainless Steel 304

Your screen should look like the image below.



Now, we are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar. You can save it under the same file created in the previous example named *Corrosion Rates*.

Analyzing the Results

Click on the **Report** tab to see the results. (You may need to use the ◀▶ buttons to move the tabs and find the Report tab).

Go to the Calculated Rates table

Calculated Rates

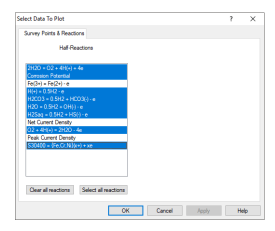
| Corrosion Rate | 2.40183e-3 | mm/yr |
|---------------------------|------------|---------|
| Corrosion Potential | -0.293505 | V (SHE) |
| Repassivation Potential | >2.0 | V(SHE) |
| Corrosion Current Density | 2.21614e-3 | A/sq-m |

The corrosion rate of 304 SS is now 0.0024 mm/year. This is a negligible value.

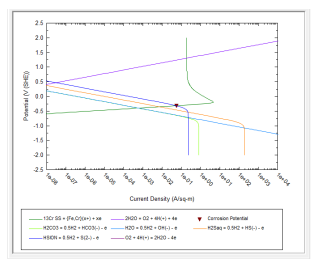
Click on the **Polarization Curve** tab (Polarization Curve).

Click on the Variables button. This will open a new window.

Remove the following variables by unchecking them. See image below. Then Click OK.



The polarization curve (after some layout modifications) should look like the image below.



The layout of the plot was changed for easy reading. To change layout, you can right-click on any part of the plot and select the option Allow Layout Changes to modify the appearance of your plot.

The corrosion potential of SS304 is -0.17 V vs SHE and the corresponding corrosion current icorr=3.3e-3 A/m². With this low corrosion current density, sour gas regeneration can be remedied using a higher-grade alloy such as SS304.

Section 9. Thermal Aging and Variations in Alloy Composition

Overview and Basic Terminology

Localized corrosion of engineering alloys is a complex function of metallurgical factors and environmental conditions. Among metallurgical factors, effects of thermal instabilities are of interest for assessing the performance and expected service life of industrial components fabricated from nominally corrosion-resistant stainless steels and nickel-base alloys.

Fabrication processes such as heat treatment and welding are known to introduce microstructural changes that may affect both the mechanical performance and corrosion resistance of an alloy. In particular, thermal instability of stainless steels and nickel-base alloys may lead to the formation of complex metal carbides of the type M₃C₂, M₇C₃, M₆C, or M₂₃C₆ in which the metallic component M represents Cr, Mo, W, and Fe. The carbide is chromium- or molybdenum-rich depending on the carbide type, which in turn depends on the alloy composition and temperature. Also, various chromium-rich intermetallic phases can form in many alloys. Precipitation of such phases may occur at temperatures ranging from 500 to 900 °C depending on alloy composition. Formation of grain boundary carbides often results in the depletion of chromium and, possibly, molybdenum in the vicinity of the grain boundary because of the slow diffusion of substitutional elements such as chromium relative to the interstitial carbon.

Similarly, the corrosion resistance of welded components may be affected by the segregation of alloying elements and precipitation of intermetallic phases, carbides, or nitrides in the solidified weld and unmixed zones as well as the precipitation of carbides and other phases in the heat-affected zone adjacent to the weld.

Sensitization of Fe-Ni-Cr-Mo alloys and its effects on intergranular attack and intergranular stress corrosion cracking is the most directly observed effect of Cr depletion. It may result in intergranular attack and intergranular stress corrosion cracking. Localized corrosion can be also affected by Cr and Mo depletion.

Corrosion Analyzer contains the following technology that can help address these issues:

- A grain boundary microchemistry model for predicting the chromium and molybdenum depletion in the vicinity of grain boundaries as a result of carbide formation.
- An electrochemical model for calculating the repassivation potential of Fe-Ni-Cr-Mo-W alloys as a function of alloy composition and environmental conditions including temperature and concentrations of aqueous solution species.
- A procedure for calculating the observable repassivation potential that corresponds to macroscopic localized corrosion by applying the electrochemical model to the depletion profiles and performing suitable integration.

More details about this technology are described by Anderko et al. (2008), Tormoen et al. (2009), Anderko et al. (2009), and Sridhar et al. (2009).

Alloy Chemistry

Simulations can be performed, in general, for alloys that belong to the Fe-Ni-Cr-Mo-W-N-C family (i.e., for stainless steels and nickel-base alloys).

Depletion profiles in the vicinity of grain boundaries and depletion parameters can be obtained for austenitic alloys (including stainless steels and Ni-base alloys). Also, the effect of Cr and Mo depletion on localized

corrosion can be calculated. This effect can be examined using the repassivation potential, which provides a threshold potential for the stabilization of localized corrosion (Anderko et al., 2009).

For other alloys from the Fe-Ni-Cr-Mo-W-N-C family, the repassivation potential can be calculated if the alloy composition is known. This also includes experimental alloys and separate phases that may be formed as a result of various forms of heat treatment (Sridhar et al., 2009).

Calculation types

Thermal aging is an additional phenomenon that can be simulated within the framework of corrosion kinetics. All calculation types and, in particular, survey types, that are supported for corrosion kinetics are also supported in conjunction with the study of thermal aging.

If it is desired to make calculations on a thermally aged sample, the thermal aging temperature and time need to be specified in the **Calc Parameters** section in the **Definition** tab. The default values are 399 °C for the thermal aging temperature and 0.0 hours for the thermal aging time. If either of these default values is used, no thermal aging effects will be predicted. Thus, by default, Corrosion Analyzer performs calculations on samples that have not been thermally aged.

Thermal Aging Temperature Survey

This calculation makes it possible to vary the thermal aging temperature within a certain range. A fixed value of thermal aging time is assumed as specified by the user. Typically, the temperatures for which thermal aging effects can be observed range from ~500 °C to ~900 °C, with the effects being most pronounced in the middle of this range.

Thermal Aging Time Survey

This calculation can be used to examine the effect of aging time at a fixed thermal aging temperature.

Output Specific to Thermal Aging

The following output can be generated:

Chromium and molybdenum depletion profiles, i.e., the variation of Cr and Mo concentration within the grain as a function of grain boundary.

The depletion parameter, which provides compact information on the extent to which the depletion process reduces the grain boundary concentration below a certain critical value of, which can be defined by well-known criteria for maintaining passivity (e.g. $x_{Cr}^* = 0.11 \ or \ 0.12$). This parameter can be calculated as the area of the depletion profile below the threshold concentration x_{Cr}^* , divided by bulk Cr concentration, x_{Cr}^0 :

$$\delta(x_{Cr}^*) = \frac{1}{x_{Cr}^0} \int_0^{z^*} (x_{Cr}^* - x_{Cr}(z)) dz$$

where z^* is the distance from the grain boundary that corresponds to the threshold concentration x_{Cr}^* .

The repassivation potential, which is a key parameter for determining whether localized corrosion can occur, may be affected by thermal aging in a rather complex way.

Thermal Aging and Variations in Alloy Composition

We will show examples of simulating the effects of thermal aging on Fe-Ni-Cr-Mo-W-C-N alloys (i.e., stainless steels and nickel-base alloys). Specifically, we will show how to predict:

Chromium and molybdenum depletion profiles in the vicinity of grain boundaries, which result from heat treatment of austenitic alloys.

Depletion parameters for sensitized austenitic alloys, which provide an indicator of whether the alloy is susceptible to intergranular corrosion.

Effect of thermal aging on the repassivation potential of austenitic alloys, which provides a threshold condition for localized corrosion (pitting or crevice corrosion).

The repassivation potential of alloys with compositional variations that may or may not result from thermal aging. This facility can also be used for bulk alloys that are not in the database or for hypothetical or experimental alloys as long as they belong to the Fe-Ni-Cr-Mo-W-C-N family. An example will be given for a duplex alloy, either annealed or thermally treated.

Example 52: Thermal aging of alloy 600

We will be studying the behavior of thermally aged alloy 600 in a dilute aqueous solution of sodium chloride, and will simulate how the time of thermal aging affects alloy 600 at a fixed thermal aging temperature of 700°C.

Starting the Simulation

Thermal Aging Calculation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Thermal Aging Calculation | | | |
|---------------------------|----------------------|-----------------------------------|-------------------------|
| Calculation | on Settings | Stream Composition and Conditions | |
| Stream Name | Thermal Aging | Stream Amount | Calculated |
| Calculation Type | Corrosion Rates | Temperature | 60 °C |
| Survey by | Thermal Aging Time | Pressure | 1 atm |
| Name Style | Display Name | H2O | Default – 55.5082 moles |
| Unit Set | Metric, Batch, Moles | NaCl | 0.04 moles |
| Framework | AQ | | |
| | | | |

Add a new Stream

Click on the new Stream and press <F2> to change the name to Thermal Aging

Select the AQ thermodynamic Framework (selected by default)

Click on the Units Manager Icon and select Metric, Batch, Moles

Click on the Names Manager Icon and select Display Name

Enter the stream composition and conditions specified in the table above

Go to the Add Calculation button and select Corrosion Rates

Note: Even though the thermal aging simulation facilities do not predict corrosion rates per se, they belong to the category of corrosion kinetics and, therefore, they are included in the Corrosion Rates section of the Corrosion Analyzer.

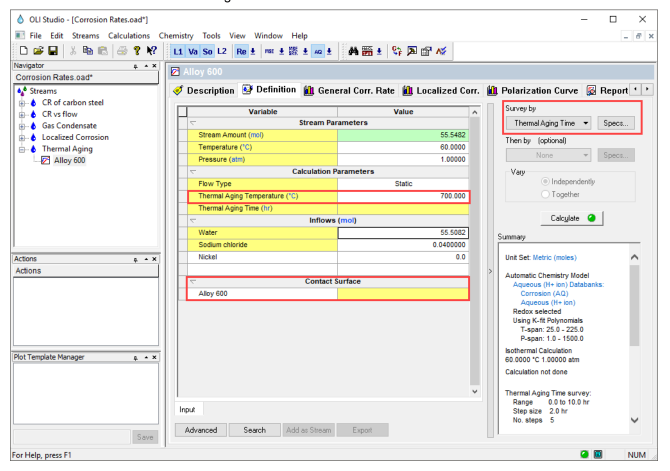
Select the Survey by Thermal Aging Time

Change the name from *Rates* to *Alloy 600* using the <F2> key or by right-mouse clicking on the object and select Rename

Select Alloy 600 from the Contact Surface grid

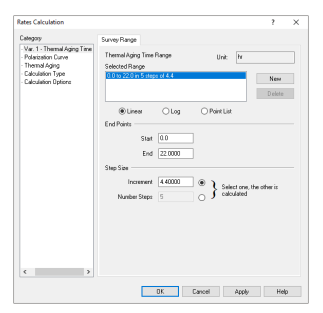
In the **Calculation Parameters** grid, change the default value for the **Thermal Aging Temperature** to 700°C. Note that the default value for the **Thermal Aging Temperature** is 399 °C, which is a low value so that, in the default case, no effects of thermal aging are being calculated.

Your screen should look like the image below.

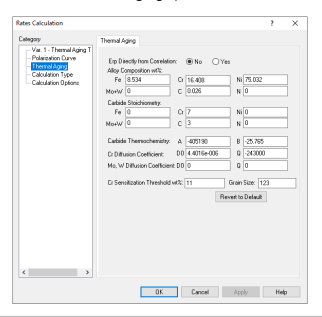


Now we need to specify the Thermal Aging Time range. Click on the Specs button. This will open a new window.

Under the **Survey Range** tab, change the range for Thermal Aging Time from 0 to 22 hours with an increment of 4.4 hours (of 5 steps).



Under the Category section, select the Thermal Aging option.



In this **Thermal Aging** Category, you can change all parameters that are necessary for calculating chromium and (if applicable) molybdenum depletion profiles. You can change:

- The alloy composition (for example, to analyze the effect of different carbon content);
- The stoichiometry of the carbide phase that may form at the grain boundary; the typical carbide stoichiometry is M_7C_3 or $M_{23}C_6$ (where M = Cr, Mo) but can be adjusted
- The parameters that define the equilibrium constant for the formation of the carbide
- The diffusion coefficient of Cr and, if applicable, Mo
- The threshold concentration of Cr for sensitization. This threshold concentration is used for calculating the depletion parameter.
- The average grain size, which affects the process of healing of chromium depletion as a function of time

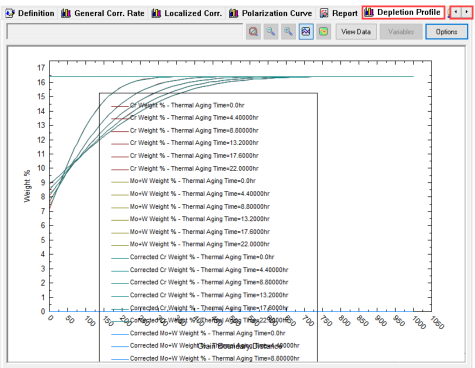
Also, you can indicate whether the repassivation potential (E_{rp}) should be calculated directly from the correlation developed by Anderko et al. (2008), which expresses E_{rp} as a function of alloy composition or not. If it is not calculated directly, then the repassivation potential is calculated first using the alloy-specific parameters for the alloy of interest and only the decrement of E_{rp} (i.e., E_{rp} , thermally aged $-E_{rp}$, bulk alloy) is obtained from the correlation. This decrement is then added to the alloy-specific E_{rp} value. This option is set by default to "No" because this maximizes the accuracy of calculations for alloys that are already in the database.

For now, we will accept the default settings. Click OK.

Now, we are ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar. You can save it under the same file created in the previous section named *Corrosion Rates*.

Analyzing the Results

Click on the **Depletion Profile** tab (Depletion Profile). If you don't see it in your screen, use the ► buttons to move the tabs.



Note: Since alloy 600 does not contain any molybdenum, the molybdenum curves will always be equal to zero.

This plot will show the concentrations of chromium and molybdenum within a grain as a function of the distance from the grain boundary (in μ m). You will see that, for each condition, there are four lines:

Cr weight %: concentration of Cr in weight %

Mo+W weight %: sum of the concentrations of molybdenum and tungsten in weight %

Corrected Cr weight %: concentration of Cr corrected for beam scattering and related effects so that it can be directly compared with experimental results. The procedure for calculating the correction is described by Anderko et al. (2009).

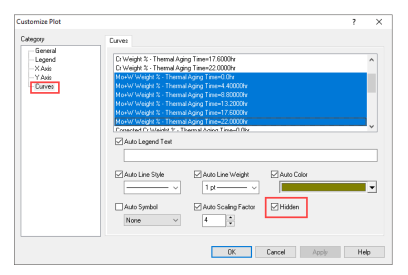
Corrected Mo+W weight %: sum of the concentrations of Mo and W corrected in the same way as those for Cr.

Let's customize the plot

Click on the **Options** button. This will open a new window.

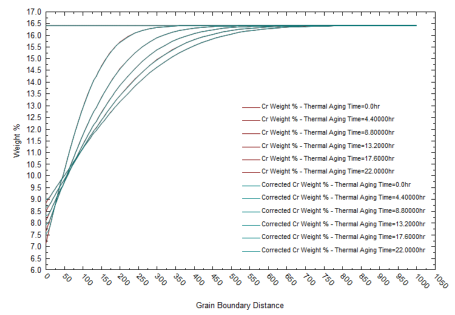
Since alloy 600 does not contain Mo or W, eliminate the Mo+W curves. To do this, go to the **Category** section and select the **Curves** option

Highlight the Mo+W weight % curves by pressing the Shift key and the down key (**Shift** $+ \downarrow$). After highlighting these curves, check the **Hidden** button.



After hiding the Mo+W weight % curves, do the same for the Corrected Mo+W weight % curves. Then click **OK**.

The plot should look like the image below.



The legend of this plot was modified for clarity. The border style of the legend was modified by double-clicking on the legend, and selecting *None* as the Border Style

Differences between Uncorrected and Corrected Depletion Profiles

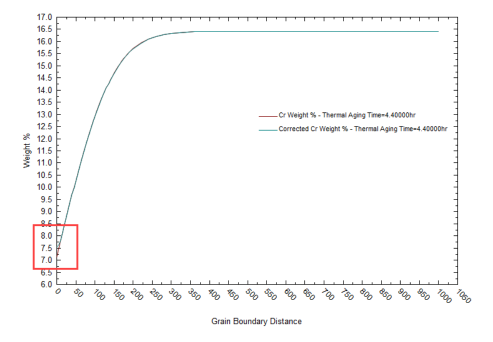
Now, let's focus on the differences between the uncorrected and corrected depletion profiles. To have a clear picture of the difference between them, we will analyze the depletion profiles results obtained at 4.4 hours of thermal aging.

Click on the **Options** button, and select the **Curves** category
Highlight all curves (hold CTRL and left-click to highlight/unhighlight) except the ones labeled as:

Cr Weight % - Thermal Aging Time = 4.40000 hr, and Corrected Cr Weight % - Thermal Aging Time = 4.40000 hr

Check the Hidden box, then click OK.

Your plot should look that the image below.



In the obtained plot, you can see the difference between the two curves for small distances from the grain boundary (roughly below 30 μ m). The corrected concentration is much more rounded in the vicinity of the minimum because of instrumental measurement effects. At larger distances from the grain boundary, there is no difference between the two curves.

Effect of Cr depletion on corrosion

After analyzing the depletion profiles, let's focus on the effects of Cr depletion on corrosion. We will do it by analyzing two parameters – the depletion parameter and the repassivation potential.

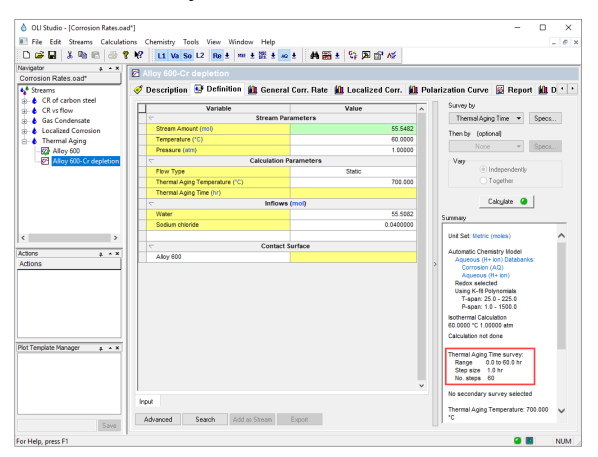
To have more meaningful results, let's expand the range of thermal aging time and reduce the interval for calculations.

Copy the Alloy 600 corrosion calculation, and then paste it under the Thermal Aging Stream Rename the new calculation as Alloy 600 – Cr depletion

Go back to the **Definition** tab and click the **Specs** button

In the **Var.1 Thermal Aging Time** category, change the Survey Range as follows: **Start**=0 hours, **End**= 60 hours with an **Increment**= 1 hour. Then click **OK**.

Your screen should look like the image below

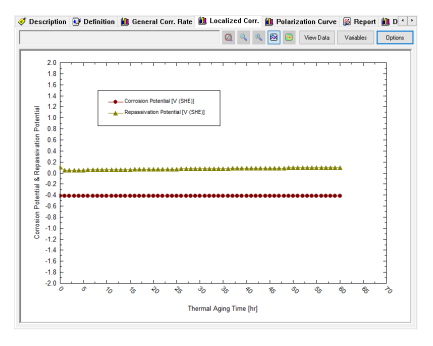


Now, we are ready to perform the calculation. Click on the Calculate button or press the <F9> key

It is time to **save** your file **(File >Save as...)** or using the **save** icon in the tool bar. You can save it under the same file created in the previous section named *Corrosion Rates*.

Analyzing the Results

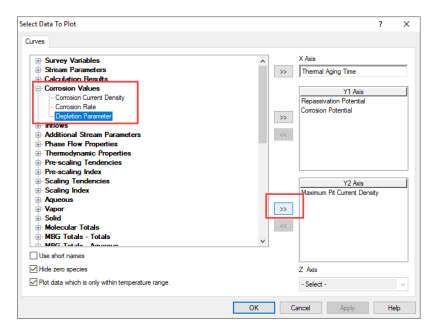
Click on the **Localized Corr.** tab (Localized Corr.). By default, this tab will show a plot of the corrosion potential and repassivation potential.



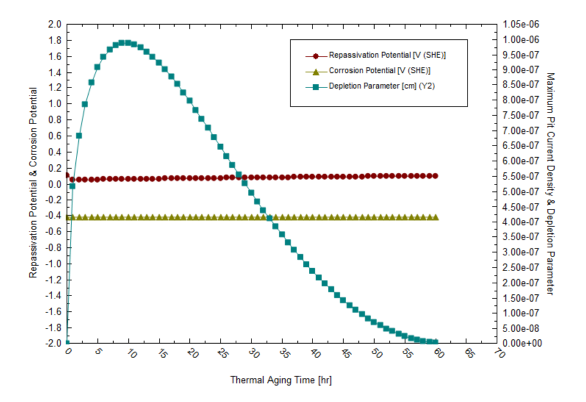
Let's add a new parameter to the plot: The **Depletion Parameter**.

Click on the **Variables** button. This will open a new window.

Look for the **Corrosion Values** section, click on the '+' icon, and select the **Depletion Parameter** option. Click on the >> button that belongs the **Y2 Axis**. This action will place the new variable under the Y2 Axis. Then click **OK**.



The plot should look like the image below.



It is clear that the depletion parameter reaches a maximum for aging time of about 10 hours. Since the depletion parameter is a measure of the susceptibility of an alloy to intergranular corrosion, we can expect that the alloy will be most susceptible to intergranular corrosion at intermediate aging times. When the depletion parameter is zero, intergranular corrosion or intergranular stress corrosion cracking are unlikely. The decrease of the depletion parameter as a function of time is a manifestation of the phenomenon of healing of Cr depletion.

The repassivation potential shows a minimum as a function of aging time (see the option **View Data**). This indicates that the tendency of the alloy to undergo localized corrosion is enhanced as a result of thermal aging. However, the effect of thermal aging on the repassivation potential of alloy 600 is small (cf. Tormoen et al., 2009, Anderko et al., 2009). The repassivation potential shows a minimum at low aging times (ca. 1-2 hours). Therefore, the susceptibility to localized corrosion is enhanced the most for these aging times. It is noteworthy that the maximum in the depletion parameter does not coincide with the minimum in the repassivation potential. This is due to the fact that intergranular corrosion (which is related to the depletion parameter) and localized corrosion (which is controlled by the repassivation potential) are subject to different mechanisms. A general discussion of these differences is given by Tormoen et al. (2008).

In general, the alloy will be susceptible to localized corrosion if the corrosion potential exceeds the repassivation potential. In the above example, the corrosion potential is low because we have no oxidizing agents in the system. Therefore, the alloy will not undergo localized corrosion at the conditions of this example. However, a rise in the corrosion potential due to the presence of oxidizing agents may cause localized corrosion.

Example 53: Thermal aging of alloy 825

Alloy 825 is appreciably different from alloy 600 because alloy 825 contains molybdenum and, also, substantially more chromium in addition to other alloying elements.

In this example, we will simulate how the temperature of thermal aging affects alloy 825 at a fixed thermal aging time of 15 hours.

Starting the Simulation

Thermal Aging Calculation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Thermal Aging Calculation | | | | |
|---------------------------|---------------------------|-----------------------------------|-------------------------|--|
| Calculation Settings | | Stream Composition and Conditions | | |
| Stream Name | Stream Name Thermal Aging | | Calculated | |
| Calculation Type | Corrosion Rates | Temperature | 95 °C | |
| Survey by | Thermal Aging Temperature | Pressure | 1 atm | |
| Name Style | Display Name | H2O | Default – 55.5082 moles | |
| Unit Set | Metric, Batch, Moles | NaCl | 2.846e-3 moles | |
| Framework | AQ | | | |
| | | | | |

Under the Thermal Aging stream add a new calculation rate. Go to the **Add Calculation** button and select **Corrosion Rates** or select the **Add Corrosion Rates** icon in the **Actions panel**

Change the name from *Rates* to *Alloy 825* using the <F2> key or by right-mouse clicking on the object and selecting rename

Select the Survey by Thermal Aging Temperature option

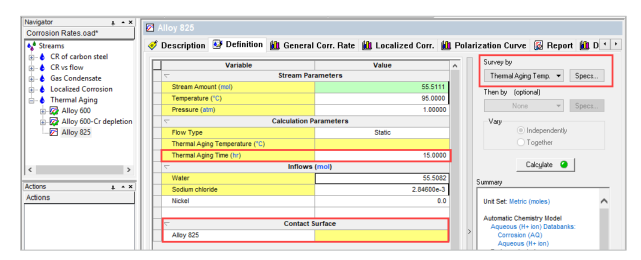
Change the stream composition and conditions to the specified values in the table above

Select Alloy 825 from the Contact Surface grid

In the Calculation Parameters grid, change the default value for the Thermal Aging Time to 15 hours

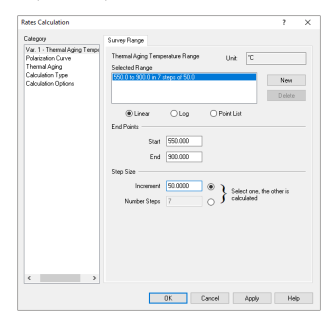
Note that the default value for the **Thermal Aging Temperature** is 0 hours, which means that no thermal aging is considered by default.

Your screen should look like the image below



Now we need to specify the Thermal Aging Temperature range. **Click** on the **Specs** button. This will open a new window.

Under the **Survey Range** tab, change the range for **Thermal Aging Temperature** from 550 to 900 °C with increments of 50 °C. We will keep the other parameters at their default values. Click **OK.**

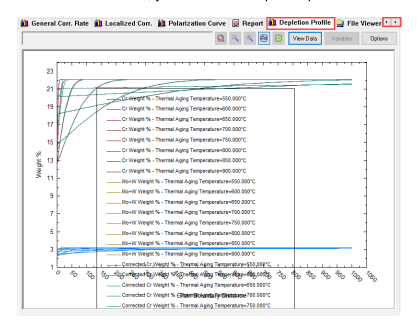


We are now ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar. You can save it under the same file created in the previous section named *Corrosion Rates*.

Analyzing the results

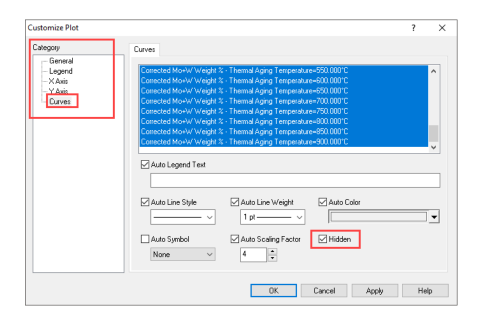
Click on the **Depletion Profile** tab (Depletion Profile). If you don't see it in your screen, use the ► buttons to move the tabs.

Since alloy 825 contains Mo in addition to Cr, you will see the depletion profiles for both Cr and Mo.

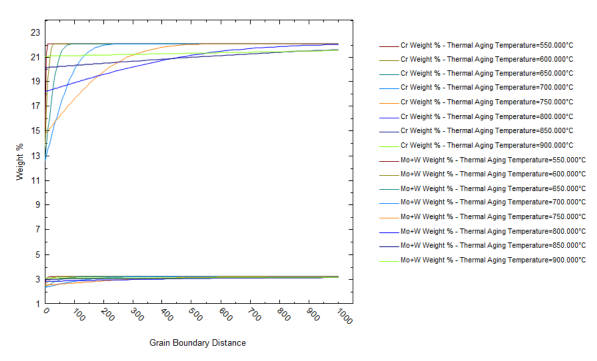


For clarity, let's customize the plot by removing the corrected depletion profiles and leaving only the uncorrected (or directly calculated) ones.

Click on the **Options** button, and select the **Curves** category, highlight the Corrected curves and check the hidden box. Then click **OK**.



Your plot should look like the image below



The plot was modified by right clicking on it and selecting the option *Allow Layout Changes*. This option allows to move and rearrange the plot and legend. Additionally, the color of the lines was changed for each temperature in order to distinguish the temperature effect.

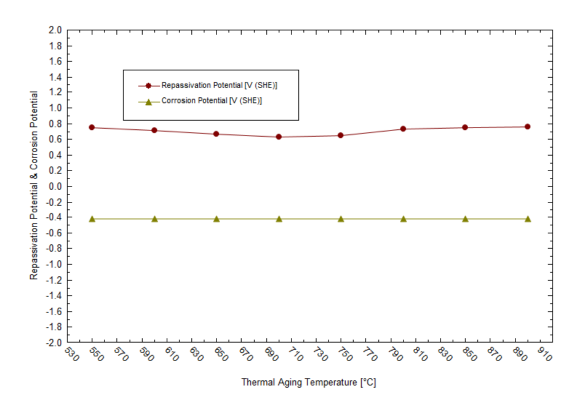
As shown in the plot above, the depletion profile is very narrow at the lowest temperature, i.e., 550 °C. On the other hand, it becomes flat at high temperatures, and it has a high minimum. The high minimum value is particularly important because it indicates that the local depletion of Cr and Mo is much less severe at high temperatures (due to much faster diffusion of substitutional elements and subsequent healing).

It should be noted that the Mo profile qualitatively parallels the Cr profile but has somewhat different slopes because of differences in diffusion coefficients of Cr and Mo.

Effect of Cr and Mo depletion on corrosion

To look at the effect of Cr and Mo depletion on corrosion

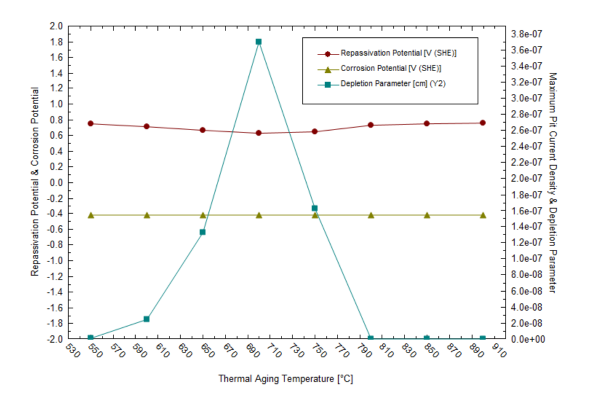
Click on the **Localized Corr.** tab (Localized Corr.). By default, this tab will show a plot of the corrosion potential and repassivation potential.



Let's add a new parameter to the plot: The **Depletion Parameter**.

Click on the Variables button. This will open a new window.

Look for the **Corrosion Values** section, click on the '+' icon, and select the **Depletion Parameter** option. Click on the >> button that belongs the **Y2 Axis**. This action will place the new variable under the Y2 Axis. Then click **OK.**



It is evident that both the depletion parameter and repassivation potential show their extreme values at intermediate temperatures of thermal aging. In the case of the depletion parameter, it is a maximum and, in the case of the repassivation potential, it is a minimum. This is in agreement with experimental data (Anderko et al., 2009). Non-zero values of the depletion parameter indicate the possibility of intergranular corrosion. A depression in the repassivation potential indicates an increased tendency for localized corrosion.

Thermal Aging and Localized Corrosion of Annealed and Thermally Aged Duplex Alloy 2324

In the previous examples, we used the Corrosion Analyzer's capabilities to predict the Cr and Mo depletion profiles for austenitic stainless steels and nickel-base alloys. However, we are not limited to such calculations.

We can also use the Corrosion Analyzer to predict the localized corrosion behavior of other alloys and other phases, including those that are not stored in the database. This facility is based on a generalized correlation for predicting the repassivation potential of Fe-Ni-Cr-Mo-W-N alloys as a function of alloy composition (Anderko et al., 2008). This correlation can be applied to both bulk alloys that are not in the database and to phases that may result from thermal aging.

In this section, we will go through two simulations:

First, we will predict the tendency for localized corrosion for the duplex alloy 2324 in an aerated chloride solution. Alloy 2324 (a.k.a. AISI 329) is not in the database, so the repassivation potential will be calculated from the generalized correlation (Anderko et al., 2008). The corrosion potential will be calculated for a similar alloy because the corrosion potential does not differ much for many Fe-Cr-Ni-Mo alloys in the passive state in neutral solutions.

Second, we will predict the localized corrosion tendency for alloy 2324 after thermal aging. Thermal aging of duplex steels in the temperature range of 900°C to 600°C leads to the formation of various phases - χ , σ , $M_{23}C_6$ -type carbide, and secondary austenite (γ_2). The secondary austenite phase is primarily responsible for the increased tendency of the alloy for localized corrosion. This is due to a very significant depletion of chromium in the secondary austenite over relatively wide spatial areas (Sridhar et al., 2009). Since the composition of the secondary austenite cannot be predicted at present, we will use experimental microstructural data (Sridhar et al., 2009) in conjunction with the generalized correlation for the repassivation potential.

Example 54: Prediction of Localized Corrosion for Alloy 2324 Before Thermal Aging in an Aerated NaCl Solution

Starting the Simulation

Use the inputs and parameters from the table below to create the stream's composition. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Thermal Aging Calculation | | | | |
|---------------------------|--------------------------------|-----------------------------------|-------------------------|--|
| Calculation Settings | | Stream Composition and Conditions | | |
| Stream Name | ame Thermal Aging – Alloy 2324 | | Calculated | |
| Calculation Type | Corrosion Rates | Temperature | 60 °C | |
| Survey by | Composition | Pressure | 1 atm | |
| Name Style | Display Name | H2O | Default – 55.5082 moles | |
| Unit Set | Metric, Batch, Moles | NaCl | 0 moles | |
| Framework | AQ | 02 | 0.02 moles | |
| | | N2 | 0.08 moles | |

Note: The oxygen and nitrogen have been added to simulate the presence of air

Add a new Stream

Click on the new Stream and press <F2> to change the name to Thermal Aging - Alloy 2324

Select the AQ thermodynamic Framework (selected by default)

Click on the Units Manager Icon and select Metric, Batch, Moles

Click on the Names Manager Icon and select Display Name

Enter the stream composition and conditions specified in the table above

Go to the Add Calculation button and select Corrosion Rates

Select the Survey by Composition

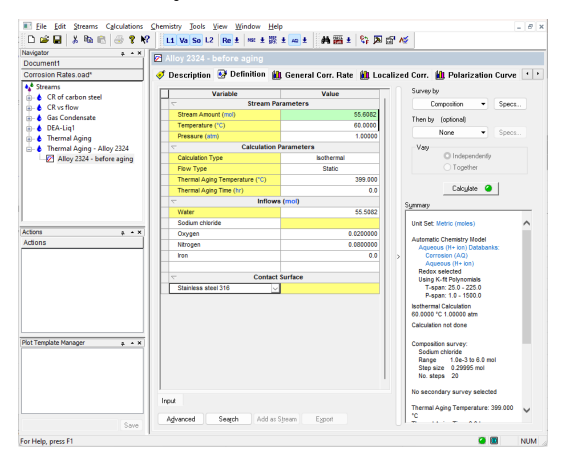
Change the name from *Rates* to *Alloy 2324 – before aging* using the <F2> key or by right-mouse clicking on the object and selecting Rename

Since alloy 2324 is not available in the database, we will select stainless steel 316 as the contact surface. This will ensure that the predicted corrosion potential is very similar to that for alloy 2324.

Select Stainless Steel 316 from the Contact Surface grid

Unlike in the previous examples, do not make any changes in the **Calc Parameters** section. We will not make Cr depletion calculations; rather, we will be specifying the **compositions of the phases**.

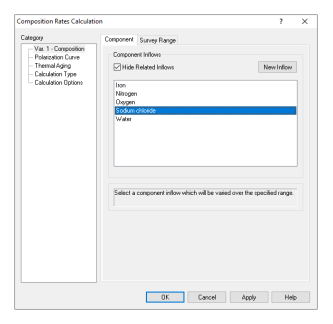
Your screen should look like the image below.



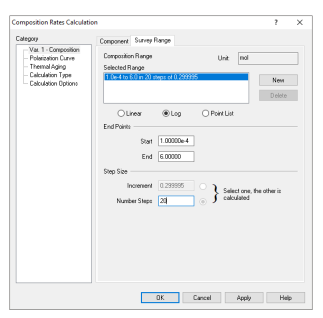
We will be running a composition survey to see how the concentration of NaCl affects the propensity for localized corrosion.

We need to specify the NaCl concentration range. Click on the Specs button. This will open a new window.

Under the Component tab select Sodium Chloride



Click on the **Survey Range** tab and change the concentration range as follows: Start=0.0001 moles, End=6 moles, and number of steps=20. Then, click on the Log radio button.



Under the **Category** section, select the **Thermal Aging** option. The screen will be populated with default parameters for type 316 stainless steel.

In the following steps we are going to enter the alloy 2324 composition and use the repassivation potential (E_{rp}) correlation (embedded in the software) to calculate the E_{rp} of the alloy. This is necessary because no parameters for alloy 2324 are stored in the databank and we have to rely exclusively on the correlation to predict the repassivation potential.

First, click on the Yes button next to "Erp Directly from Correlation:"

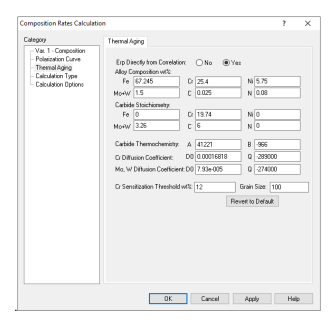
Then, enter the composition of alloy 2324 by replacing the default values for alloy 316.

Enter the following composition of alloy 2324:

| Element | Composition (wt%) |
|---------|-------------------|
| Fe | 67.245* |
| Cr | 25.4 |
| Ni | 5.75 |
| Mo+W | 1.5 |
| С | 0.025 |
| N | 0.08 |

*which is the balance that includes many minor elements

The remaining parameters in the Thermal Aging screen can remain the same because we will not be using them in this example (i.e., we will not be calculating any depletion profiles). After entering the values, the screen should look as follows:



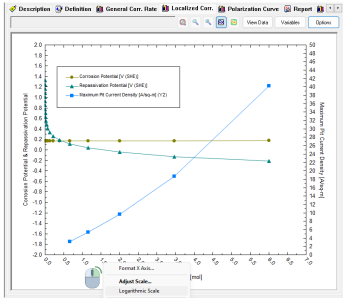
Click **OK** to accept the changes

We are now ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar. You can save it under the same file created in the previous section named *Corrosion Rates*.

Analyzing the results

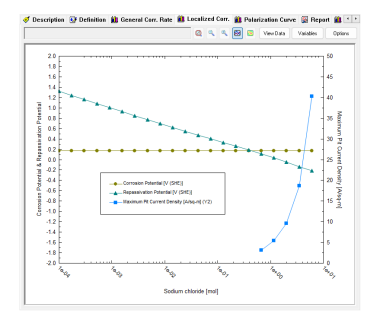
Click on the **Localized Corr.** tab (Localized Corr.). By default, this tab will show a plot of the corrosion and repassivation potentials as a function of NaCl concentration. In the Y2 axis the Maximum Pit Current Density is also plotted.



To visualize the results better, change the horizontal axis to a logarithmic scale.

Right-mouse click on the X Axis and select Logarithmic Scale

The plot should look like the image below.



These results indicate that alloy 2324 is susceptible to localized corrosion in aerated solutions when the chloride concentration exceeds \sim 0.3 molal. Above this concentration, the repassivation potential drops below the corrosion potential and, therefore, localized corrosion can be stabilized at these conditions.

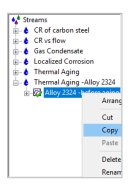
Example 55: Prediction of Localized Corrosion for Alloy 2324 After Thermal Aging

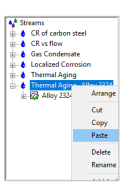
Now, we will perform the same prediction of localized corrosion calculations for thermally aged alloy 2324. We will run the same concentration survey as in the previous case so that we can compare the results for bulk alloy 2324 with those for a heat-treated sample.

Sridhar et al. (2009) found that after aging alloy 2205 at 700 °C for 24 hours, the formation of a chromium-depleted secondary austenite phase is responsible for a much-increased tendency for localized corrosion. We will use this experimental data to approximate the localized corrosion tendency of alloy 2324.

Starting the Simulation

Copy the Alloy 2324-before aging calculation and paste it under the Thermal Aging - Alloy 2324 stream





Rename the calculation as *Alloy 2324 – after aging* by using the **<F2>** key or by **right-mouse clicking** on the object and selecting **Rename**

Set Thermal Aging Temperature to 700°C. Set Thermal Aging Time to 24 hr.

Click on the **Specs** button. This will open a new window.

Under the **Category** section, select the **Thermal Aging** option. The screen will be populated with default parameters for type 316 stainless steel.

Click on the **Yes** button next to "Erp Directly from Correlation:" – because we will be running calculations for a completely new phase

Then, enter the composition of the secondary austenite phase by replacing the default values for alloy 316.

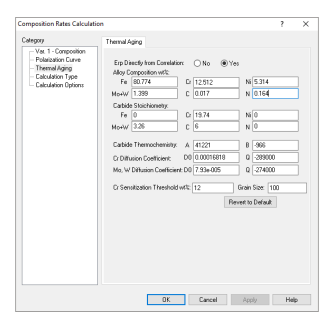
Note: we are using experimental data obtained for alloy 2205 for illustration purposes. For more details on how these compositions were determined, see the paper of Sridhar et al. (2009).

Enter the following composition:

| Element | Composition (wt%) |
|---------|-------------------|
| Fe | 80.774* |
| Cr | 12.512 |
| Ni | 5.134 |
| Mo+W | 1.399 |
| С | 0.017 |
| N | 0.164 |

^{*}which is the balance that includes many minor elements

The remaining parameters in the Thermal Aging screen can remain the same because we will not be using them in this example (i.e., we will not be calculating any depletion profiles). After entering the values, the screen should look like the image below.



Click **OK** to accept the changes

We are now ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key

It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar. You can save it under the same file created in the previous section named *Corrosion Rates*.

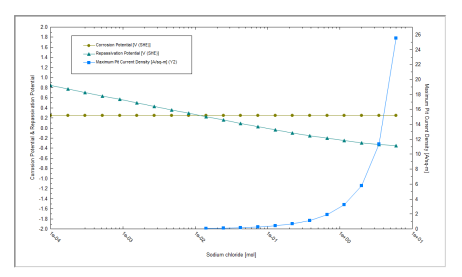
Analyzing the results

Click on the **Localized Corr.** tab (Localized Corr.). By default, this tab will show a plot of the corrosion and repassivation potentials as a function of NaCl concentration. In the Y2 axis the Maximum Pit Current Density is also plotted.

To visualize the results better, change the horizontal axis to a logarithmic scale.

Right-mouse click on the X Axis and select Logarithmic Scale

The plot should look like the image below.



The obtained plot shows that the repassivation potential is much lower than that for bulk alloy 2324. Because of the strong depression of the repassivation potential, the corrosion potential exceeds the repassivation potential at chloride concentrations of ~ 0.0007 m. Thus, localized corrosion is predicted to be possible at concentrations above 0.0007 m. This indicates a very strong increase in the propensity for localized corrosion compared with bulk alloy 2324, for which the predicted threshold is ~ 0.3 m.

In general, you can use this facility to predict the repassivation potential for any alloys, including unknown and experimental ones, as long as they belong to the Fe-Ni-Cr-Mo-W-N-C family.

Chapter IV – OLI Studio: EVS Analyzer

Section 10. EVS Overview

Overview and Theoretical Foundation

Engineering systems may have many pits and corroded areas of varying degree of severity. The first perforation, whose time and location will be a matter of probability, may cause the failure of the construction. Accordingly, the probability of such failure must be known as accurately as possible.

Extreme value statistics (EVS) is one of the most powerful statistical techniques that have been used extensively to extrapolate damage (maximum pit depth) from small samples in the laboratory to larger area samples in the field (see, for example, Eldridge G. 1957, Shibata T. et al. 1988, Kowaka et al. 1994). Thus, it was shown (Shibata T. et al. 1988) that probability of failure of a construction, P_f, i.e. the probability that at least one pit reaches the critical dimension, d, (for example wall thickness) in the system with area S, is described by the equation:

$$P_f = 1 - \exp\left\{-\frac{\exp\left[-d - \left(u + \alpha \ln\frac{S}{S}\right)\right]}{\alpha}\right\} \tag{1}$$

where location parameter, u, and scale parameter, α , are measured by using small samples with constant area, s. Equation (1) is used to extrapolate corrosion damage from a small reference area, such as a coupon to a larger operation area, S. This is the classical use of Extreme Value Statistics.

Experimental studies demonstrate that both the shape and location parameters are time-dependent. However, those dependencies must be established empirically, and since no theory contained within classical EVS is available for the functional forms of u(t) and $\alpha(t)$, it is necessary to know the answer (prediction) in advance for predicting the damage at long times. This has proven to be a severe constraint of the applicability of classical EVS.

This problem can be overcome by applying a damage function analysis (DFA) method that considers propagation of corrosion damage by drawing an analogy between the growth of a pit and the movement of a particle (Engelhardt and Macdonald, 2004). In many cases, DFA yields an analytical expression for u and α in terms of time of the hyperbolic form:

$$u = \frac{a_1 t}{1 + a_2 t}$$
 and $\alpha = a_3 t$ (2)

where a₁, a₂, and a₃ are readily determined by calibration from short term data in order to predict damage over the longer time. Namely, equations (2) are used now by OLI software for predicting damage in corroding systems. It must be noted that a different (power) form of such dependencies has been used by Laycock et al., 1990.

Input and Output Specific to EVS

For applying this technique, the user has to provide a set of experimental data (x_i, t_i, s_i) , i = 1, 2, ..., N, where x_i is the depth of the deepest pit over area s_i , of a metal exposed to corrosion attack. The separate area, s_i , could be distinct coupons from a designed experiment or random samples at various times from different locations in the system. Experiments must be performed for at least two different times.

The output of the code yields the probability of failure as a function of time for a large system with area S. The code also allows the user to answer several engineering questions: for example, what service life, t, will the pipe have with the width, d, and length L in order to ensure acceptable performance (probability of failure, P_f).

Advantages and Disadvantages of EVS

The advantage of this approach is self-evident. The prediction of corrosion damage for long times will be done by using experimental data for short times without requiring the explicit determination of any information about the kinetic parameters of the system. However, such approach has evident disadvantages, as follows:

The results of the analysis cannot be transferred for predicting corrosion damage to other systems (for example, pipelines) due to the different technological and environmental conditions that generally exist. The results cannot be used for predicting damage in the same system if technological and environmental conditions change.

We can expect that when the depth of the pit increases some critical value, the nucleation of cracks can occur. A purely statistical method cannot predict such a transition. This method also cannot predict any catastrophic event.

This method cannot be used for design of new constriction because it relies upon calibration upon a preexisting system.

Extreme Value Statistics for Predicting Pitting Damage

We will show examples of applications of Extreme Value Statistics for predicting pitting damage. Specifically, we will show how to predict:

The depth of the deepest pit in the engineering structure or laboratory systems as a function of time and the surface area of the system

Probability of failure for a given penetration depth and the area of the system as a function of observation time

Probability of failure for a given observation time and the area of the system as a function of penetration depth

Probability of failure for a given penetration depth and observation time as a function of the area of the system

Foundations of Extreme Value Statistics can be found in the following references: Aziz, 1956; Kowaka et al., 1994; Laycock et al., 1990; Engelhardt and Macdonald, 2004.

Example 56: Corrosion of Aluminum Alloy in Tap Water

In this example, we will consider the classical data for pitting corrosion (Aziz, 1956). In this paper, we can find particularly the experimental data for the maximum pit depths developed on Alcan 2S-O coupons with area s≈129 cm² immersed in Kingston tape water at 25 °C. The experimental data is summarized in the table below.

Maximum pit depth (in μ m) developed on Alcan 2S-O coupons with immersed in Kingston tape water for different observation times. Area of all coupons is $s\approx129~\text{cm}^2$

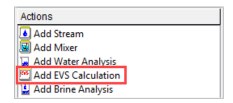
| Coupon # | One Week | One Month | Three Months | Six Months | One Year |
|----------|----------|-----------|--------------|------------|----------|
| | 7 days | 30 days | 90 days | 180 days | 365 days |
| 1 | 180 | 460 | 480 | 620 | 640 |
| 2 | 266 | 500 | 578 | 620 | 680 |
| 3 | 290 | 510 | 610 | 620 | 700 |
| 4 | 306 | 580 | 610 | 680 | 760 |
| 5 | 334 | 580 | 610 | 680 | 800 |

| 6 | 340 | 640 | 660 | 720 | 810 |
|----|-----|-----|-----|-----|-----|
| 7 | 340 | 654 | 690 | 740 | 820 |
| 8 | 410 | 680 | 718 | 740 | 840 |
| 9 | 410 | 692 | 760 | 760 | 840 |
| 10 | 545 | 692 | 798 | 760 | 900 |

Calculating the Depth of the Deepest Pit

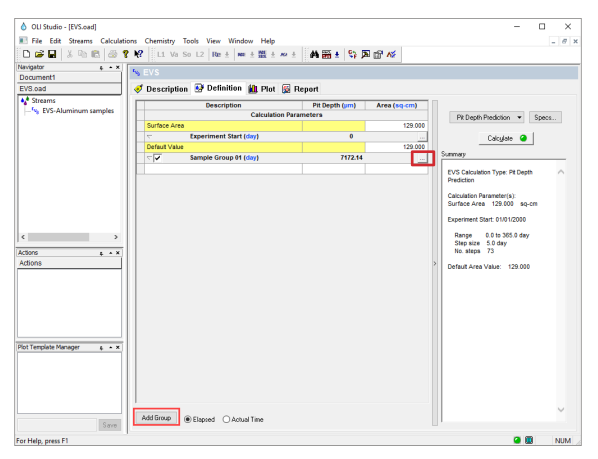
Starting the Simulation

Add a new EVS calculation. Select the Add EVS Calculation icon in the Actions Pane



Click on the new EVS calculation and press **<F2>** to change the name to *EVS-Aluminum samples*. Notice that the default calculation is the *Pit Depth Prediction*.

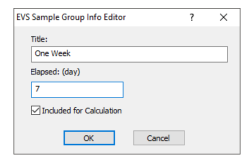
All samples have a constant area s = 129 cm². Under the **Calculation Parameters** grid, enter 129 cm² for the **Surface Area** and **Default Value** cells.



There are experimental data sets for 5 different periods of time. This means that we need to enter 5 different sample groups.

By default, **Sample group 01** is created. Click on the button to edit this field. This will open a new window.

Change the title to One Week, enter the number 7 for the number of elapsed days, and check the box 'Include for Calculation'. Then click OK.



Enter the Pit Depth data (second column) obtained for 1 week

| Description | Pit Depth (µm) | Area (sq-cm) | | |
|------------------------|----------------|--------------|--|--|
| Calculation Parameters | | | | |
| Surface Area 129. | | | | |
| Experiment Start (day) | 0 | | | |
| Default Value | | 129.000 | | |
| √ One Week (day) | 7.0 | | | |
| 1 | 180.000 | 129.000 | | |
| 2 | 266.000 | 129.000 | | |
| 3 | 290.000 | 129.000 | | |
| 4 | 306.000 | 129.000 | | |
| 5 | 334.000 | 129.000 | | |
| 6 | 340.000 | 129.000 | | |
| 7 | 340.000 | 129.000 | | |
| 8 | 410.000 | 129.000 | | |
| 9 | 410.000 | 129.000 | | |
| 10 | 545.000 | 129.000 | | |

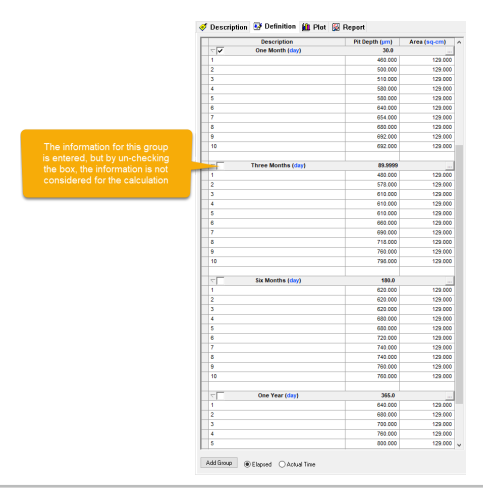
Now we need to add 4 more groups.

Click on the Add Group button (located at the bottom of the window). This will open a new window.

Change the title to One Month, enter the number 30 for the number of elapsed days, and check the box 'Include for Calculation'. Then click OK.

Enter the Pit Depth data (second column) obtained for 1 month.

Click on the **Add Group** button, and repeat the same procedure for one month, three months, six months and one year.



Note the following:

Here, for the description of different experiments, we simply used the number of the corresponding row in Table 1. However, this description can be done in an arbitrary form.

For each group, the order of samples' relative depth can be arbitrary (not necessary in ascending order as in Table 1)

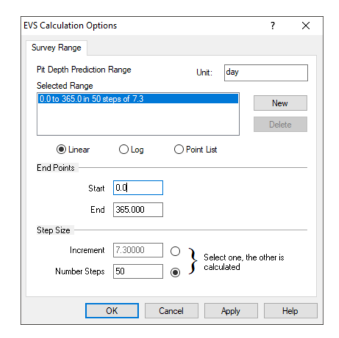
If coupons have different areas, each area must be specified in the column Area

Radio button **Elapsed** means the time after corrosion attack is used in calculations and, namely, this time is used usually in scientific publication. However, it is possible to use also **Actual Time** of the experiments.

At this point, the experimental data has been entered. Now we need to define the period of time for which we want to do the prediction of failure due to pitting.

Click on the Specs button. This will open a new window.

Under the **Survey Range** tab, enter the range for **Pit Depth Prediction** as follows: Start=0 days, End=365 days, and number of steps=50. Then click **OK**



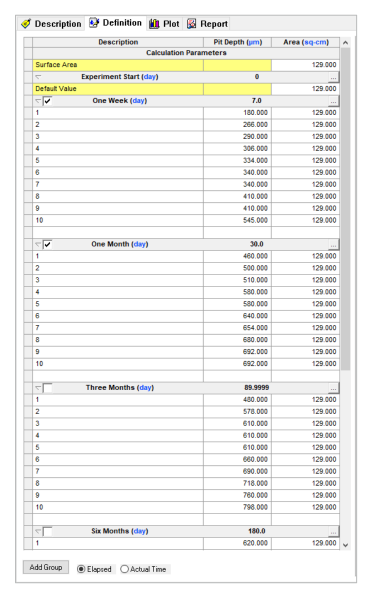
Before running the calculation, we are going to study two different cases:

The first case will include the experimental data for 1 week and 1 month – short term experiments

The second case will include all the experimental data – long term experiments

Predicting the Worst Pit Depth for 1 year of service life using short term experiments (Experimental Data for 1 week and 1 month)

Check the boxes for 1 week and 1 month only to include them into the calculations



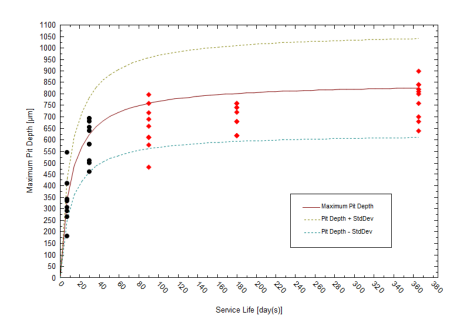
We are now ready to perform the calculation. Click on the Calculate button or press the <F9> key

It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar. Save this new file as *EVS calculations*.

Analyzing the results

Click on the **Plot** tab (\coprod **Plot**). You will see a plot of the predicted mean value of the depth of the deepest pit, x_m and the plots of values $x_m - \sigma$ and $x_m + \sigma$, where σ is the standard deviation of x_m .

It is important to note that only data for short term experiments (for 1 week and 1 month) are shown as black circles in the plot. The additional data are shown as red diamonds, only for demonstrating the accuracy of prediction.



Click on the Report tab, and to the Calculation Results table.

The predicted depth of the deepest pit at 365 days is $610.43\mu m \le 825.373\mu m \le 1040.31\mu m$ (based on the short-term experimental data).

Predicting the Worst Pit Depth for 2 years of service life using long term experiments (all data sets)

The accuracy of prediction increases when additional group of experiments are included into consideration. For this case, we are going to add the experimental data for 3 months, 6 months, and 1 year.

Go back to the **Definition** tab

Include (check) the experimental data for 3 months, 6 months, and 1 year

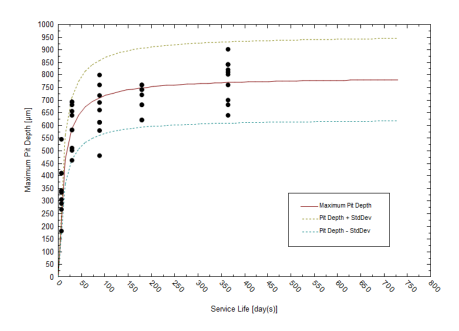
Click on the Specs button. This will open a new window

Under the **Survey Range** tab, enter the range for **Pit Depth Prediction** as follows: Start=0 days, End=730 days, and number of steps=50. Then click **OK**

Then Click on the Calculate button or press the <F9> key

Analyzing the results

Click on the **Plot** tab (Plot).



Click on the Report tab, and to the Calculation Results table.

The predicted depth of the deepest pit at 730 days (2 years) is 617.57 μ m \leq 780.873 μ m \leq 944.18 μ m.

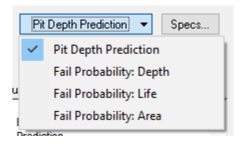
Calculating Fail Probabilities

When doing engineering design, information regarding the probability of failure is of importance to consider. The next set of calculations will involve calculating the Probability of Failure for a given:

Penetration depth

Observation time (life)

Area of the system



Fail Probability: Depth

Let's start with the calculation of Probability of Failure for a given penetration depth and the area of the system as a function of observation time.

We are going to answer the following question:

Which thickness, d, does an aluminum pipe with the area of $S=1 \text{ m}^2$ have to have in order to ensure acceptable performance (probability of failure $P_f < 5\%$ at design service life, $t_s = 5$ years)?

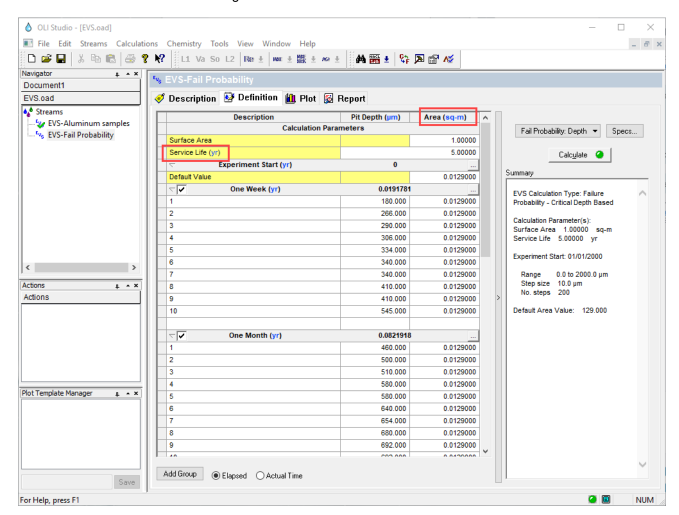
Starting the Simulation

Use the data entered in the example above. Change the Calculation Type to Fail Probability: Depth

Under the **Calculation Parameters** grid, enter 1 m² for the **Surface Area** and 5 years for the **Service Life**. Change units by clicking on the **blue** hyperlinks.

Make sure all data sets are selected (checked)

Your screen should look like the image below.

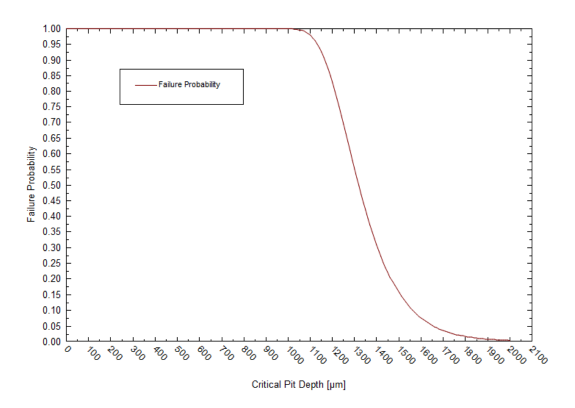


We are now ready to perform the calculation. Click on the Calculate button or press the <F9> key

It is time to save your file (File >Save as...) or use the save icon in the tool bar.

Analyzing the results

Click on the Plot tab (Plot). You will see a plot of predicted probability of failure



Click on the **Report** tab, and to the **Calculation Results** table.

We can conclude that the acceptable pipe wall width is reached at d > 1950 µm.

Fail Probability: Life

Let's continue with the calculation of probability of failure for a given service time (life) and the area of the system as a function of penetration depth.

We are going to answer the following question:

What service life, t, will the aluminum pipe have with the width,\ d=1850 μ m and area S=10 m² in order to ensure acceptable performance, i.e. probability of failure, Pr< 5 %?

Starting the Simulation

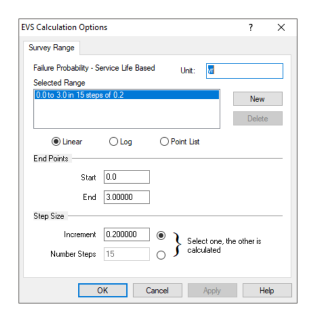
Change the Calculation Type to Fail Probability: Life

Under the **Calculation Parameters** grid, enter 10 m² for the **Surface Area** and 1850 μ m for the **Critical Pit Depth.** Change units by clicking on the **blue** hyperlinks (If needed).

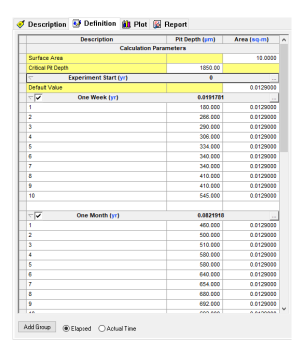
For this calculation, we need to specify the period of time at which we want to bound the calculation.

Click on the **Specs** button. This will open a new window.

Change the **Service Life** range as follows: Start=0 years, End=3 years and Increments=0.2 years. Then click **OK**.



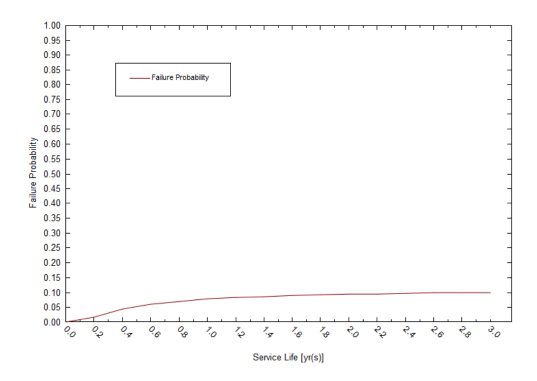
Your screen should look like the image below.



We are now ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar.

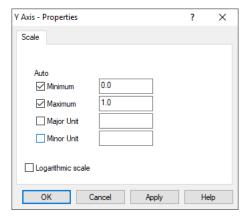
Analyzing the results

Click on the **Plot** tab (Plot). You will see a plot of predicted probability of failure. We need to modify the plot to see the results in more detail.

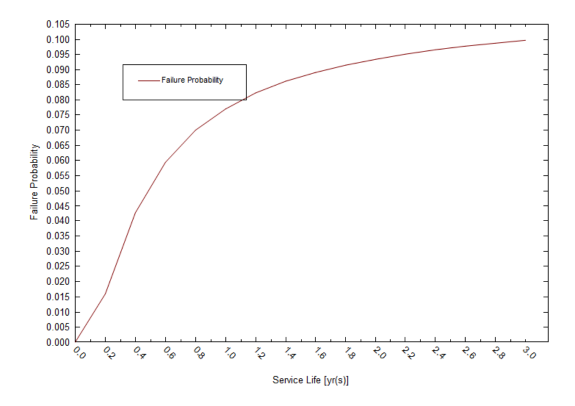


Double-click on the **Y axis**. This will open a new window.

Check the boxes Minimum and Maximum. Then click OK.



The plot should look like the image below.



The maximum probability of failure seems to be at around 10%.

Click on the **Report** tab, and to the **Calculation Results** table.

We can conclude that the lifetime of this pipe before failing is <0.6 years.

Fail Probability: Area

Finally, let's do the the calculation of Probability of failure for a given penetration depth and service time as a function of the area of the system.

We are going to answer the following question:

What area, S, can the aluminum pipe have with the width, d=2000 μ m and service life t=5 years in order to ensure acceptable performance, i.e. probability of failure, $P_f < 5 \%$?

Starting the Simulation

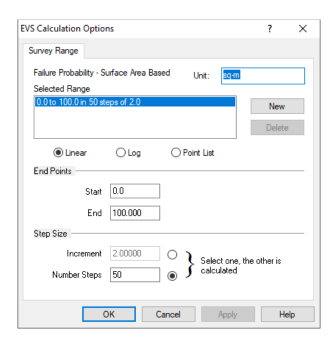
Change the Calculation Type to Fail Probability: Area

Under the **Calculation Parameters** grid, enter 2000 μm for **Critical Pit Depth** and 5 years for the **Service Life.** Change units by clicking on the **blue** hyperlinks (If needed).

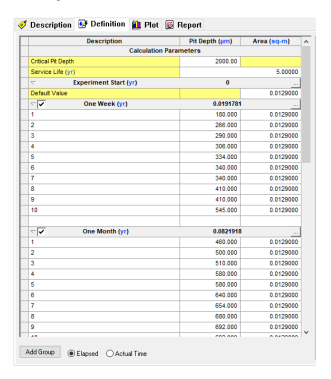
For this calculation we need to specify the surface area range that we want to survey.

Click on the **Specs** button. This will open a new window.

Change the **Surface Area** range as follows: Start=0 sq-m, End=100 sq-m and Number of steps=50. Then click **OK**.



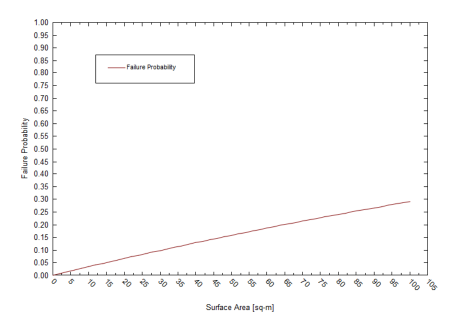
Your screen should look like the image below.



We are now ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar.

Analyzing the results

Click on the **Plot** tab (Plot). You will see a plot of predicted probability of failure. We need to modify the plot to see the results in more detail.



The maximum probability of failure seems to be at around 30%.

Click on the **Report** tab, and to the **Calculation Results** table.

We can conclude that acceptable performance is reached at <14 m².

Example 57: Failure Analysis due to Corrosion in Pipelines

In this example, we will show how in some cases reliable prediction of corrosion damage can be done by using a very limited number of experimental points.

In the table below, you can see the results of direct measurements of the depth of the deepest pits in the pipeline between Samara and Moscow [Zikerman, 1972].

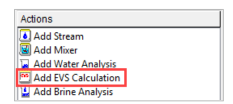
Pit depths (mm) for pipeline between Samara and Moscow

| | Group 1 | Group 2 | Group 3 | Group 4 | Group 5 | Group 6 | Group 7 | Group 8 |
|--------|---------|---------|---------|---------|---------|---------|---------|---------|
| Sample | 1440 h | 5040 h | 5760 h | 8959 h | 12624 h | 17688 h | 28032 h | 28272 h |
| 1 | 0.1 | | | 1.4 | 1.7 | 1.9 | | 2.1 |
| 2 | | 0.49 | | 1.95 | 2.1 | 2.08 | | 2.25 |
| 3 | 0.3 | 1.6 | | 1.8 | | | | |
| 4 | 0.4 | | | 1.6 | 1.65 | 1.88 | | |
| 5 | 0.9 | | 1.57 | 2.1 | 2.21 | 2.4 | 2.4 | |
| 6 | 0.3 | | 1.2 | 1.4 | 1.4 | 1.55 | 1.71 | |

Calculation the Depth of the Deepest Pit

Starting the Simulation

Add a new EVS calculation. Select the Add EVS Calculation icon in the Actions Pane



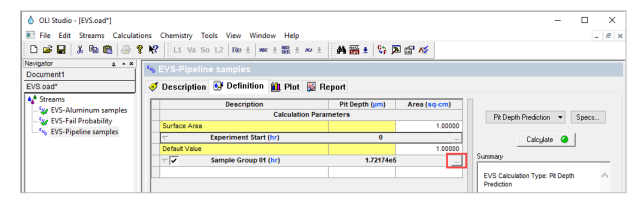
Click on the new EVS calculation and press **<F2>** to change the name to *EVS-Pipeline samples*. Notice that the default calculation is the *Pit Depth Prediction*.

Under the Calculation Parameters grid, enter 1 cm² for the Surface Area and Default Value cells.

Note: Because the area of the pipelines metal was not changed with time, the information about this area is not needed for extrapolation of corrosion damage in time.

Since the data is in hours, click on the blue hyperlink day and change it to hours (hr) for both the **Experiment** Start and the Sample Group.

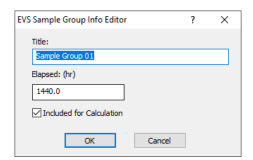
Your screen should look like the image below:



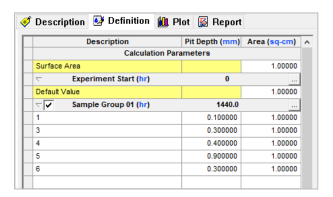
There are experimental data sets for 8 different periods of time. This means that we need to enter 8 different sample groups.

By default, **Sample group 01** is created. Click on the [...] button to edit this field. This will open a new window.

Leave the default name as Sample Group 01, enter the number 1440 for the number of elapsed hours, and check the box 'Include for Calculation'



Enter the Pit Depth data (second column) obtained for 1440 hours.



Now we need to add 7 more groups.

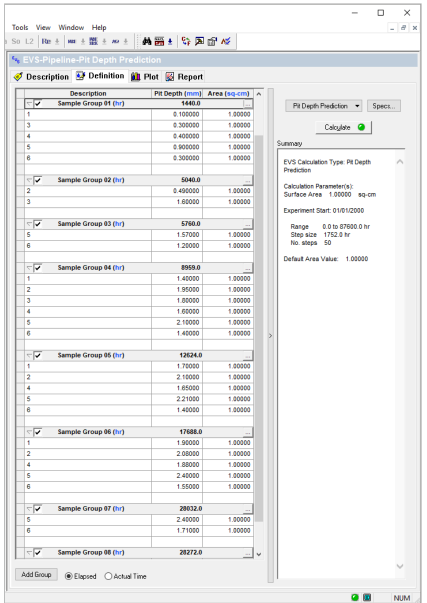
Click on the Add Group button (located at the bottom of the window. This will open a new window.

Leave the default name as **Sample Group 02**, enter the number 5040 for the number of **elapsed hours**, and **check** the box '**Include for Calculation**'. Then click **OK**.

Enter the Pit Depth data (second column) obtained for 5040 hours.

Click on the **Add Group** button, and repeat the same procedure for 5760, 8959, 12624, 17688, 28032, and 28272 hours.

Once you have entered all the experimental data, your screen should look like the image below.

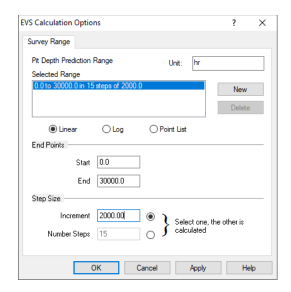


Make sure that all the groups are selected (checked) to be considered for the calculation.

At this point, the experimental data has been entered. Now we need to define the period of time for which we want to do the prediction of failure due to pitting.

Click on the Specs button. This will open a new window.

Under the **Survey Range** tab, enter the range for **Pit Depth Prediction** as follows: Start=0 hours, End=30000 hours, and Increment=2000 hours. Then click **OK**



Before running the calculation, we are going to study two different cases: The first case will include the experimental data from groups 01 to 05 The second case will include all the experimental data from groups 01 to 08

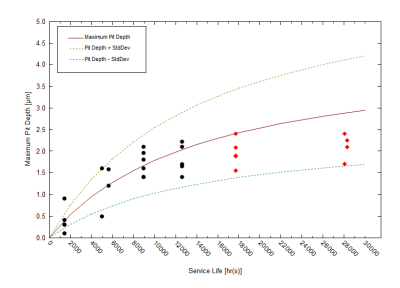
Calculation including Groups 01 to 05

Check the boxes for Group 01 to 05

We are now ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar. Save it under the same file that we previously created as *EVS calculations*.

Analyzing the results

Click on the **Plot** tab (M Plot). You will see a plot of the predicted mean value of the depth of the deepest pit, x_m , and the plots of values $x_m - \sigma$ and $x_m + \sigma$, where σ is the standard deviation of x_m .



As previously noted, only the points in black mean that were used for predicting propagation of corrosion damage. The red diamonds mean that were not taken into account for the calculation and are shown only for demonstrating the accuracy of prediction.

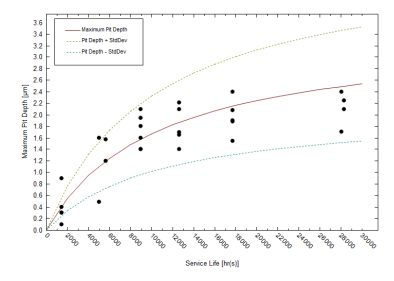
Calculation including Groups 01 to 08

Check the boxes for Group 01 to 08

We are now ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar. Save it under the same file that we previously created as *EVS calculations*.

Analyzing the results

Click on the **Plot** tab (Plot).



The plot above shows how the predicted results improved with an increased number of subsequent inspections.

Note: The predictions can be substantially improved if they were obtained on the same part of the pipe where conditions are approximately the same.

Click on the Report tab, and to the Calculation Results table.

You can see here that at 30,000 hours the maximum pit depth predicted is 1.55 µm ≤2.54 µm ≤3.53 µm.

Example 58: Predicting the Pit Depth with Insufficient Data

In some cases, the data that provided by the user may be insufficient for reliable prediction of corrosion damage. The table below shows experimental data (obtained from Laycock et al., 1990) for depths of the deepest pits that were measured on 316L coupons ($2 \times 2 \times 1/2$ in) in a 10% ferric chloride solution at 50 °C.

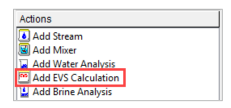
Pit Depth Data for an area of $s=25.8 \text{ cm}^2$, depth in μm

| Sample | Grp 1 | Grp 2 | Grp3 | Grp 4 | Grp 5 | Grp 6 | Grp 7 | Grp 8 | Grp 9 |
|--------|--------|----------|----------|---------|---------|---------|----------|---------|---------|
| | 40.5 h | 144.17 h | 215.33 h | 292.5 h | 331.0 h | 378.5 h | 453.25 h | 477.0 h | 528.0 h |
| 1 | 775 | 1326 | 1036 | 912 | 1361 | 1613 | 2101 | 1722 | 1714 |
| 2 | | 1176 | 1199 | 1173 | 1534 | 1641 | 2024 | 1798 | 1767 |
| 3 | | | | | | | | 1496 | 1775 |

Calculation the Depth of the Deepest Pit

Starting the Simulation

Add a new EVS calculation. Select the Add EVS Calculation icon in the Actions Pane



Click on the new EVS calculation and press **<F2>** to change the name to *EVS-Insufficient Data*. Notice that the default calculation is the *Pit Depth Prediction*.

Under the Calculation Parameters grid, enter 25.8 cm² for the Surface Area and Default Value cells.

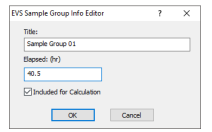
Since the data is in hours, click on the blue hyperlink **day** and change it to hours (**hr**) (for both: The **Experiment Start** and the **Sample Group**.

Your screen should look like the image below:



There are experimental data sets for 9 different periods of time. This means that we need to enter 9 different sample groups.

By default, **Sample group 01** is created. Click on the ____ button to edit this field. This will open a new window. **Leave** the default name as **Sample Group 01**, enter the number **40.5** for the number of **elapsed hours**, and **check** the box 'Include for Calculation'



Enter the **Pit Depth** data (second column) obtained for 40.5 hours.

Now we need to add 8 more groups.

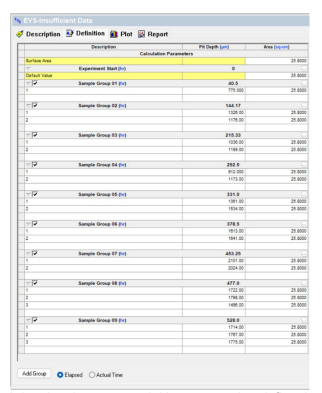
Click on the Add Group button (located at the bottom of the window. This will open a new window.

Leave the default name as **Sample Group 02**, enter the number 144.17 for the number of **elapsed hours**, and **check** the box '**Include for Calculation**'. Then click **OK**.

Enter the Pit Depth data (second column) obtained for 144.17 hours.

Click on the Add Group button, and repeat the same procedure for the rest of the experimental data (groups)

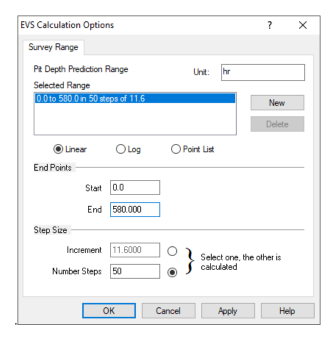
Once you have entered all the experimental data, your screen should look like the image below.



At this point, the experimental data has been entered. Now we need to define the period of time for which we want to do the prediction of failure due to pitting.

Click on the Specs button. This will open a new window.

Under the **Survey Range** tab, enter the range for **Pit Depth Prediction** as follows: Start=0 hours, End=580 hours, and number of steps=50 hours. Then click **OK**



Before running the calculation, we are going to study two different cases:

The first case will include the experimental data from groups 01 to 03 The second case will include all the experimental data from groups 01 to 09

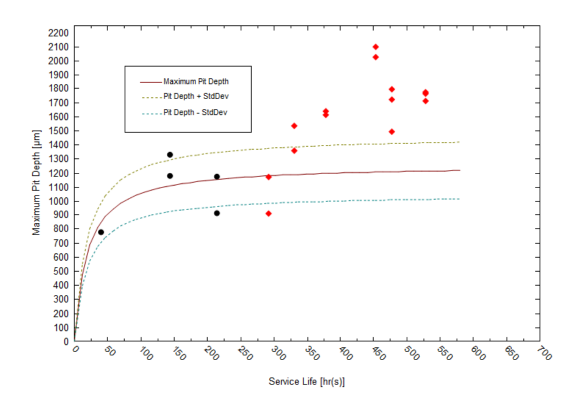
Calculation including Groups 01 to 03

Check the boxes for Group 01 to 03

We are now ready to perform the calculation. Click on the Calculate button or press the <F9> key

It is time to **save** your file **(File >Save as...)** or use the **save** icon in the tool bar. Save it under the same file that we previously created as *EVS calculations*.

Click on the Plot tab (Plot).



We see that in this case the prediction cannot be considered satisfactory. The reason is that for the first, second, and third observation times the observed mean value of deepest pit decreases. Obviously, such behavior of maximum pit depth has no physical foundation. Generally speaking, such situation is the result of an insufficient numbers of experiments (used coupons) for given observation times. Accordingly, we can expect that after increasing the number of used coupons the situation can improve.

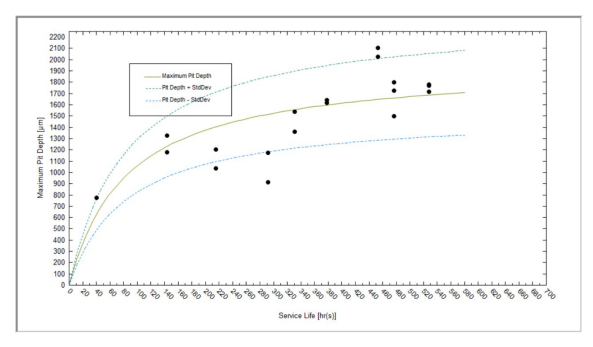
Calculation including Groups 01 to 09

Check the boxes for Group 01 to 09

We are now ready to perform the calculation. **Click** on the **Calculate** button or press the **<F9>** key It is time to **save** your file **(File >Save as...) or** use the **save** icon in the tool bar. Save it under the same file that we previously created as *EVS calculations*.

Analyzing the results

Click on the **Plot** tab (Plot).



The plot above shows how the predicted results improved with an increasing number of subsequent inspections.

The plot above also shows that the results of approximation of the full set of available experimental data from Laycock et al., 1990) can be reasonably approximated by using EVS approach.

Additionally, these results show that insufficient number of coupons (measurements at given observation times) can be compensated by increasing numbers of observation at different times.

Click on the **Report** tab, and to the **Calculation Results** table.

You can see here that at 580 hours the maximum pit depth predicted is 1331.62 μ m \leq 1706.34 μ m \leq 2081.05 μ m.

Chapter V – OLI Studio: ScaleChem

ScaleChem is a module within the OLI Studio. A separate license enables this module.

Why Use OLI Studio: ScaleChem?

Scale problems arise when fluid, initially in equilibrium with its environment, is disturbed and becomes unstable. The unstable fluid results in H₂O, CO₂, and H₂S partitioning across the water, oil, and gas phases, corrosion of metal surfaces, and precipitation/dissolution of solids. think simulation | getting the chemistry right recognizes all three effects, which are all important to the oil and gas production professional.

Precipitates form when mineral-forming elements increase the concentration of produced waters beyond supersaturation, or the saturation point. The primary causes of supersaturation are pressure, temperature, phase partitioning, and fluid mixing. think simulation | getting the chemistry right quantifies the effects on mineral scale potential while calculating the physical and chemical properties of fluid and gas phases. Industry professionals can use the software's calculations to help determine the best methods to deal with scaling situations.

think simulation | getting the chemistry right can be used to calculate scaling at one or more user specified temperatures and pressures. Other calculation options include the ability to mix waters at user specified ratios to find compatible waters, and the ability to saturate a water with respect to one or more solids to simulate reservoir conditions.

There are different types of inputs and calculation objects that can be used in think simulation | getting the chemistry right. A brief definition of each type of calculation is given below.

Input Objects

There are three different types of input objects, also referred to as analysis types, in the think simulation | getting the chemistry right software: Brine Analysis, Oil Analysis and Gas Analysis.



think simulation | getting the chemistry right refers to all waters and aqueous samples as brines. This option is added by double clicking on the **Add Brine Analysis**. A brine can be a surface water, an injection water, a formation water, a production water or any other type of aqueous fluid you can create. Brine compositions are entered in terms of ionic concentrations. In addition, the brine pH, total inorganic carbon and alkalinity can also be specified.



Add Oil Analysis allows you to enter an oil (non-aqueous phase). The oil sample may consist of pure component hydrocarbons (e.g., alkanes), distillation data, pseudocomponents or all three.



Add Gas Analysis allows you to enter any hydrocarbon mixture which may or may not contain water, carbon dioxide or hydrogen sulfide. The default hydrocarbon is methane (CH₄) but the hydrocarbon list may be expanded to include higher carbon numbers.

Calculation Objects

There are five different types of calculations that can be carried out in think simulation | getting the chemistry right: Saturator, Facilities, Scale Scenario, Scale Contour, and Mixing Water. A brief definition of each type of calculation is given below.



Add Saturator object combines fluids at the specified temperature and pressure and saturates the combined phases with the selected minerals.



Add Facilities mixes and separates fluids. It is used to simulate production operations.



Add Scale Scenario object calculates the scaling of minerals from a fluid as temperature and pressure changes, e.g. at different production locations..



Add Scale Contour object calculates the scaling of minerals from a fluid over a matrix of temperature and pressure. It creates a 2D contour plot visual.



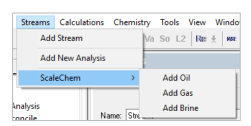
Add Mixing Water object mixes two potentially incompatible brines together to identify at what ratios the scale will form.

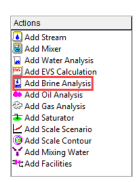
In this chapter we will model a hypothetical field and learn how to use and set up each one of the objects presented above.

Section 11. Entering a Brine Analysis in think simulation | getting the chemistry right

Entering Data for a Brine Analysis

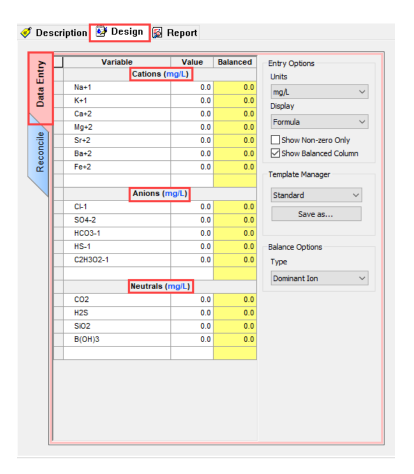
To add a **Brine Analysis**, go to the toolbar menu and click on **Streams > ScaleChem > Add Brine**, or by selecting the **Add Brine Analysis** icon in the Actions Pane.





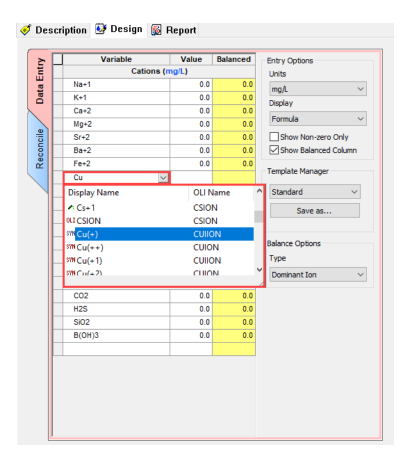
You will see three different tabs for this analysis. The **Description**, **Design** and **Report** tab.

The Brine Analysis open in the **Design**, which has 2 different sub-tabs: **Data Entry** and **Reconcile**, where we need to enter the laboratory analysis information.

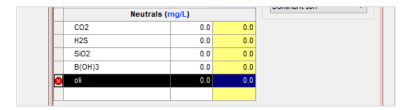


We will start in the **Data Entry** (red) sub-tab. Here you enter the concentration of Cations, Anions, and Neutrals. This sub-tab comes already prepopulated with some of the most common cations, anions, and neutrals species found in laboratory water/brine analyses (by default the units are in **mg/L**). If your species is not present in the prepopulated grid, simply click on the white grid and type the ion or neutral of interest. If it is a **cation**, type the element followed by a plus (+) sign and the corresponding oxidation state, e.g., Cu+2. If it is an **anion**, type the element followed by a minus (-) sign and the corresponding oxidation state, e.g., Br-1. If it is a **neutral**, simply type the species either using the formula name or its chemical name, as has been shown in the previous sections.

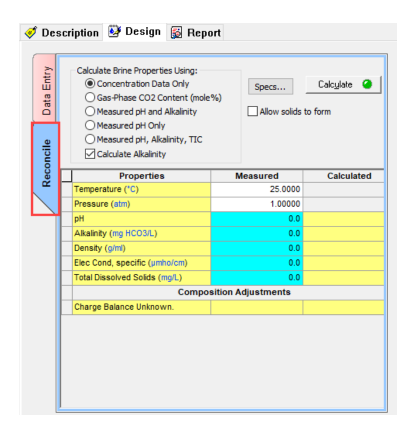
The Brine Analysis grid also contains search aids to find a specific cation, anion, or neutral species. The first search aid is the drop list located in each the cation, anion, and neutral grid sections. The list is alphabetic and is activated using the drop-down arrow within the cell, after the first few letters of the ion is typed.



If a name is misspelled or if the text is unrecognized, then a red 'X' appears to the left of the name. This name needs to be corrected or the row deleted before proceeding. To delete the row, simply select the wrong entry (which will turn black) and hit the key **<Delete>**.



The **Reconcile** sub-tab is used to define the measured properties of the brine. In this **Reconcile** sub-tab, we will tell the software how to reconcile the brine.



The first step, however, is to enter the brine measured properties and conditions. To do this we need to start in the **Properties | Measured | Calculated** table.

| Properties | Measured | Calculated | | | |
|-------------------------------|----------|------------|--|--|--|
| Temperature (°C) | 25.0000 | | | | |
| Pressure (atm) | 1.00000 | | | | |
| pH | 0.0 | | | | |
| Alkalinity (mg HCO3/L) | 0.0 | | | | |
| Density (g/ml) | 0.0 | | | | |
| Elec Cond, specific (µmho/cm) | 0.0 | | | | |
| Total Dissolved Solids (mg/L) | 0.0 | | | | |
| Composition Adjustments | | | | | |
| Charge Balance Unknown. | | | | | |

By default, the values of temperature, and pressure are 25°C and 1 atm. The aqua-blue cells indicate the values that you need to enter if they were measured at the temperature and pressure specified, these are: Measured pH, Measured Alkalinity ¹³, Density, Specific Electrical Conductivity and Total Dissolved Solids (TDS).

You can always change the units of these properties by clicking on the units highlighted in **blue**. This action will open the Units Manager Window.

If you don't have a measured property value, for example, the Specific Electrical Conductivity, just leave it blank.

¹³ This is generally a reliable value, unless solids have precipitated in the sample. Alkalinity is often but not always the same value as the bicarbonate ion (HCO₃⁻). ScaleChem Analyzer can reconcile on a measured alkalinity by adjusting the solution composition.

The yellow cells, under the **Calculated** column, will contain calculated values by the software and will be shown once the simulation is run.

When reviewing laboratory analysis of brine samples, it is quite common for the positive ions (cations) and the negatively charged ions (anions) in solution to not balance. This may be due to the precision limits of the various experimental procedures used to measure the ions - i.e. some ions may not have been analyzed. These solutions must have a neutral charge, think simulation | getting the chemistry right will adjust/modify inflows in order to balance the charges and make the solution neutral. This adjusting procedure is referred to as **Reconciliation**.

There are 5 different types of **Reconciliation Options**:

Concentration Data Only
Gas-phase CO2 content (mole%)
Measured pH and Alkalinity
Measured pH Only
Measured pH, Alkalinity, TIC

Additionally, there is the option to calculate the alkalinity value by checking the Calculate Alkalinity box.



When the Reconciliation Option is selected, the software will calculate the properties of the brine

A more detailed description of each reconciliation option is given in Section 11.3. Reconciliation Options in the Brine Analysis below.

Brine Analyses - Reporting Elements

Brine analysis data obtained from ICP measurements will contain concentrations for B, P, S, and Si. These elements do not exist in the water, rather they exist as dissolved ions. If they are part of your analysis, then you should convert them to the following before entering them into the Brine Analyses object.

Converting element concentration to species for Brine Analysis

| | Aqueous Species | Formula to enter | Formula weight multiplier |
|---------------|----------------------|---------------------------------|---------------------------------------|
| B, boron | Boric Acid | H3BO3 | B (mg/l) × 5.72 = H3BO3 (mg/l) |
| Si, Silicon | Silica | SiO2 | Si (mg/l) × 2.14 = SiO2 (mg/l) |
| P, Phosphorus | Dihydrogen Phosphate | H2PO4-1 | $P (mg/I) \times 3.13 = H2PO4 (mg/I)$ |
| | | HS-1 or SO4-2 | S (mg/l) × 1.03 for HS-1 (mg/l) |
| S, Sulfur | Sulfate or Sulfide | (cannot tell from total S only) | or |
| | | (damiet ten nem tetal e emy) | S (mg/l) × 3.0 for SO4-2 (mg/l) |

Reconciliation Options in the Brine Analysis - Definitions

When reconciling a Brine Analysis, there are five options for reconciliation:

Concentration Data Only: The software will run an electroneutrality reconciliation only, and then compute the water properties such as pH, density, etc., based on the entered concentration of neutral, cations, and anions species. In the **Concentration Data Only** option you may allow the program to pick the species to adjust for electroneutrality or you may manually choose the species to perform the adjustment. (See electroneutrality options).

Gas-phase CO2 content (mole%): Frequently it is simpler and more stable to measure the gas-phase CO₂ that is separated from the brine at the sampling point. When matched with another measured variable, usually alkalinity, the concentration of the carbonate species and the pH can be calculated. think simulation | getting the chemistry right performs a CO₂ gas fraction calculation by taking the P_{CO2} and the calculated alkalinity (based on the water analysis data) to reconcile the system for pH and carbonate properties. The CO₂ is adjusted to match a saturated gas composition.

Measured pH Only: Many brines analyses report a measured pH. This pH may or may not match the pH calculated by the software. The cause may be an incomplete and/or inaccurate brine description. The software will run both an electroneutrality and pH reconciliation. This type of reconciliation will match your recorded pH. Additionally, the software will compute the water properties such as, density, electrical conductivity, etc. The pH of the solution is automatically adjusted by the software by adding either HCl or NaOH, or you may select your preferred acids and bases to adjust the pH.

Measured pH and Alkalinity: The purpose of the Measured pH and Alkalinity reconciliations is to match the computed pH and alkalinity values with those you measured. The software will run an electroneutrality, pH and alkalinity reconciliation. Additionally, the software will compute the water properties such as density, electrical conductivity, etc. The pH of the solution is automatically adjusted by the software by adding either HCl or NaOH or you may select your preferred acids and bases to adjust the pH. The Alkalinity is automatically calculated by the software, using CO₂ as the alkalinity titrant, H₂SO₄ as the alkalinity pH titrant and 4.5 as the alkalinity end point pH. You can also change to a different alkalinity titrant if you prefer.

Measured pH, Alkalinity, TIC: The purpose of this reconciliation is to match the measured pH, total alkalinity, and the total inorganic carbon (TIC). The Total Inorganic Carbon (TIC) is adjusted using CO_2 as the alkalinity titrant, H_2SO_4 as the alkalinity pH titrant and 4.5 as the alkalinity end point pH. The software adjusts the acetate concentration (organic acids) to match the total Alkalinity value by adding or removing acetic acid. You cannot however change the CO_2 or acetic acid for the alkalinity adjustment. These are fixed by the software. The target pH is obtained simultaneously by HCl or NaOH. You may select your preferred acids and bases to adjust the pH instead of the default HCl and NaOH.

| Additionally, there is the option to Calculate Alkalinity: Calculate Alkalinity | It is important to note, that this is only |
|---|--|
| an alkalinity calculation based on the concentration entered, it is not an a | alkalinity reconciliation. |

A Basic Brine Analysis

A brief introduction to the brine analysis tool will be shown in the example below. As we go through the example, the basic definitions, functionalities and reporting for the *Brine Analysis object* will be shown.

Example 59: Basic Brine Analysis

In this example, we will input dissolved species concentrations and other measured properties into a **Brine Analysis** object to model a hypothetical water sample. We will calculate the **pH** and **scaling tendency** of a brine sample based upon its measured composition at 1 atm and 25 °C.

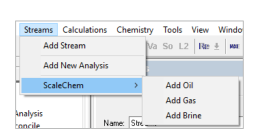
Starting the Simulation

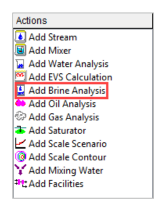
Use the inputs and parameters from the table below to create the brine analysis. Certain inputs, such as the name style, units, etc. will require further adjustments, and will be described as necessary.

| Brine Analysis Composition | | | | | | | | |
|----------------------------|----------|---------------|-------|-----------------|-----|---------------------------|-----|--|
| Cations | s (mg/L) | Anions (mg/L) | | Neutrals (mg/L) | | Measured Properties | | |
| Na+1 | 20400 | CI-1 | 37000 | CO2 | 150 | рН | 7.1 | |
| K+1 | 400 | SO4-2 | 2200 | H2S | 15 | Alkalinity as HCO3-, mg/L | 715 | |
| Ca+2 | 3800 | HCO3-1 | 0 | | | | | |
| Mg+2 | 830 | HS-1 | 0 | | | | | |
| Sr+2 | 15 | C2H3O2-1 | 715 | | | | | |
| Ba+2 | 0.7 | | | | | | | |
| Fe+2 | 10 | | | | | | | |

Setting the Water Analysis

Add a **Brine Analysis**. Go to the toolbar menu and click on **Streams > ScaleChem > Add Brine**, or by selecting the **Add Brine Analysis** icon in the Actions Pane





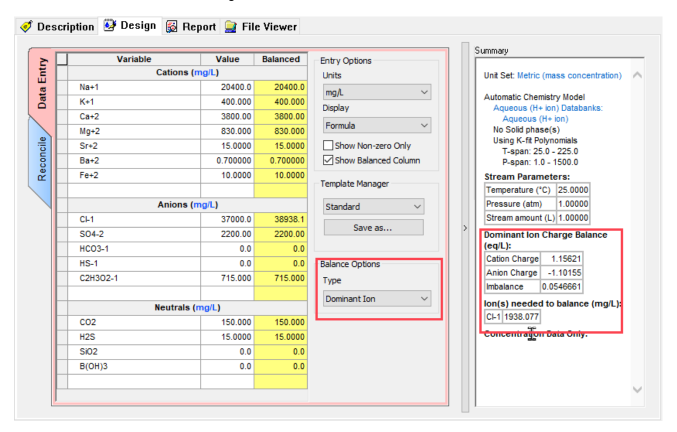
Click on the new Brine and press <F2> to change the name to Brine Analysis

Select the AQ thermodynamic Framework (remember that MSE is the default so please switch it)

Click on the Names Manager Icon and select the Formula option (default)

Click on the Units Manager Icon, and select Metric, Concentration (default)

Under the **Design** Tab, enter the **Cation, Anion,** and **Neutrals** composition given in the table above. Your screen should look like the image below:



Note: It is highly unusual for the data to be electrically neutral. Therefore, the software reconciles the sample for **electroneutrality**. After entering each species concentrations, you will notice that the software shows a yellow column named **Balanced**. The software balances charges according to the selected option and displays this in the yellow column.

There are 4 different Balance Options:

- Dominant Ion: This is the default method. The largest concentration ion is used to adjust the electroneutrality. In our example, 1938.077 mg/L of CI-1 are added since there is an excess of positive charge (see the above image).
- 2) **Prorate Cations**: All cations are adjusted up or down equally
- 3) **Prorate Anions**: All anions are adjusted up or down equally
- 4) **Make-up Ion**: This option allows for a single ion species to be adjusted. User selects an ion to increase or decrease.

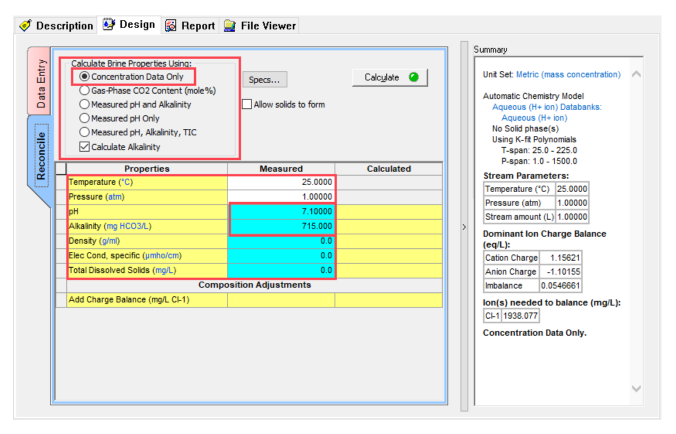
Select **Dominant Ion** as balance option

Click on the **Reconcile** (vertical blue tab)

Notice that the five different types of reconciliations are available in in the upper left corner of the window (see image below). Also, you can see the **Calculate Alkalinity box** (Calculate Alkalinity) option, which allows you to compute the alkalinity, based on the concentration entered.

Select the **Concentration Data Only** option. This option means that the software will compute the water properties based on the current concentration of neutral, cation, and anion species. The calculation will not use the measured pH, or the measured alkalinity entered (if any).

Enter the measured properties: pH = 7.1 and Alkalinity = 715 as mg HCO3/L

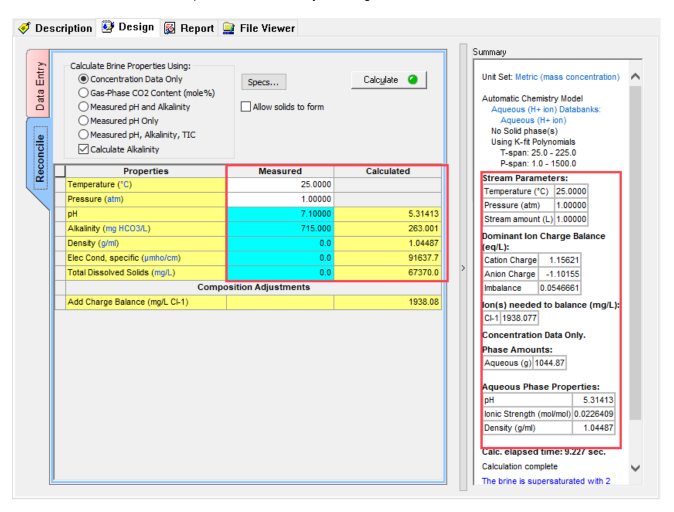


Now, we are ready to perform the calculation. Click on the Calculate button or press the <F9> key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. Create a new file and name it: *ScaleChem Analysis Calculations*.

Analyzing the Results

Once the calculation is done, the **Calculated** column displays the results obtained based on the concentrations entered in the **Data Entry** tab. Notice that the calculated results are pH=5.31 and Alkalinity=263.0 mg HCO3-1/L vs the measured values of pH=7.1 and Alkalinity=715 mg HCO3-1/L.



To the right you can see the results summarized in the **Summary Box**.

The top section contains the **Stream Parameters** information. In this example, the software performed the calculation at 25 °C, 1 atm, and a total Stream amount of 1 L.

The **Dominant Ion Charge Balance** section shows the type of balance that was used to reach electroneutrality, which cation/anion was deficient, and how much of it was added.

The **Phase amounts** section shows the distribution of species in the different phases. This analysis contains two phases: aqueous and vapor.

The **Aqueous Phase Properties** information shows the computed pH, ionic strength, and density of the solution. It is important to note here that the measured pH is 7.1 and the computed pH is 5.31. The density of this solution is 1.04487 g/ml. Remember, for this example the software used only the concentration of neutrals, cations, and anions in solution to do a reconciliation.

Click on the Report tab and scroll down to the Pre and Post Scaling Tendencies table.

Pre and Post Scaling Tendencies

| Scale Mineral | Pre-scaling | Pre-index | Post-scaling | Post-index |
|------------------------|-------------|------------|--------------|------------|
| BaSO4 (Barite) | 9.41306 | 0.973731 | 9.41306 | 0.973731 |
| CaCO3 (Calcite) | 0.0252328 | -1.59803 | 0.0252328 | -1.59803 |
| CaSO4.2H2O (Gypsum) | 0.936931 | -0.0282926 | 0.936931 | -0.0282926 |
| CaSO4 (Anhydrite) | 0.738602 | -0.131590 | 0.738602 | -0.131590 |
| FeCO3 (Siderite) | 2.71223e-3 | -2.56667 | 2.71223e-3 | -2.56667 |
| FeS (Pyrrhotite) | 16.3375 | 1.21319 | 16.3375 | 1.21319 |
| KCI (Sylvite) | 6.72353e-4 | -3.17240 | 6.72353e-4 | -3.17240 |
| FeS (Mackinawite) | 0.278931 | -0.554504 | 0.278931 | -0.554504 |
| NaCl (Halite) | 0.0122286 | -1.91262 | 0.0122286 | -1.91262 |
| SrCO3 (Strontianite) | 2.96599e-4 | -3.52783 | 2.96599e-4 | -3.52783 |
| SrSO4 (Celestine) | 0.200644 | -0.697574 | 0.200644 | -0.697574 |

From this table, you can see that Barite and Pyrrhotite solids have been predicted to be supersaturated.

For more detailed information about scaling tendencies, you can go to Scaling section on page 179.

Scroll down to the **Brine Composition** table

Brine Composition

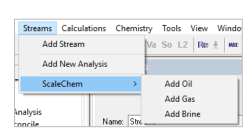
| Cations | Value (mg/L) | Anions | Value (mg/L) | Neutrals | Value (mg/L) |
|---------|--------------|-----------|--------------|----------|--------------|
| K(+1) | 400.000 | CI(-1) | 38938.1 | CO2 | 107.419 |
| Na(+1) | 20400.0 | HCO3- (*) | 59.0364 | H2S | 15.0000 |
| Ba(+2) | 0.700000 | SO4-2 | 2200.00 | | |
| Ca(+2) | 3800.00 | Acetate- | 545.141 | | |
| Fe(+2) | 10.0000 | | | | |
| Mg(+2) | 830.000 | | | | |
| Sr(+2) | 15.0000 | | | | |

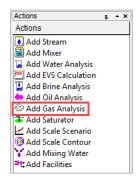
This table shows the composition of the brine after the reconciliation (calculation) has taken place.

Section 12. Entering a Gas Analysis

Entering Data for a Gas Analysis

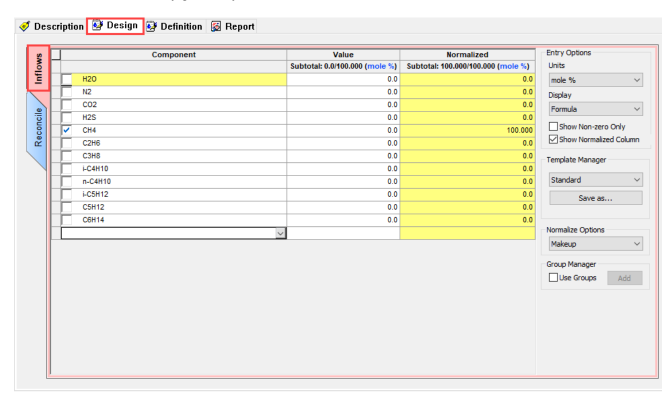
To add a **Gas Analysis**, go to the toolbar menu and click on **Streams > ScaleChem > Add Gas**, or by selecting the **Add Gas Analysis** icon in the Actions Pane.





You will see four different tabs for this analysis. The **Description, Design, Definition** and **Report** tab.

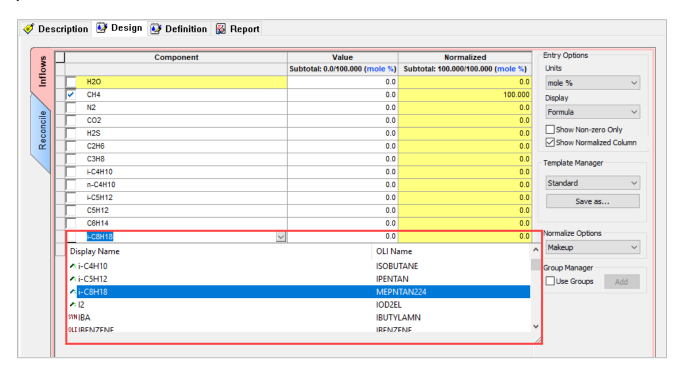
The gas analysis opens in the **Design** tab, which has 2 different sub-tabs: **Inflows** and **Reconcile**, where we need to enter the laboratory gas analysis information.



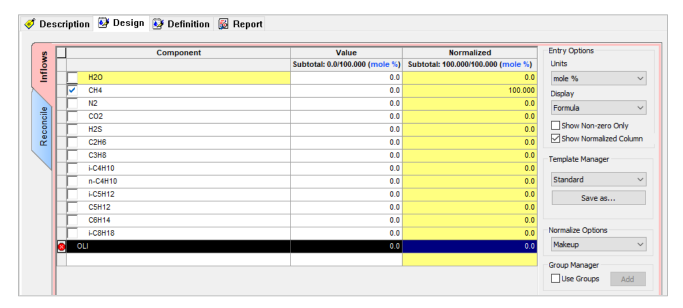
We will start in the **Inflows** (red) sub-tab. Here you enter the concentration of a pure-component hydrocarbon gas, in **mole** % units. The standard list of component extents to C6 alkanes.

If there is a component that is not present in the prepopulated grid, simply click on the white grid and type the species of interest, using the formula name or its chemical name, as has been shown in the previous sections. Let's say for example, you want to add Isooctane. Type isooctane or i-C8H18 in the white cell to add it to the list.

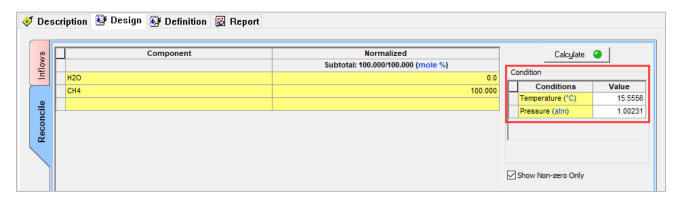
There is also the option of using the drop-down arrow, that allows you to search for the specific components you want to add.



If a name is misspelled or if the text is unrecognized, then a red 'X' appears to the left of the name. This name needs to be corrected or the row deleted before proceeding. To delete the row, simply select the wrong entry (which will turn black) and hit the key **<Delete>**.



The **Reconcile** sub-tab is used to calculate the properties of the gas at the temperature and pressure conditions specified. By default, the values of temperature, and pressure are 60 °F (15.56°C) and 14.7 psia (1.002 atm).



A Basic Gas Analysis

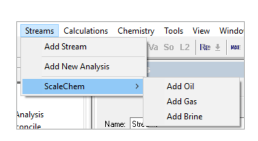
A brief introduction to the gas analysis tool will be shown in the example below. As we go through the example, the basic definitions, functionalities, and reporting for the *Gas Analysis object* will be shown.

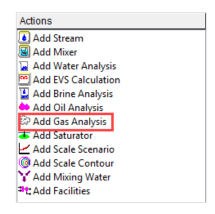
Example 60: Basic Gas Analysis

The Gas Analysis object provides a quick way to enter a pure-component hydrocarbon gas.

Setting the Gas Analysis

Add a Gas Analysis. Go to the toolbar menu and click on Streams > ScaleChem > Add Gas, or by selecting the Add Gas Analysis icon in the Actions Pane





Click on the new Gas Analysis and press <F2> to change the name to Gas Analysis

Select the AQ thermodynamic Framework (Remember that MSE is the default so you need to change it)

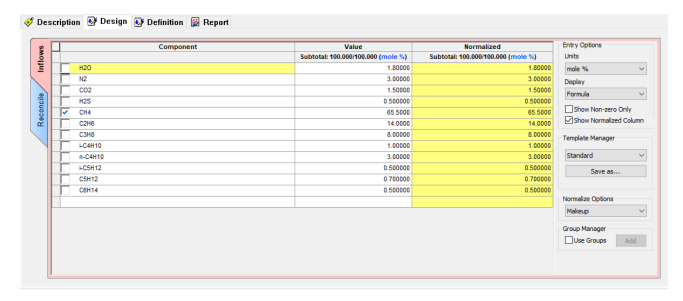
Click on the Names Manager Icon and select the Formula option (default)

Click on the Units Manager Icon, and select Metric, Mole Frac. (default)

Under the Inflows Tab, enter the gas composition given in the table below

| Gas Analysis Composition | | | | | | |
|--------------------------|------------------|--------|--|--|--|--|
| Formula | Component Name | Mole % | | | | |
| H2O | Water | 1.80 | | | | |
| N2 | Nitrogen | 3.00 | | | | |
| CO2 | Carbon Dioxide | 1.50 | | | | |
| H2S | Hydrogen Sulfide | 0.50 | | | | |
| CH4 | Methane | 65.5 | | | | |
| C2H6 | Ethane | 14.0 | | | | |
| C3H8 | Propane | 8.00 | | | | |
| i-C4H10 | Isobutane | 1.00 | | | | |
| n-C4H10 | n-Butane | 3.00 | | | | |
| i-C5H12 | Isopentane | 0.50 | | | | |
| C5H12 | n-Pentane | 0.70 | | | | |
| C6H14 | n-Hexane | 0.50 | | | | |

Your screen should look like the image below:



Note: There are several options to the right of the data entered.

Entry options: Here you can select the preferred units, and display name. You also have the option to show the non-zero values only – sometimes the composition of species is zero – and also show the normalized (yellow) column – this option is useful when the values don't add up to 100 mole%.

Template manager: Here you can select the standard (default) template or the extended template – which includes hydrocarbons with longer chains. You can also create and save your own template for future use.

Normalize options: When your analysis does not add up 100 mole% you can select between two options: Make-up and Prorate. For the Make-up option, the default gas used is CH4.

Click on **Reconcile** (vertical blue tab). The reconciliation calculation will be calculated at 15.556 C and 1.00231 atm (standard conditions).

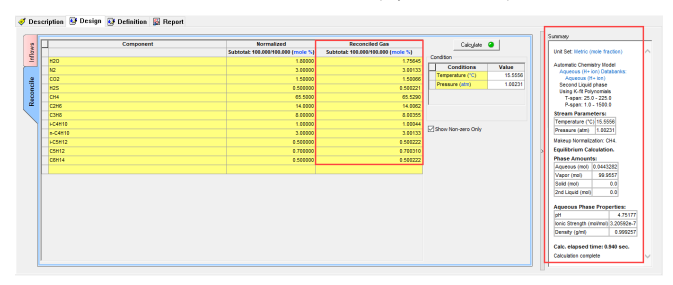


Now, we are ready to perform the calculation. Click on the Calculate button or press the <F9> key

It is time to **save** your file (**File >Save as...**) **or** using the **save** icon in the tool bar. Create a new file and name it: *ScaleChem Analysis Calculations*.

Analyzing the Results

Once the calculation is done, the **Reconciled Gas** column displays the results at equilibrium conditions.



To the right you can see the results summarized in the **Summary Box**.

The top section contains the **Stream Parameters** information. In this example, the software performed an equilibrium calculation at 15.556°C and 1.00231 atm.

The **Phase amounts** section shows the distribution of species in the different phases. This analysis contains two phases: aqueous and vapor.

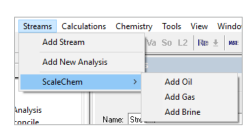
The Aqueous Phase Properties information shows the computed pH, ionic strength, and density of the solution.

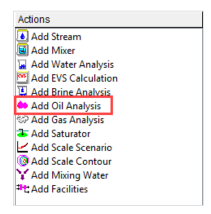
Click on the **Report** tab. Here you will see a more complete representation of the results at equilibrium.

Section 13. Entering an Oil Analysis

Entering Data for an Oil Analysis

To add an **Oil Analysis**, go to the toolbar menu and click on **Streams > ScaleChem > Add Oil**, or by selecting the **Add Oil Analysis** icon in the Actions Pane.



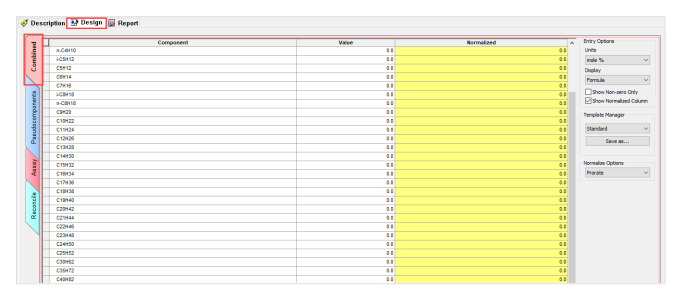


You will see three different tabs for this analysis. The **Description**, **Design**, and **Report** tab.

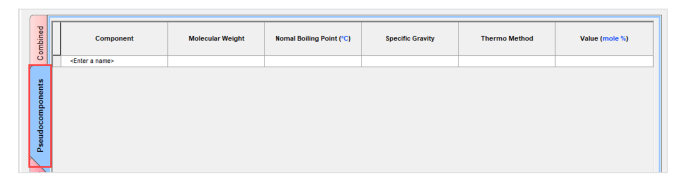
The oil analysis opens in the **Design** tab, which has 4 different sub-tabs: **Combined, Pseudocomponents, Assay** and **Reconcile.**

In the **Combined** (red) sub-tab, pure components (organic and inorganic) are entered. If there is a component that is not present in the prepopulated grid, simply click on the white grid and type the species of interest, using the formula name or its chemical name, as has been shown in the previous sections. Let's say for example, you want to add Isooctane. Type isooctane or i-C8H18 in the white cell to add it to the list.

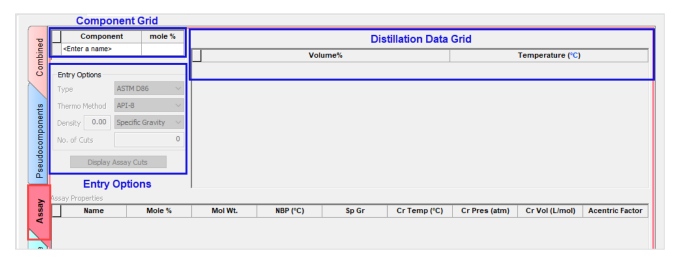
Also, you can use the **show non-zero only** option to hide all the zero values – since they are not needed.



In the **Pseudocomponents** (blue) sub-tab, you can enter pseudocomponents. Here you need to provide the molecular weight, NBP, SG, Thermodynamic Method, and the mole% information.



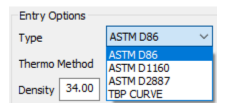
In the **Assay** (red) sub-tab, distillation curves can be entered. The Assay screen contains three data entry grids: Component, Entry Options and Distillation Data.



Component Grid: In the component grid you can name your assay. No more than 5 letters are allowed for the name. The mole% instructs the software that the mole% entered represents the total hydrocarbon mass.

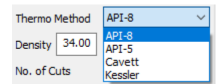
Entry Options Grid: In this grid there are four pieces of information that the user needs to fill out: The Assay Type, the Thermo Method, Density and No. of Cuts.

There are four Assay types. These are experimental methods used to create distillation curves.



ASTM D86 runs at atmospheric pressures and is used for all oil types **ASTM D1160** runs at vacuum pressure and is used for heavy oils **ASTM D2887** runs on a gas chromatograph and is used for light oils **TBP** is the true boiling point curve

There are also four methods for calculating thermodynamic properties, referred to as **Thermo Method** in the software, and these are: API-8, API-5, Cavett and Lee-Kessler.



The API-8 and API-5 are methods that use specific gravity to determine critical parameters. Cavett is a method to determine critical parameters using the API gravity Lee-Kessler is a method to determine critical parameters by using the Watson K

There are three, average bulk density options: Specific Gravity, API gravity and Watson K.



Specific Gravity (SG) is the ratio of the material density to water. think simulation | getting the chemistry right requires specific gravity to be between 0.228 and 1.6.

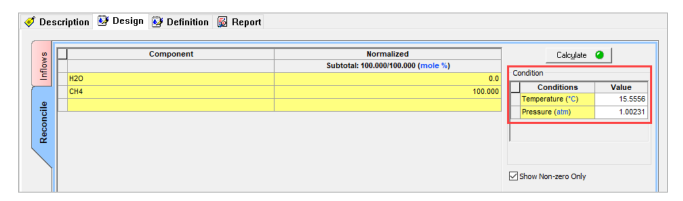
API Gravity is defined as API Gravity = (141.5/SG) – 131.5

Watson K is a method that relates density to boiling point

A more detailed description of the different assay types, thermodynamic methods and density calculation methods is found in the Basic Definitions for the Oil Analysis Tool section on pages 393-396.

Distillation Data Grid: In this section the distillation data is entered.

The **Reconcile** tab is where the equilibrium calculations are performed and is used to calculate the properties of the gas at the temperature and pressure conditions specified. By default, the values of temperature, and pressure are 60 °F (15.556°C) and 14.7 psia (1.002 atm).



Basic Definitions for the Oil Analysis Tool

Hydrocarbon Petroleum Fractions

Frequently a hydrocarbon analysis is the only data available for entry into the software. This analysis is usually a distillation curve where the volume distilled as a function of temperature of a petroleum fraction has been analyzed. This information must be turned into a vapor, organic and aqueous component for use in the simulator.

ASTM D86

Used for light and medium petroleum products and is carried out at atmospheric pressure. The results are converted internally in the OLI model generator to a TBP (True Boiling Point Curve). This curve is then fit to a spline to smooth the curve. The cuts are taken from the spline.

ASTM D1160

Used for heavier petroleum products and is often carried out under vacuum. Sometimes as low as 1 mm Hg. The results are converted internally in the OLI model generator to a TBP (True Boiling Point Curve). This curve is then fit to a spline to smooth the curve. The cuts are taken from the spline.

ASTM D2887

Uses gas chromatography to produce the distillation curve and is applicable to a wide range of petroleum products. The results are always reported on a volume percent basis. The results are converted internally in the OLI model generator to a TBP (True Boiling Point Curve). This curve is then fit to a spline to smooth the curve. The cuts are taken from the spline

True Boiling Point (TBP)

These curves, in practice, are difficult to obtain. The other methods are usually used instead.

Thermodynamic Methods (pseudo-components and petroleum fractions)

The methods used within OLI to calculate the critical temperature and pressure are the API, Cavett and Lee-Kesler methods.

API Method

Uses the specific gravity to estimate the critical parameters. The specific gravity, if not entered, can be estimated from the API gravity or the Watson K. The boiling points are taken from the assay data. API version 5 (API-5) and API version 8 (API-8) are currently supported.

The API method is selected by the user. Tc, Pc, and Vc are calculated as follows:

$$T_c = 24.2787 \times T_b^{0.58848} \times SG^{0.3596}$$

$$\ln P_c = \frac{3.12281\rho + 9}{T_b^{2.3125}SG^{2.3201}} \ if \ T_b < 1000 \ ^{\circ}\text{F}$$

Otherwise

$$\begin{split} & \ln P_c = 8.3634 - \frac{0.566}{SG} - \left(0.24244 + \frac{2.2898}{SG}\right) + \left(\frac{0.11857}{SG^2}\right) \times 0.001 \times T_b \\ & + \left(1.4685 + \frac{3.648}{SG} + \frac{0.47127}{SG^2}\right) \times \left|\rho - 7 \times T_b^2 - \left(0.42019 + \frac{1.6977}{SG^2}\right)\right| \rho - 10 \times T_b^3 \end{split}$$

$$V_c = \frac{Z_c \times T_c \times 10.73}{P_c}$$

Where.

Tc = Critical temperature in Rankine

Pc = Critical pressure in psia

Zc = Critical compressibility factor

$$Z_c = \frac{1}{3.43 + 6.70 - 9 \times \Lambda^2}$$

And

$$\Delta = \frac{8.75 + 1.978(\log T_b) \times T_b}{1.8} \text{ if } T_c < 536.67 R$$

If 536.67<Tc<593 R, the above result for Δ is multiplied by f:

$$f = \left(T_c - \frac{536.67}{T_c - T_b}\right)^{0.38}$$

If Tc>593 R,

$$\Delta = \left(\frac{(0.98907 \times SG)(\Delta - 592.4439)}{MW}\right)^{0.5}$$

Cavett

This method uses the API gravity method to determine the critical properties. The API gravity, if not entered can be estimated from the actual specific gravity or the Watson K. The boiling points for the pseudo-components are taken from the assay.

If the Cavett method is chosen by the user, Tc, Pc and Vc are calculated as follows:

$$\begin{split} T_c &= 768.0712 + 1.7133693 \times T_b - 0.0010834 \times T_b^2 - 0.008921258 \times T_b \times API + (3.8890584\rho - 7) \\ &+ T_b^3 \times 5.309492\rho - 6 \times T_b^2 \times API + 3.27116\rho - 8 \times (T_b \times API)^2 \end{split}$$

$$\begin{split} log P_c = 2.829046 + 0.0009412 \times T_b - 3.047475 \rho - 5 \times T_b^2 - 2.087611 \rho - 5 \times API \times T_b + 1.5184103 \rho - 9 \times T_b^3 \\ + 1.1047809 \rho - 8 \times API \times T_b^2 - 4.82716 \rho - 8 \times API^2 \times T_b + 1.3949619 \rho - 10 \times (API \times T_b)^2 \\ V_c = \frac{Z_c \times T_c \times 10.73}{P_c} \end{split}$$

Where,

Tc = Critical temperature in Rankine

Pc = Critical pressure in psia

Zc = Critical compressibility factor

$$Z_c = \frac{1}{3.43 + 6.7\rho - 9 \times \Lambda^2}$$

And

$$\Delta = \frac{8.75 + 1.978(\log T_b) \times T_b}{1.8} if T_c < 536.67 R$$

If 536.67<Tc<593 R, the above result for Δ is multiplied by f:

$$f = \left(T_c - \frac{536.67}{T_c - T_b}\right)^{0.38}$$

If Tc>593 R,

$$\Delta = \left(\frac{(0.98907 \times SG)(\Delta - 592.4439)}{MW}\right)^{0.5}$$

Lee-Kesler

This method uses the Watson K and the specific gravity (which can be estimated via the Watson K) to determine the critical parameters.

If the Lee-Kesler method is selected by the user, Tc, Pc, and Vc are calculated as follows:

$$T_c = 341.7 + \frac{811}{SG} + (0.4244 + 0.1174 SG) \times T_b + (0.4669 - 3.2623 SG) + \frac{100,000}{T_b}$$

$$\begin{split} \ln P_c &= 8.3634 - \frac{0.566}{SG} - \left(0.24244 + \frac{2.2898}{SG} + \frac{0.11857}{SG^2}\right) \times 0.001 \times T_b \\ &+ \left(1.4685 + \frac{3.648}{SG} + \frac{0.47127}{SG^2}\right) \times \left|\rho - 7 \times T_b^2 - \left(0.42019 + \frac{1.6977}{SG^2}\right)\right| \rho - 10 \times T_b^3 \end{split}$$

$$V_c = \frac{Z_c \times T_c \times 10.73}{P_c}$$

Where,

Tc = Critical temperature in Rankine

Pc = Critical pressure in psia

Zc = Critical compressibility factor

$$Z_c = \frac{1}{3.43 + 6.7\rho - 9 \times \Delta^2}$$

And

$$\Delta = \frac{8.75 + 1.978(\log T_b) \times T_b}{1.8} if T_c < 536.67 R$$

If 536.67<Tc<593 R, the above result for Δ is multiplied by f:

$$f = \left(T_c - \frac{536.67}{T_c - T_b}\right)^{0.38}$$

If Tc>593 R,

$$\Delta = \left(\frac{(0.98907 \times SG)(\Delta - 592.4439)}{MW}\right)^{0.5}$$

Average Bulk Density

Specific Gravity

Unitless, relative to pure water (H2O) at 15 °C which has a density of 1.0 g/mL

API Gravity

Degrees API (°API). This is calculated via the following equation:

$$API(60F) = \left(\frac{141.5}{s.g.(60F)}\right) - 131.5$$

where, SG is the specific gravity at 60 °F.

Watson K

The Watson K has no units but is calculated via:

$$K = \left(\frac{NBP^{1/3}}{SG}\right)$$

where NBP is the normal Boiling point.

Putting together a complete calculation

Now that we have defined some terms, we are now ready to begin entering the information required to run a calculation. In this calculation we will be entering the concentrations of a single brine.

Task 1 – Create a Brine Analysis

In this task we will show the main interface of a Brine Analysis, followed by an example of a brine (water analysis) and calculate its scaling tendency. Steps for the simulation will be given with bullet points.

Add Brine Analysis from Actions Panel.



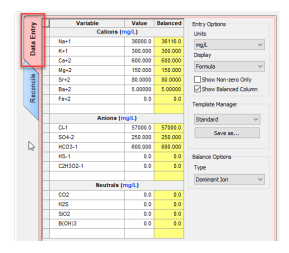
Click on the description Tab. If the **Description** tab is not currently displayed, click on the tab.



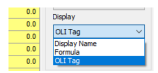
The chemistry of the brine needs to be entered. This information includes concentrations, alkalinity, pH and density. The table below describes the hypothetical water that we will create. Choose the AQ thermodynamic framework for these calculations.

| Cations | s mg/L | Anions mo | g/L | Measured Properties | |
|---------|--------|-----------|-------|-------------------------------|-------|
| Na+1 | 36000 | CI- | 57000 | Temperature | 25 C |
| K+1 | 300 | SO4-2 | 250 | Pressure | 1 atm |
| Ca2+ | 600 | HCO3-1 | 600 | pН | 7.67 |
| Mg2+ | 150 | | | Alkalinity | 600 |
| Sr+2 | 80 | | | Density (mg/L) | 1.064 |
| Ba+2 | 5 | | | Total Dissolved Solids (mg/L) | 96280 |

The screen should look like the image below

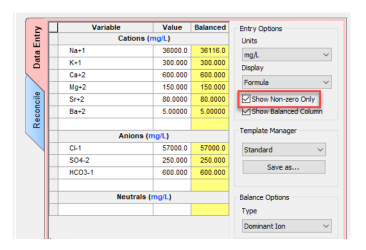


Select Formula



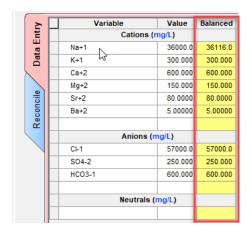
You can display an easier-to-read table by showing only the species that have actual concentrations.

Select the Show non-zero only box

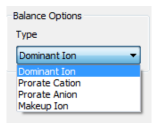


Another thing to consider is which **Balance Option** to select.

It is highly unusual for the data to be electrically neutral. Therefore, samples are reconciled for Electroneutrality. After entering each species concentrations, you will notice that Balanced values show up in the column next to values. The Column header says **Balanced**.

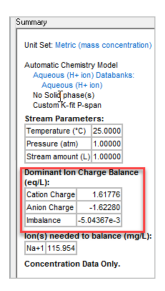


When adding or removing ions to balance charge, the solute mass is altered. We must decide as to whether we keep the mass of the solution constant (thereby adjusting the amount of water) or keeping the amount of water constant and adjusting the solution mass.



Select **Dominant Ion** balance option type.

You can review the Dominant Ion Charge and Ions needed to balance in the tables presented in the Summary Box.



The summary box shows additional detail about the brine's chemistry and balance option. The stream parameters table shows default values. The Dominant Ion Charge Balance shows the total cations or anions, and the total imbalance.

Now is time to enter the measured property data in the *Reconcile Tab*.

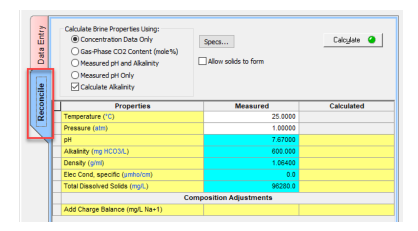
Select Concentration Data Only Option

Check the Calculate Alkalinity box

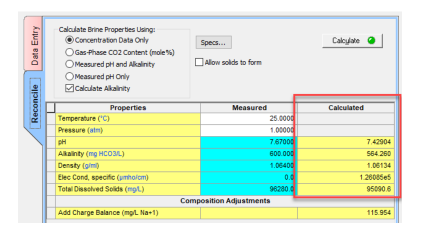
Enter the measured pH (7.67), density (1.064 g/mL) and TDS (96280 mg/L)

Leave the Allow solids to form unchecked.

We will leave the **Allow solids to form** unchecked because generally the industry uses acid to preserve water samples, which prevent solids to form. Acidified samples contain the unprecipitated ion concentration, which is what we entered into the brine. In this step, we will eliminate solids because allowing solids to form would change the brine's composition significantly.



Select the Calculate button or press the <F9> key



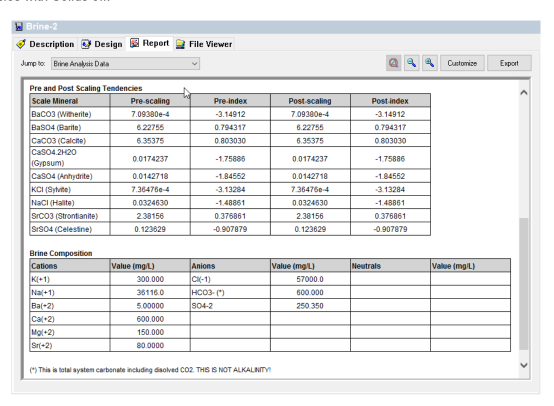
Once the calculation is done, the calculated column displays results based on the concentration that were entered in the Data Entry Tab. The calculated results of note are pH=7.43 and 564.26 mg/L as HCO3 in total alkalinity.

Experimental inaccuracies in sample measurement may, on occasion, result in a calculation that indicates that solid is supersaturated. This can normally be ignored providing that the super saturation is not excessive.



More details of the Brine composition can be found in the **Report** Tab.

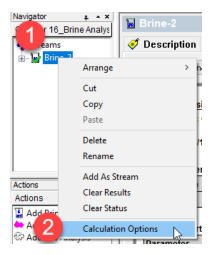
In the Report tab, scroll down to the Pre and Post Scaling Tendencies, where you will find the results of Scaling Tendencies with Solids off.



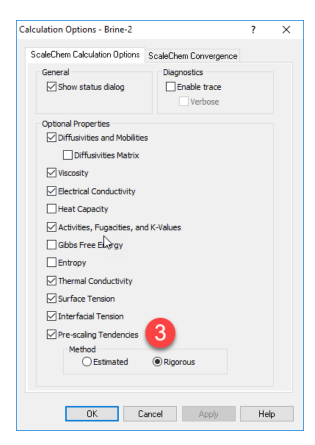
If we would have allowed solids to form, these concentrations would not be as close to the initial values.

Note: If scaling tendencies did not show up in the Report Tab, follow these steps:

Right click on the brine analysis icon Select *Calculation Options*



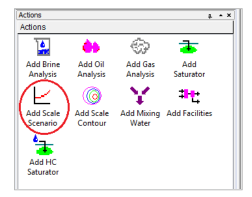
Enable Pre-scaling Tendencies



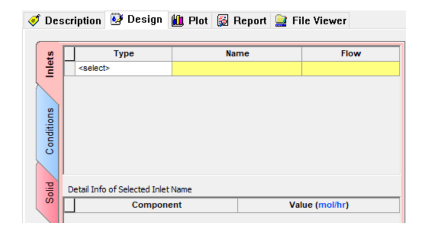
Task 2 - Create a Scaling Scenario

The next task is to calculate the brine scaling tendencies. Once the Brine Analysis data is entered and the sample reconciled, we can begin the **Scaling Scenario** calculation.

1. Select **Add Scale Scenario** from Actions panel.



Once we add a **Scale Scenario** object, the software opens to a new **Design** screen containing three vertical tabs: **Inlets, Conditions**, and **Solid**. We will work in each of these sections to set up the scaling calculation.



Inlet Tab

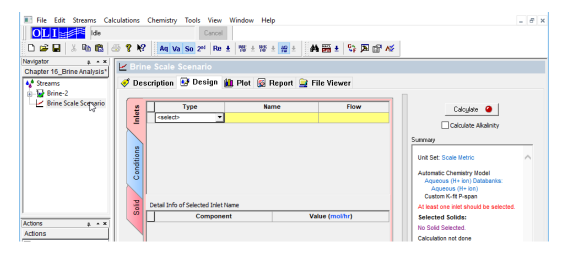
Type You can select the brine, gas or hydrocarbon of interest

Name Click in the Name field. As you position the cursor in the field, a Down Arrow will appear. You can then select from a list of brines, gases or oils already entered into this think simulation | getting the chemistry right document.

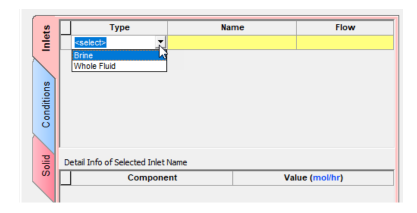
Flow Enter the flow rate for the gas, hydrocarbon or brine.

- 2. Click on the **Description** tab and rename the Scale Scenario as "Brine Scale Scenario".
- 3. Then click the **Design** tab.

Your screen should now look like image below.



- 4. Select the Inlets tab
- 5. Select the right corner of the first cell in the Type column then select Brine (bbl/day). You may need to change units by clicking the blue hyperlink.



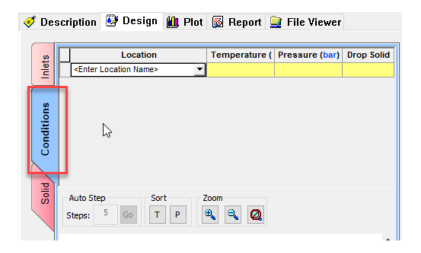
6. Select Brine from the drop-down menu in the Name column.



7. Enter 1400 (bbl/day) in the Flow Column. Please change units if necessary.

Conditions Tab

8. Select the Conditions Tab (vertical blue tab on the left-hand side of the Design screen)



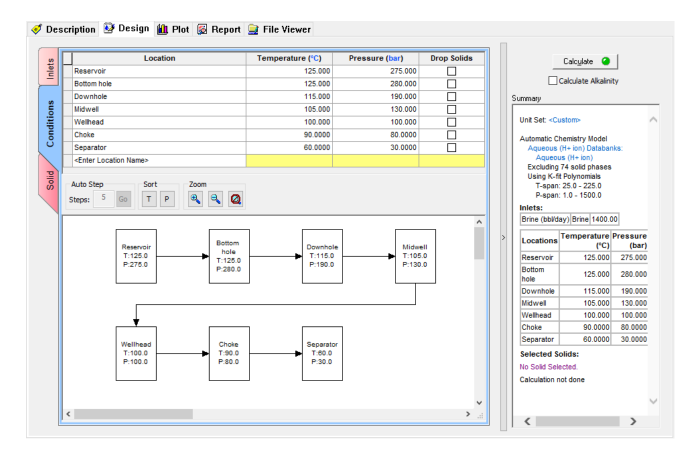
The *Conditions* section is where we enter points along the production scheme. The first column is Location. We can use the dropdown menu in each row to select a location or type in our own location name.

9. Type in the following conditions or use the dropdown menu within the Location cells.

Scale Scenario Locations

| Location | Temperature (C) | Pressue (bar) |
|------------|-----------------|------------------|
| Reservoir | 125 | 275 |
| Bottomhole | 125 | 280 |
| Downhole | 115 | 190 |
| Midwell | 105 | 130 |
| Wellhead | 100 | 100 |
| Choke | 90 | 80 |
| Separator | 60 | 30 |

The diagram within the Conditions screen and the list of locations in the Summary window both show the locations in the order that we entered. This screen has options for zooming, auto steps, sorting by temperature or pressure, and dropping solids at specified conditions.



The graphical view clearly shows the five locations and their Temperature and Pressure conditions.

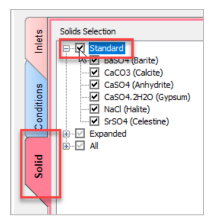
The Drop Solids checkbox column is designed to help the users decide if they want to carry forward solids from certain locations or not.

Solid Tab

- 10. Select the Solid Tab
- 11. Make sure that the solid button in the menu bar is selected.

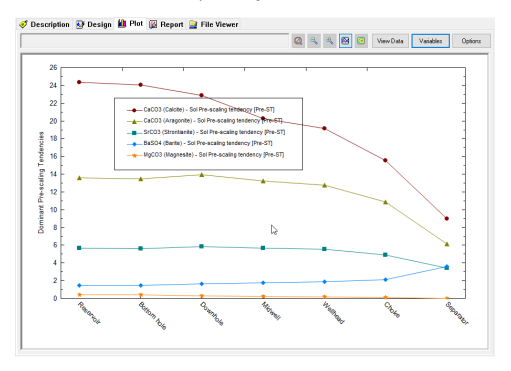


12. Select the Standard checkbox



- 13. Press the Calculate button of select the <F9> key
- 14. Select the Plot Tab

You can customize which curves are visible by selecting the *Variables* button.



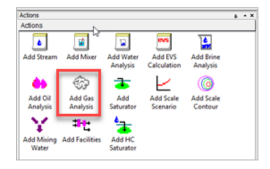
15. Select the View Data button



The *View Data* button shows the pre-scaling tendency at each location. We can copy the results of this table to a program like Excel by selecting the top left cell then selecting <Ctrl>+<C> on the keyboard.

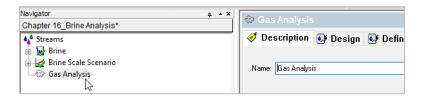
Task 3 - Create a Gas Analysis

Double click on the Add Gas Analysis object in the Actions Panel



This adds a Gas object to the Navigator panel.

Select the **Description** tab, then rename the object Gas Analysis.

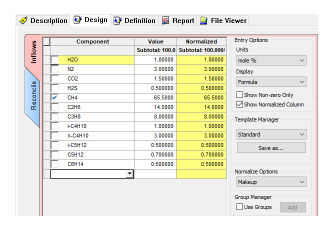


Enter the following composition and values in the Inflows grid:

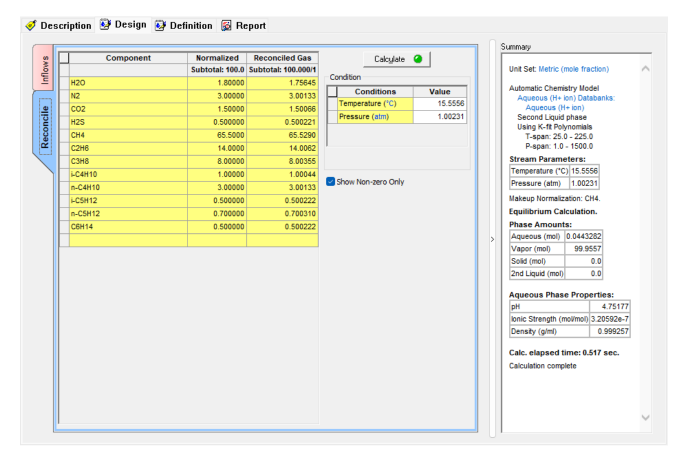
Gas Analysis

| Formula | Component Name | mole % | Formula | Component Name | mole % |
|---------|-------------------|--------|---------|-------------------|--------|
| H2O | Water | 1.80 | C3H8 | Propane | 8.00 |
| N2 | Nitrogen | 3.00 | i-C4H10 | Isobutane | 1.00 |
| CO2 | Carbon dioxide | 1.50 | n-C4H10 | n-Butane | 3.00 |
| H2S | Hydrogen sulfide | 0.50 | i-C5H12 | Isopentane | 0.50 |
| CH4 | Methane | 65.5 | n-C5H12 | n-Pentane | 0.70 |
| C2H6 | Ethane | 14.0 | n-C6H14 | n-Hexane | 0.50 |

The Inflows grid will look like one below..



This is the extent of the gas entry step. There are modifications that can be part of the gas analysis entry step (in the Reconcile Tab), but in this case, they are not considered. Now that the gas is entered, the scale scenario considered in Task-2 will be recomputed.



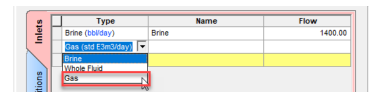
Task 4 - Recalculate Scale Scenario with Gas

The gas analysis will be added to the Scale Scenario so that it is considered during the scale evaluation.

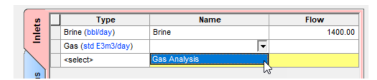
Select the Brine Scale Scenario icon in the navigator panel



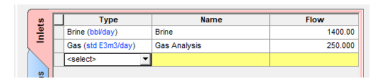
Select the **Design** Tab (horizontal tab) if not automatically sent there Select the *Inlets* tab (vertical) if not automatically sent there In the *Type* Column Add *Gas*



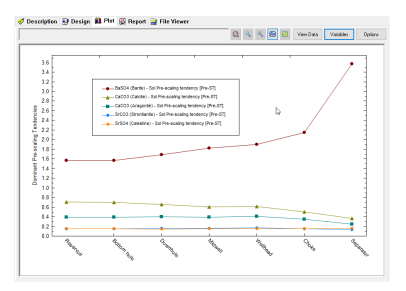
In the Name column select Gas Analysis



Enter a flowrate of 250 std E3m3/day in the flow cell



Calculate (Press <F9>) Select <u>Plot</u> Tab

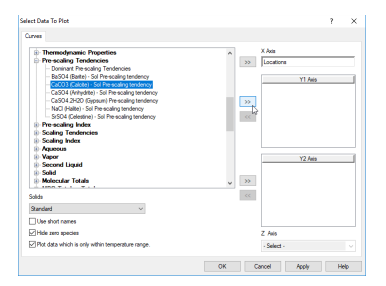


Select Variables button

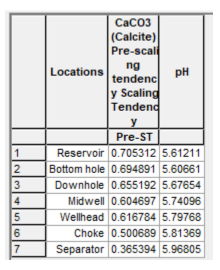
Remove all the variables from the Y1 Axis and the Y2 axis by double-clicking on the variables (this removes them from the list). Alternatively, highlight each variable and click on the double-arrow.

You will replace the existing variables with the calcite pre-scaling tendency.

Expand the Pre-scaling Tendencies button by clicking the + sign Double-click on the CaCO3 (Calcite) variable to move it to the Y1 Axis Expand the Additional Stream Parameters and add pH to the Y2 axis Click <u>OK</u>



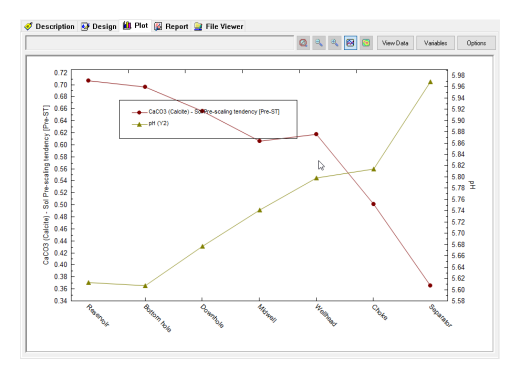
If you are in the View Data view, then the table should be similar to the one below.



Calcite is sub-saturated (S<1) at all locations.

Select the View plot button

The decreasing calcite saturation trend and the increasing pH is easier to see from the plot view.

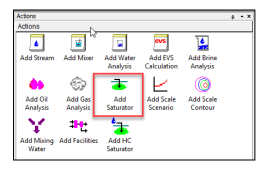


Task 5 - Creating a Saturator Object

You will use the *Saturator* object to set the reservoir minerals at equilibrium with the produced fluid. It is reasonable, in some cases, to saturate the produced fluid with common evaporite and secondary minerals like CaCO3, CaSO4.H2O, FeCO3, and NaCl. This assumes that as the reservoir fluid flows through the rock pores, there is sufficient time to interact with the surface minerals. This is based on a second assumption that the vertical column of liquid and gas are in complete equilibrium and that the saturating minerals are distributed throughout the reservoir matrix, such that complete water-mineral contact is possible.

In this case we are going to simulate a particular oil & gas well: ABC. This well produces from calcite-cemented sandstone. Barite is also present in minor quantities in the rock matrix. You will, therefore, create a reservoir called ABC Reservoir and saturate the Brine (previously calculated) with calcite and barite. You will then recalculate the Brine Scale Scenario.

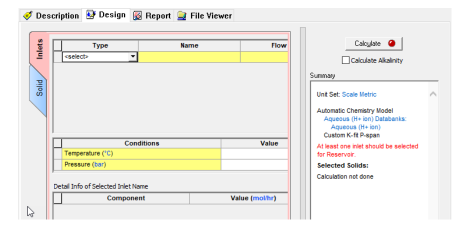
1) Add a Saturator Object



2) Select the **Description** Tab to rename the Object **Reservoir**



3) Select the Design Tab

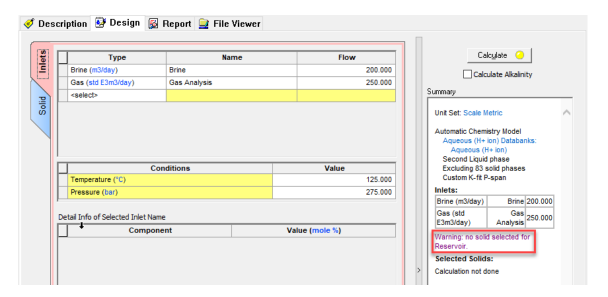


The Saturator has two vertical tabs: Inlets and Solid.

In the Inlets Tab the Fluids are entered in the top grid and the Conditions are entered in the bottom grid.

Solids are selected and saturated in the **Solid** Tab.

- 4) Enter the Brine calculated in Task 1 (see the image below for the flows)
- 5) Enter the Gas calculated in Task 3 (see the image below for the flows)
- 6) Enter the Reservoir conditions: 125 C and 275 bar in the conditions section.



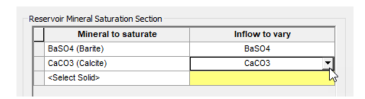
The Saturator's calculate button is yellow. This is because solids have not been selected yet. A warning also appears in the summary box. A yellow button is allowable, and a calculation may continue; it is merely a warning that the case is incomplete.

- 7) Select the **Solid** Tab (vertical tab)
- 8) Select the Standard checkbox

This last step instructs the software to allow the selected solids to precipitate – to include solid-liquid equilibrium equation in the calculation. It **does not** instruct the software to saturate these solids. Rather, if the phase is supersaturated, then it will precipitate. It is in the lower section that saturated solids are chosen.

This section contains two columns. The first (Solid) is the mineral to be saturated, and the second (inflow) is the cation and anion to be adjusted. If, for example, Barite is subsaturated in the existing fluid, then Ba+2 and SO4-2 are added. The amount of each added is stoichiometrically equivalent; one Ba+2 with one SO4-2, which maintains charge balance. If, for example, (again) Calcite is supersaturated, then equal mole amounts of Ca+2 and CO3-2 will be removed from the fluid. This inflow will be positive if the solid is subsaturated in the existing fluid, or it will be negative if the solid is supersaturated.

- 9) Use the Solid drop-down menus to select BaSO4 (Barite) and CaCO3 (Calcite) as solids to saturate
- 10) Use the Inflow drop-down menu to select BaSO4 and CaCO3 as solids to saturate (it should have been selected automatically).



At this point, the Calculate button is green indicating that the specifications are complete.

- 11) Calculate (or press the <F9> key)
- 12) Click the Report tab

The Saturator Report tab contains several tables: Saturation Details, Inlet Summary, Stream/Phase Properties, Pre and Post Scaling Tendencies, Brine Composition, Gas Composition, and if selected an inlet summary.

Scroll down to the Pre and Post Scaling Tendencies Table

Pre and Post Scaling Tendencies

| Formula | Mineral | Excess Solute | Excess Solute | Pre-Scale | Pre-Scale | Post-Scale | Post-Scale |
|------------|--------------|---------------|---------------|------------|-------------|------------|------------|
| | | mg/L | lb/1000bbl | S, ST | SI, Index | S, ST | SI, Index |
| Mg(OH)2 | Brucite | | | 4.98480e-4 | -3.30235 | 4.98479e-4 | -3.30235 |
| NaCl | Halite | 0.0 | 0.0 | 0.0245553 | -1.60985 | 0.0245553 | -1.60985 |
| BaSO4 | Barite | 2.64658e-5 | 9.27644e-6 | 1.00000 | 1.82516e-6 | 1.00000 | 0.0 |
| SrSO4 | Celestine | 0.0 | 0.0 | 0.148191 | -0.829178 | 0.148191 | -0.829178 |
| SrCO3 | Strontianite | | | 0.209852 | -0.678087 | 0.209852 | -0.678087 |
| CaCO3 | Calcite | 1.13495e-5 | 3.97806e-6 | 1.00000 | -4.82687e-8 | 1.00000 | 0.0 |
| KCI | Sylvite | | | 2.14382e-4 | -3.66881 | 2.14382e-4 | -3.66881 |
| CaSO4.2H2O | Gypsum | 0.0 | 0.0 | 0.0179366 | -1.74626 | 0.0179365 | -1.74626 |
| CaSO4 | Anhydrite | 0.0 | 0.0 | 0.0934633 | -1.02936 | 0.0934633 | -1.02936 |

Excess solute or Max Scale: The solids amount forming at equilibrium.

Pre-Scale: The saturation ratio before solids precipitate

Post-Scale: The saturation ratio AFTER solids precipitate (if solids are selected)

S, ST - Saturation, Scale Tendency: The ratio of the concentration (activity) to its solubility (S=1).

SI - Scale Index: Log(S).

The Excess Solute column shows that trace amounts of BaSO4 and CaCO3 are present in the water. These exceedingly small values (in the parts-per-trillion range) are the specifications or targets for the calculation. The software sets these target solid concentrations and then adjusts up or down, the BaSO4 and CaCO3 inflows until it reaches these targets. These part-per-trillion values may look arbitrary, but in fact the values are equivalent to 1e-10 moles of solid/kg water.

Notice also, that the pre-scale tendency for BaSO4 and CaCO3 are set to 1.0, by saturation definition. That is, these are not calculated, they are defined. The remaining scale tendencies are computed.

Task 6 – Recalculate the Scale Scenario with the Saturated Reservoir

Considerable time is spent creating what is hoped to be a more representative brine. The Brine is reconciled for charge balance, and where needed, alkalinity and pH; the gas is saturated with water as needed (though not done in this case), and lastly the fluids are set to equilibrium with important reservoir minerals. It is only after these steps are complete, and that there is confidence that the assumptions and calculations are representative of the process, that a final scaling calculation is run.

The **Reservoir** calculation contains the representative fluid. It is, therefore, the output of this calculation that will be used in all subsequent calculations. To access this output, a new Type of fluid, Whole Fluid is defined. The Whole Fluid type represents 100% of the contents of a previous calculation, regardless of phase.

You will use the Whole Fluid from the Reservoir to rerun the scaling calculations.

Select *Brine Scale Scenario* in the Navigator Pane and rename it **Reservoir Scale Scenario** Select the **Design** tab

In the *Inlets* tab in the **Type** column delete the Brine and Gas from the Type column (click the small gray box to the left of the name which will highlight the entire row, then press the delete key)

Add Whole Fluid

In the Name column select Reservoir

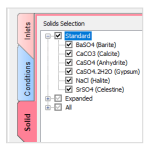
In the Flow column select < Automatic>



Note: The Whole Fluid type refers to all the phases within the fluid. The Automatic option takes the final flow rate phase of the separator and automatically enters this as the flow rate in the Scale Scenario calculation.

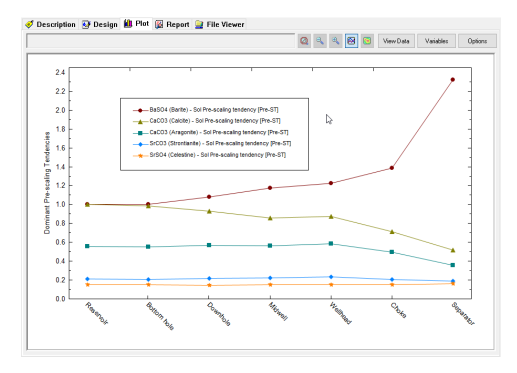
Delete the gas inlet

Go to the Solid tab and make sure that the Standard box is checked



Calculate (Press <F9>)

Select <u>Plot</u> tab and select *View Plot* (note: you may have to delete the existing entry and then reselect the dominant pre-scaling tendencies from the variables list)



The Saturator is a tool for normalizing the thermodynamics of a reservoir system. The purpose of this calculation is to apply the constraint of solid-liquid equilibrium on a system to create a specific compositional starting point.

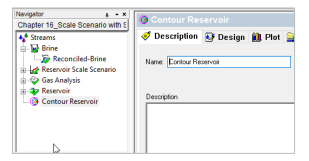
The basis of this calculation is that water in the pores of a rock matrix is at equilibrium with the mineral surface. Some questions to consider are whether it is reasonable to assume that a reservoir gas, oil, water, and rock are in equilibrium. Furthermore, we should also consider over what vertical and radial distance we can make this claim.

Regardless of the inconsistencies, the industry widely accepts this approach. It is up to the analyst to decide if the Saturator produces a more representative fluid than the original data.

Task 7 – Add a Contour Diagram

An alternative to the *Scale Scenario* calculation is the *Contour Diagram*. The conditions defined in the Scale Scenario are specific to production locations, and generally no more than ten are entered. By comparison, the conditions defined in the Contour Diagram are a range of temperatures and pressures. The object then computes a matrix of temperature and pressure conditions and creates a presentation-quality contour diagram.

Add Scale Contour
Rename the Object Contour Reservoir

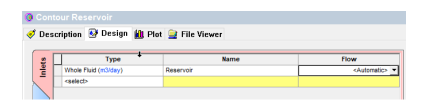


Select the **Design** Tab Select the **Inlets** (vertical) tab

Select the bottom right corner of the 1st cell in the Inlet grid and choose Whole Fluid

Select Reservoir in the 2nd column

Keep the **<**Automatic> flow rate option in the 3rd column

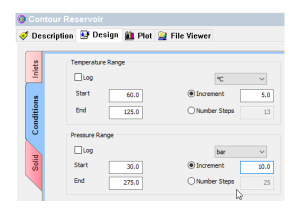


Click Conditions (vertical) tab

The minimum and maximum conditions are at the Separator 60C, 30 bar and Reservoir 125C, 275 bar. These are the start and end range of the study. To create a reasonable number of calculations (~300) and to keep the matrix somewhat symmetric, we will use increments of 2 C and 10 bar.

Change the Start to 60
Change the End to 125
Select the Increment radio button

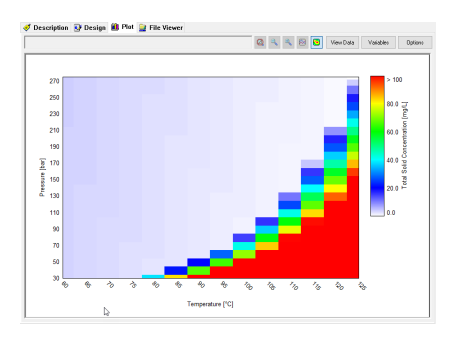
Enter 5 as the increment
Uncheck the Log box (if it is checked)
In the Pressure Range box, change the Start to 30
Change the End to 275
Select the Increment radio button (uncheck the Log box first)
Change the Increment to 10



This produces a matrix of 13 temperature and 25 pressure steps. This produces a 14x26 matrix, or 364 calculations.

| | Temperature | Pressure |
|--------------|-------------|----------|
| Units | °C | bar |
| Start | 60.0 | 30.0 |
| Stop | 125.0 | 275.0 |
| Step Size | 5.0 | 10.0 |
| No. Steps | 13.0 | 25.0 |
| Calculations | 364 | |

Select the *Solid* Tab
Check the Standard box
Calculate. The calculation should take less than 5 min to complete
When calculation is done, go to the <u>Plot</u> tab



The default plot is the total solids forming. The color scale represents a different numerical range. The red cells indicate heavier solid precipitation, and the faint purple-white in the center and left of the screen indicate that little to no scale forms. The upper right corner is the Reservoir conditions (125C and 275 bar). The lower left represents Separator conditions (60C and 30 bar). At both points, minimal solids are computed to form. The fluid traveling through the piping traverses this plot as temperature and pressure decreases. Layering production conditions (e.g., early to late life) over this plot provides an indication as to whether production will be at risk now or in the future. For example, if the pressure decline is significant (e.g., 60 bar), then the fluid will be unstable at the higher temperatures (bottom of well).

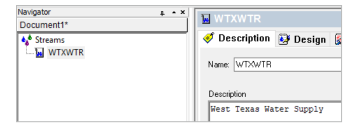
Overview of Brine, Gas and Oils and Introducing Mixer, and Saturator Objects

Other calculation options include the ability to mix waters at user specified ratios to find compatible waters, and the ability to saturate a water with respect to one or more solids to simulate reservoir conditions.

This chapter will cover the same calculation objects using fewer instructions and will introduce *Mixing*, *Saturator* and *Facility* Calculations. Its purpose is to reinforce what was learned in the previous chapter and to promote anticipation of next steps when using the interface.

Task 1 - Add a Brine

Add a *Brine Analysis* from the Action Panel Rename the Brine in the **Description** Tab, and name it WTXWTR



Go to the **Design** Tab and enter the following composition in the **Data Entry** (vertical) tab:

Name: WTXWTR
Type of water: Aquifer Water

Comment: West Texas Water Supply

| Species | Concentration | | |
|--------------------|---------------|------|--|
| Na⁺ | 3074 | mg/L | |
| Ca ⁺² | 910 | mg/L | |
| Mg ⁺² | 249 | mg/L | |
| Fe ⁺² | 0.77 | mg/L | |
| CI ⁻¹ | 4474 | mg/L | |
| SO ₄ -2 | 2960 | mg/L | |
| HCO₃⁻ | 439 | mg/L | |
| HS ⁻¹ | 146.2 | mg/L | |

Check the Show non-zero Only box (under Entry Options)

Go to the Reconcile (vertical) tab

Select the measured pH and alkalinity Option

Enter the following conditions (change units if necessary)

• To change units just click on the hyperlink next to the variable that you want to change.

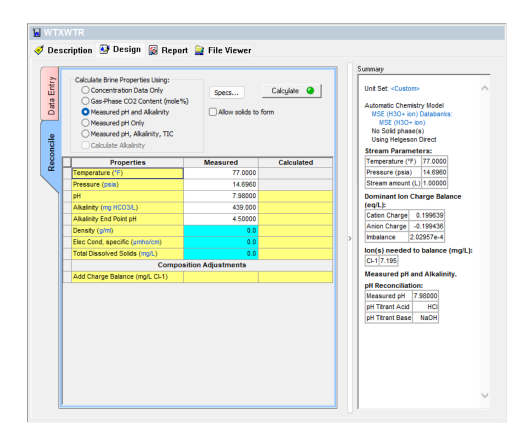


Temperature 77 F
Pressure 14.7 psia pH 7.98

Alkalinity end Point pH 439 mg/L as HCO3-

Titration pH 4.5

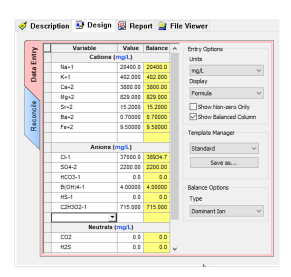
Density (will be estimated)



Click the Calculate Button

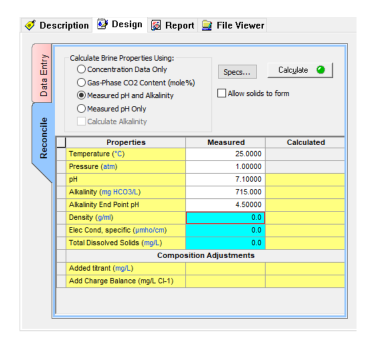
Task 2 - Add a Second Brine

Add a Brine Analysis, name it SSC-Brine Enter the composition given below in the **Data Entry** (vertical) Tab



Go to Reconcile tab

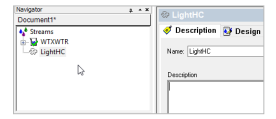
Select pH and Alkalinity calculation type, and enter the information provided in the figure below.



Click the Calculate button

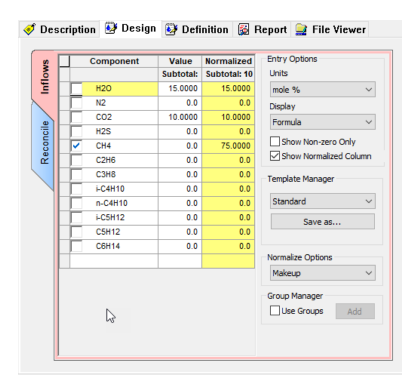
Task 3 - Add a Gas

From the Actions Panel click on *Add Gas Analysis* Input the name in the **Description** tab. Name it *LightHC*



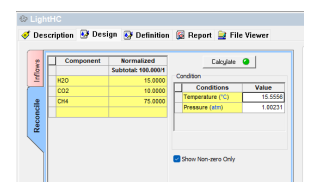
Go to the <u>Design</u> Tab and enter the following composition in the *Inflows* (vertical) tab Enter the following composition:

| Component | Composition |
|----------------------|-------------|
| Carbon dioxide (CO2) | 10 mole % |
| Water (H2O) | 15 mole % |



think simulation | getting the chemistry right will assume that all of the hydrocarbon gas is methane (CH4). So, in this case the software normalizes the gas adding 75 mole% of CH4.

Go to the Reconcile (vertical) Tab

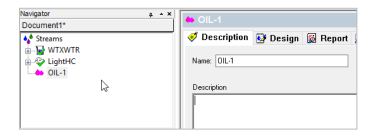


Leave the default conditions for Temperature and Pressure Calculate

Task 4 - Add an Oil

From the Actions Panel click *Add Oil Analysis*.

Enter the name in the **Description** Tab. Name it *OIL-1*



Go to the **Design** tab

This tab is divided into:

- Combined Tab: Here pure components (organic and inorganic) are entered.
- Pseudocomponent Tab
- Assay Tab: Here distillation curves are entered.
- Reconcile Tab

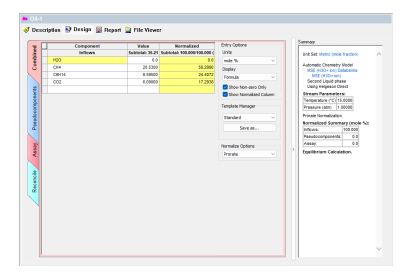
For this example, we will enter pure component and pseudocomponent data.

Go to the Combined tab

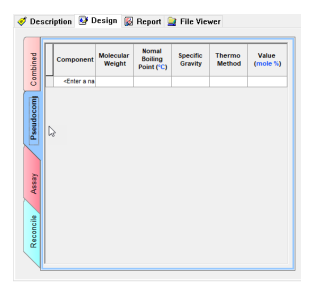
Enter the following composition:

Methane (CH4) 20.53mole % Hexane (C6H14) 8.595mole % CO2 6.09mole %

Note: You will have to scroll down to find the CO2 entry.



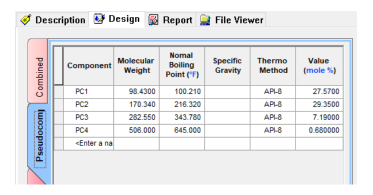
Go to the **Pseudocomponent** tab



Add the following pseudocomponents:

| Name | MW | nBP (F) | Thermo | Value |
|------|---------|---------|--------|--------|
| | (g/mol) | | Method | mole % |
| PC1 | 98.43 | 100.21 | API-8 | 27.57 |
| PC2 | 170.34 | 216.32 | API-8 | 29.35 |
| PC3 | 282.55 | 343.78 | API-8 | 7.19 |
| PC4 | 506.0 | 645.00 | API-8 | 0.68 |

The completed input looks like this:

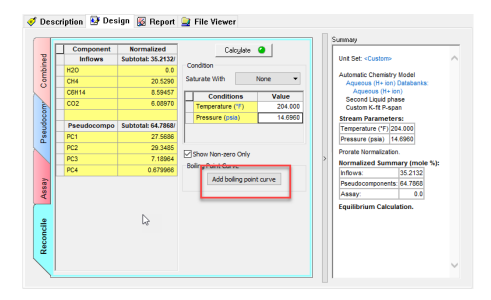


Go to the *Reconcile* (vertical) tab

Here you will see how well the Pseudocomponent will predict the phase behavior of the hydrocarbon sample.

In the Condition option (below calculate button) enter T=204 F and P=1 atm.

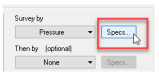
Click on the Add boiling point curve (red square shown)



A new calculation appears in the navigation panel below OIL1 Object.



This is a survey by Pressure. Click on the specs button and specify the conditions below. Change pressure units if necessary.

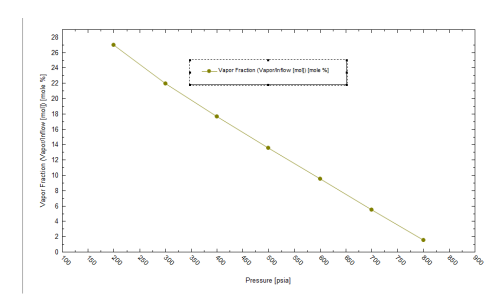


Enter a pressure range:

Start 200 psia End 2000 psia Increment 100 psia

Click the Calculate button.

Click on the plot tab, you will see the following graph: You could adjust the parameters by clicking on curves.



If the curve does not meet expectations, you will have to adjust the mole percentages or the pseudocomponent properties. This is a manual iterative approach. You can see the actual data by clicking the View data button.

| | Pressure | Vapor Fraction (Vapor/Inflow [mol]) |
|----|----------|---|
| | psia | mole % |
| 1 | 200.000 | 26.9611 |
| 2 | 300.000 | 21.9231 |
| 3 | 400.000 | 17.6477 |
| 4 | 500.000 | 13.5638 |
| 5 | 600.000 | 9.53472 |
| 6 | 700.000 | 5.52958 |
| 7 | 800.000 | 1.54051 |
| 8 | 900.000 | |
| 9 | 1000.00 | |
| 10 | 1100.00 | |
| 11 | 1200.00 | |
| 12 | 1300.00 | |
| 13 | 1400.00 | |
| 14 | 1500.00 | |
| 15 | 1600.00 | |
| 16 | 1700.00 | |
| 17 | 1800.00 | |
| 18 | 1900.00 | |
| 19 | 2000.00 | |

Frequently the hydrocarbon is saturated with water. We will simulate this part.

Go back to OIL-1 in the Action panel



Click on the **Design** tab

In the Conditions option use T= 77 F and 14.7 psia.

Click on the Calculate button.

When done, the software will create a *Reconciled OIL-1* in the Navigator pane.

The Brine, Gas and Oil that we already calculated will be used for the upcoming Mixer, Saturator and Facilities Calculations.

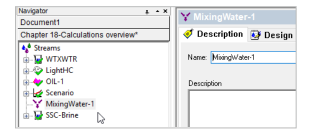
Task 5 – Adding a Mixer Calculation

The Mixer calculation determines if two waters (brines) can be mixed. Frequently the mixing of two waters will cause precipitates to form which were not present in original brines. This can lead to the plugging of a formation when an injection water is mixed with the natural fluids in the formation.

From the Actions Panel, click on the Add Mixing Water logo.



After double clicking the logo, you will the object in the Navigation Panel.



Click on the **Design** tab to enter more information.

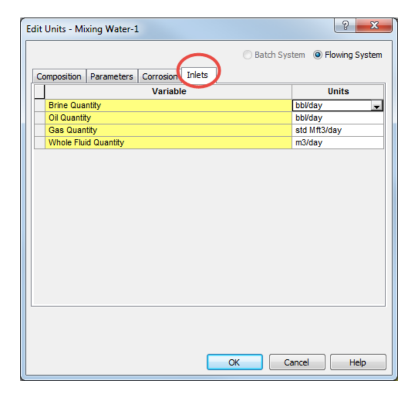
Select your Brines, Gas and Oil s as shown in the figure below. Enter the indicated brine flow rate of 1000 bbl/day, gas flow of 230 stdMft3/day and oil flow of 7 bbl/day.

Note: To change units on brine, oil and gas, click on the hyperlinked (**blue**) units inside of the bracket next to Inlets First Brine, Second Brine etc. This is under the column heading Type.

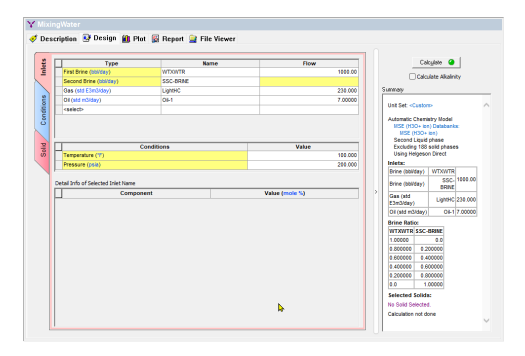


This should pop the following window up. Click on Inlets tab and change the units.

Inlets tab looks like below. Changing units on Brine:



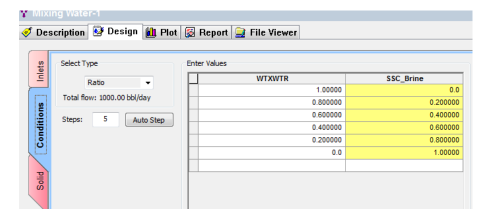
The final Mixer with inflows entered should look like the image below:



Go to the *Conditions* tab.

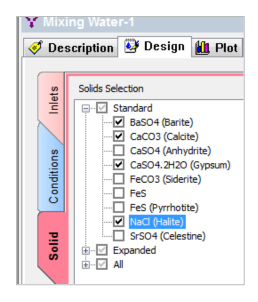
In this example, we are mixing the brines at 100 F and 200 psia. The first brine specified (WTXWTR) is the one we compare to when evaluating the ratios. In this case we start out with all brine WTXWTR and none of the SSC_Brine and end up with none of brine WTXWTR and all of the SSC_Brine.

Leave the default values



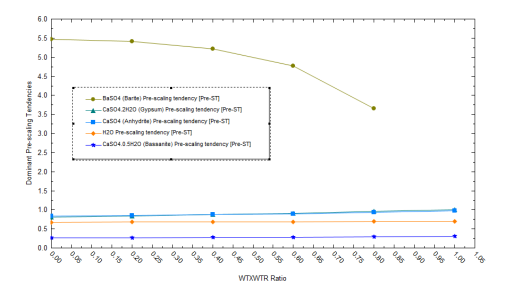
Go to the Solid tab

For this calculation we are only selecting a few of the possible solids. Mark a check box next to desired solids (BaSO4, CaCO3, CaSO4.2H2O, NaCl).



Click on the Calculate button.

Click on the Plot tab.



The ratio is relative to the first brine specified. This means at a ratio of 0.0 (all the first brine and none of the second) we have possible BaSO4 scaling. As we add the second brine, the amount of BaSO4 pre-scaling index decreases. These waters are perhaps incompatible.

Task 5 - Adding a Saturator Calculation

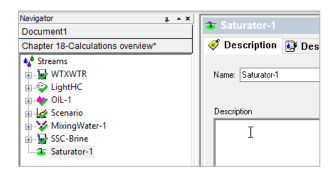
Water and gas samples at the surface are not necessarily representative of conditions in the reservoir. The processing of the samples may involve significant changes in the chemistry.

The Saturate option (often referred to as "Saturate at reservoir conditions") allows the user to "Back-calculate" the conditions downhole.

Select Add Saturator from the Actions Panel.



Rename the object in the descriptions tab. Name it Saturator-1



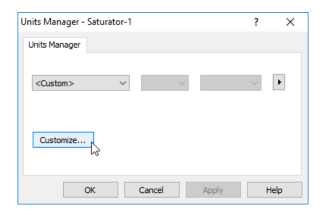
Go to **Design** tab And go to **Inlets** (vertical) tab

We will select objects that we have already defined.

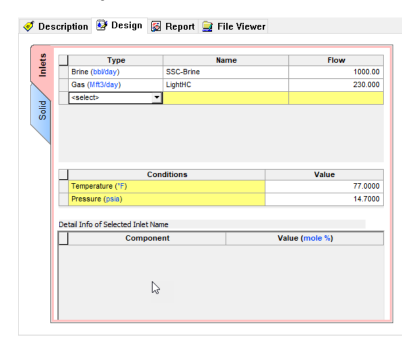
Enter a Brine flow of 1000 bbl/day (SSC-brine) Enter a Gas flow of 230 std Mft3/day (LightHC) Enter T=77 F and P=14.7 psia

Set these units as custom units for all new objects in Units manager.





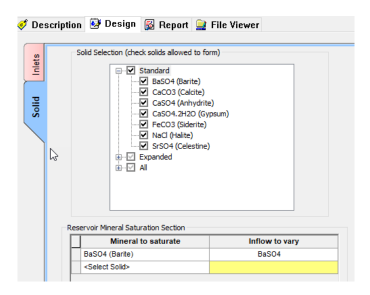
The window should look like the image below.



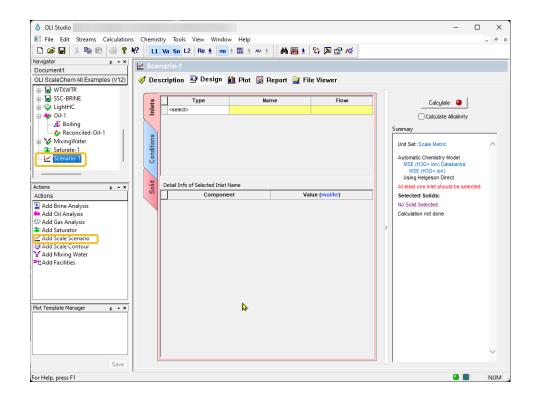
Go to **Solid** (vertical) tab Check the Standard box

We need to select solids inflow to vary for Saturator1. Under the table *Select Mineral to saturate*, choose the solid to vary from the dropdown list.

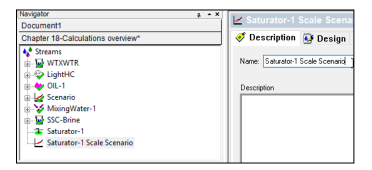
Select BaSO4 (solid) and hit enter. Automatically the inflow to vary BaSO4 will appear.



To study the Saturator-1 (Brine and Gas mix) at various locations we need to add a Scaling Scenario object.

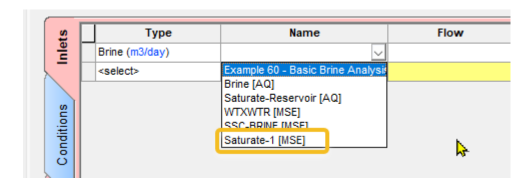


Name the new Scale Scenario as Saturator-1 Scale Scenario under the Description tab.



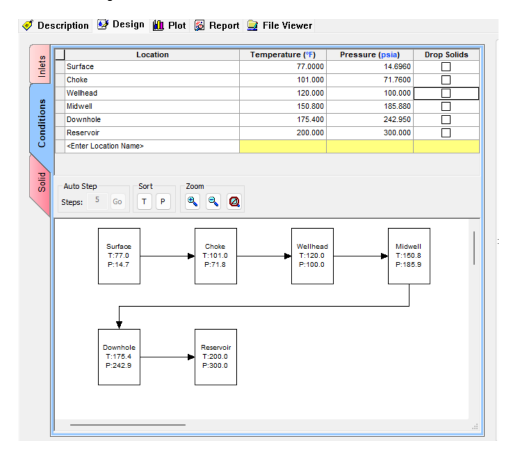
Go to the **Design** Tab

Under the **Inlets** (vertical) tab, select *Brine* under the *Type column* Make sure to select Saturator-1, as is shown in the picture below:

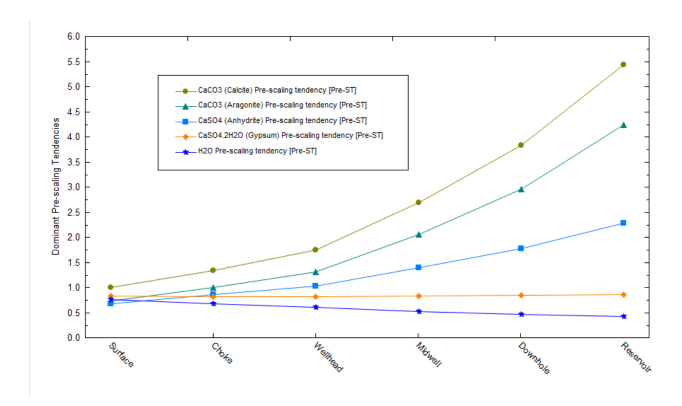


Flow will be automatically controlled. Locations can be input under the *Conditions* tab.

Go to **Conditions** (vertical) tab, and enter the information provided in the figure below:



Go to **Solid** tab and check the Standard box Click the Calculate Button Go to **Plot** tab



After the calculation is complete, the plot for scale scenario shows that other solids are appearing.

Go to the Report tab, and look for Pre and Post Scaling Tendencies

Pre and Post Scaling Tendencies

| Formula | Mineral | Excess Solute | Excess Solute | Pre-Scale | Pre-Scale | Post-Scale | Post-Scale |
|------------|--------------|---------------|---------------|------------|-------------|------------|------------|
| | | mg/L | lb/1000bbl | S, ST | SI, Index | S, ST | SI, Index |
| NaCl | Halite | 0.0 | 0.0 | 0.0114804 | -1.94004 | 0.0114804 | -1.94004 |
| BaSO4 | Barite | 0.0 | 0.0 | 1.00000 | 1.33309e-11 | 1.00000 | 0.0 |
| SrSO4 | Celestine | 0.0 | 0.0 | 0.191184 | -0.718548 | 0.191184 | -0.718548 |
| SrCO3 | Strontianite | | | 0.0155378 | -1.80861 | 0.0155378 | -1.80861 |
| CaCO3 | Calcite | 0.0 | 0.0 | 1.00000 | 3.02182e-11 | 1.00000 | 0.0 |
| CaSO4.2H2O | Gypsum | 0.0 | 0.0 | 0.825816 | -0.0831167 | 0.825816 | -0.0831167 |
| CaSO4 | Anhydrite | 0.0 | 0.0 | 0.678180 | -0.168655 | 0.678180 | -0.168655 |
| KCI | sylvite | | | 5.69739e-4 | -3.24432 | 5.69739e-4 | -3.24432 |

Excess solute or Max Scale: The solids amount forming at equilibrium.

Pre-Scale: The saturation ratio before solids precipitate

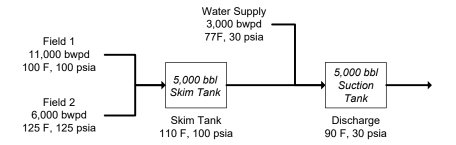
Post-Scale: The saturation ratio AFTER solids precipitate (if solids are selected).

 $S,\,ST-Saturation,\,Scale\,Tendency:\,The\,\,ratio\,\,of\,\,the\,\,concentration\,\,(activity)\,\,to\,\,its\,\,solubility\,\,(S=1).$

SI - Scale Index: Log(S).

Introducing the Facilities Object

This section presents a new calculation object: *Facilities*. The Facilities is a simplified process simulator; it mixes and separates. think simulation | getting the chemistry right has the ability to link together several individual calculations to create a flow sheet facility. An example of a facility calculation is shown in the figure below.



It is a simple process in which two field brines mix in a skim tank. The discharge from this tank then mixes with a water supply in a discharge tank. Below are the compositions and conditions of the inlet fluids.

| Name | Field 1 mg/L | Field 2 mg/L | Water Supply mg/L |
|---------------------------|-----------------|-----------------|----------------------|
| Na+ | 38209 | 27078 | 3074 |
| Ca+2 | 6600 | 4480 | 910 |
| Mg+2 | 1531 | 1191 | 249 |
| Fe+2 | 120 | 6.6 | 0.77 |
| CI-1 | 73150 | 51134 | 4474 |
| SO4-2 | 2453 | 1840 | 2960 |
| HCO3- | 421 | 677 | 439 |
| HS-1 | 244 | 146.2 | 0 |
| | | | |
| Temperature | 100 F | 125 F | 77 F |
| Pressure | 100 psia | 125 psia | 30 psia |
| pН | 6.97 | 7.53 | 7.98 |
| Alkalinity (As HCO3 mg/L) | 855 | 919 | 439 |
| Alkalinity End Point pH | 4.5 | 4.5 | 4.5 |

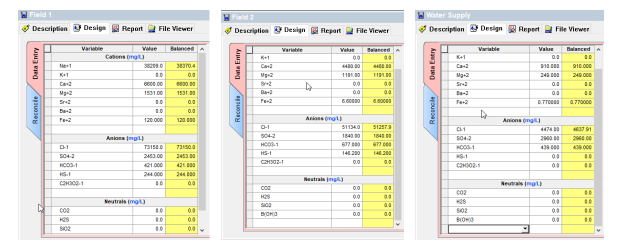
Follow the steps below:

Open a new Window for Simulation

Create the above brines: Field 1, Field 2 and Water Supply

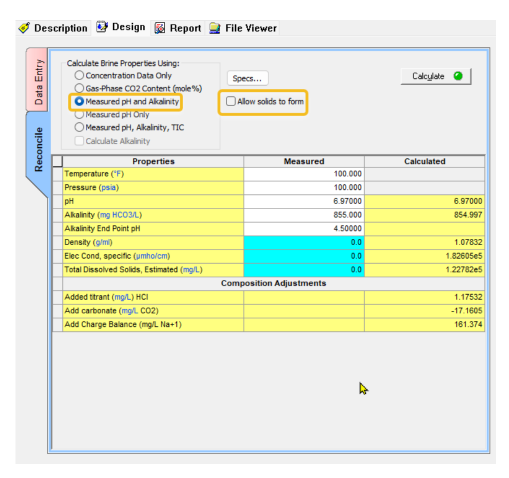
Use the Add Brine Analysis object as you have done before to create them.

The **Data Entry** Windows should look like this:



Reconcile these three brines for measured pH and alkalinity.

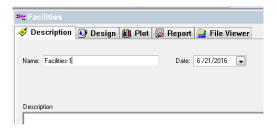
Make sure that the *Allow solids to form* box is uncheck at the bottom of the reconciliation options for all the brines.



Select Add Facilities from the Action Panel.



Go to the **Description** Tab and rename the object as Facilities-1

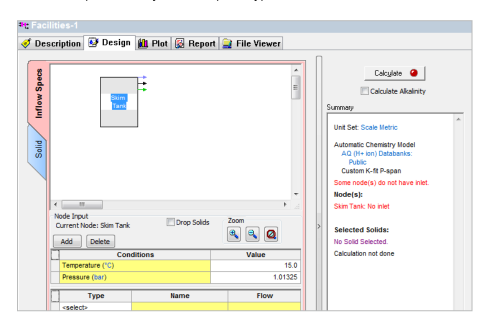


The facilities calculation is based upon transferring information between calculations through nodes. These nodes can be thought of as pseudo brines and gases. These pseudo brines are not stored as individual brine rather they are used internally in the calculation. The concentration and flow rates for these nodes can be viewed in the output.

Go to the **Design** tab
Go to the **Inflow Specs** (vertical tab)

Within Inflow Specs, we can add Nodes via Node input options. There will be one default node added.

Double click on the node name (where it says: Node 1) and type "Skim Tank".



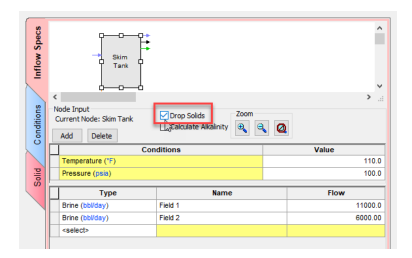
Enter the name, description, conditions, and streams for the Skim Tank shown in the table below:

| | Flow | Temperature, F | Pressure, psia |
|---------|---------------|----------------|----------------|
| Field 1 | 11000 bbl/day | 100 | 100 |

| Field 2 | 6000 bbl/day | 125 | 125 |
|--------------|--------------|-----|-----|
| Water Supply | 3000 bbl/day | 77 | 30 |
| Skim Tank | 5000 bbl/day | 110 | 100 |
| Suction Tank | 5000 bbl/day | 90 | 30 |

Make sure to select drop solids checkbox at the Skim Tank node.

When complete, your screen should look like this.

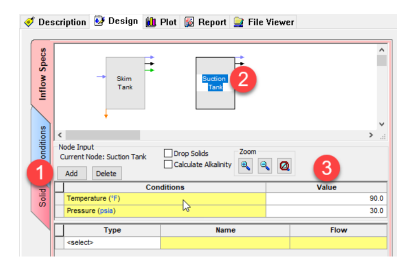


The output of this calculation will go to the next node.

Next, enter the information for the second node, Suction Tank. We will select the output brine of the Skim tank, which is a *brine from node*. The temperature and pressure as well as the flow of the brine are calculated.

Click on the Add button to add a second Node Change the name of the node to Suction Tank Change the conditions of this node to 90 F and 30 psia.

When complete your screen should look like the image below:



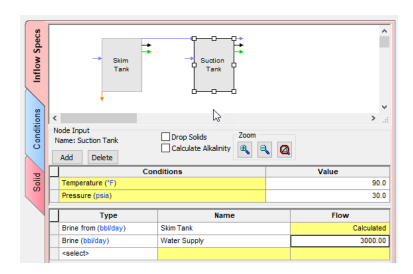
When a brine is calculated in a facilities calculation, we have the option of allowing any produced solids to be considered (that is they traveled along with the brine) or to **eliminate** them as they precipitate out. We will eliminate the solids in this case. The orange downward arrow from Skim Tank indicate dropped solids.

We are also adding the **Water Supply** to this tank.

In the Type column and first row select *Brine from*. In the Name column select *Skim Tank*. The *Flow* is calculated (since it comes from the Skim Tank).

In the Type column and second row select *Brine*. In the Name column select *Water Supply*. Enter 3000 bbl/day in the Flow column.

When complete the screen should look like the image below.



Click on the Calculate button, or press <Ctrl+F9>

Go to the **Report** tab, and check for the Pre-scaling Tendencies and Scaling Tendencies See the Results

Pre-Scaling Tendencies

Column Filter Applied: Values > 1.0e-4

Temperature Filter Applied: Active TRange Only.

| Nodes | CaCO3 (Aragonite) | CaCO3 (Calcite) | CaSO4.0.5H2O (Bassanite) | CaSO4.2H2O (Gypsum) | CaSO4 (Anhydrite) | FeCO3 (Siderite) |
|---------------|----------------------|-----------------|-----------------------------|------------------------|----------------------|------------------|
| Temp Range °C | Valid | Valid | Valid | Valid | Valid | Valid |
| Skim Tank | 19.2093 | 25.7515 | 0.334191 | 0.993874 | 1.08735 | 3.78201 |
| Suction Tank | 17.9342 | 24.1847 | 0.286645 | 0.972050 | 0.944722 | 4.06263 |

| Nodes | Fe(OH)2 (Amakinite) | FeS (Pyrrhotite) | H2O | FeS (Mackinawite) | MgCO3 | Mg(OH)2 (Brucite) |
|---------------|------------------------|------------------|----------|----------------------|------------|----------------------|
| Temp Range °C | Valid | Valid | Valid | Valid | Valid | Valid |
| Skim Tank | 2.72119e-4 | 37011.9 | 0.654343 | 714.959 | 8.91947e-3 | 1.88702e-4 |
| Suction Tank | 2.52127e-4 | 40916.2 | 0.696414 | 728.914 | 6.49943e-3 | 1.07033e-4 |

Based on these results, it is clear to see that the software predicts the formation of several solids in this process. Solids with a Pre-Scaling Tendency > 1 are predicted to form.