TC-PYTHON QUICK INSTALL GUIDE

This quick guide helps you to get a working TC-Python API installation.

There is a PDF guide included with your installation. In the Thermo-Calc menu, select Help → Manuals Folder. Then double-click to open the Software Development Kits (SDKs) folder.

Note: A license is required to run TC-Python.

1.1 Step 1: Install a Python Distribution

If you already have a Python distribution installation, version 3.5 or higher, skip this step.

These instructions are based on using the Anaconda platform for the Python distribution. Install version 3.5 or higher to be able to work with TC-Python, although it is recommended that you use the most recent version.

1.1.1 Install Anaconda

2. Click to choose your OS (operating system) and then click Download. Follow the instructions. It is recommended you keep all the defaults.

1.2 Step 2: Install Thermo-Calc and the TC-Python SDK

Note: TC-Python is available starting with Thermo-Calc version 2018a.

1. Install Thermo-Calc and choose a Custom installation.
   See Custom Standalone Installation in the Thermo-Calc Installation Guide.
   If you have already installed Thermo-Calc, you need to find the installation file (e.g. Windows *.exe, Mac *.zip and Linux *.run) to relaunch the installer and then continue with the next steps.
2. On the Select Components window, click to select the TC-Python check box.
3. On the Install TC-Python window, click Next.
4. When the installation is complete, the TC-Python folder opens and includes the *.whl file needed for the next step. There is also an Examples folder with Python files you can use in the IDE to understand and work with TC-Python.

The installation location for this API is the same as for other SDKs and based on the OS. For details, see Default Directory Locations in the Thermo-Calc Installation Guide.

1.3 Step 3: Install TC-Python

On Windows, it is recommended that you use the Python distribution prompt (i.e. Anaconda, . . .), especially if you have other Python installations. **Do not use Virtual Environments unless you have a good reason for that.**

1. Open the command line. For example, in Anaconda on a Windows OS, go to **Start→Anaconda→Anaconda Prompt**.

2. At the command line, enter the following. Make sure there are no spaces at the end of the string or in the folder name or it will not run:

   ```
   pip install <path to the TC-Python folder>/TC_Python-<version>-py3-none-any.whl
   ```

   For example, on a Windows OS Standalone custom installation, when you install for all users, the path to the TC-Python folder is `C:\Users\Public\Documents\Thermo-Calc\2018b\SDK\TC-Python\`

   Details for Mac and Linux installations are described in Default Directory Locations in the Thermo-Calc Installation Guide. Note that on Linux typically `pip3` is used.

3. Press `<Enter>`. When the process is completed, there is a confirmation that TC-Python is installed.

1.4 Step 4: Install an IDE (Integrated Development Environment)

Any editor can be used to write the Python code, but an IDE is recommended, e.g. PyCharm. These instructions are based on the use of PyCharm.

Use of an IDE will give you access to the IntelliSense, which is of great help when you use the API as it will give you the available methods on the objects you are working with.

2. Click to choose your OS and then click **Download**. You can use the **Community** version of PyCharm.
3. Follow the instructions. It is recommended you keep all the defaults.

**Note:** For Mac installations, you also need to set some environment variables as described below in Mac OS: Setting Environment Variables.

1.5 Step 5: Open the IDE and Run a TC-Python Example

After you complete all the software installations, you are ready to open the IDE to start working with TC-Python.

It is recommended that you open one or more of the included examples to both check that the installation has worked and to start familiarizing yourself with the code.
1.5.1 Open the TC-Python Project in PyCharm

When you first open the TC-Python project and examples, it can take a few moments for the Pycharm IDE to index before some of the options are available.

1. Open PyCharm and then choose File→Open. The first time you open the project you will need to navigate to the path of the TC-Python installation as done in Step 4.

   For example, on a Windows OS Standalone custom installation, when you install for all users, the path to the TC-Python folder is C:\Users\Public\Documents\Thermo-Calc\2018b\SDK\TC-Python\Details for Mac and Linux installations are described in the Default Directory Locations section in the Thermo-Calc Installation Guide.

2. Click on the Examples folder and then click OK.

3. From any subfolder:
   • Double-click to open an example file to examine the code.
   • Right-click an example and choose Run.

1.5.2 Fixing potential issues with the environment

In most cases you should run TC-Python within your global Python 3 interpreter and not use Virtual Environments unless you have a good reason to do so. If there are problems with the interpreter settings, you can resolve them in the settings window:

1. Go the menu File→Settings.
2. Navigate in the tree to Project.YourProjectName and choose Project Interpreter.
3. Click on the settings symbol close to the Project Interpreter dropdown menu and choose Add.
4. Now choose System Interpreter and add your existing Python 3 interpreter.
5. Select your added interpreter and confirm.

Note: If you are not following the recommended approach and create a new project (File→New Project...), you need to consider that by default the options to choose the interpreter are hidden within the Create Project window. So click on Project Interpreter: New Virtual Environment and in most cases choose your System Interpreter instead of the default New Virtual Environment.

Note: If you really need to use a Virtual Environment, please consider the hints given in the Best Practices chapter.

1.6 Updating to a newer version

When updating to a newer version of Thermo-Calc, you always need to also install the latest version of TC-Python. It is not sufficient to run the installer of Thermo-Calc. The procedure is generally identical to Step 3:

```
pip install <path to the TC-Python folder>/TC_Python-<version>-py3-none-any.whl
```

In case of problems you may wish to uninstall the previous version of TC-Python in advance:
pip uninstall TC-Python
pip install <path to the TC-Python folder>/TC_Python-<version>-py3-none-any.whl

However, that should normally not be required. Note that on Linux usually pip3 is used.

You can check the currently installed version of TC-Python by running:

```
pip show TC-Python
```
MAC OS: SETTING ENVIRONMENT VARIABLES

In order to use TC-Python on Mac you need to set some environment variables.

TC18B_HOME=/Applications/Thermo-Calc-2018b.app/Contents/Resources

If you use a license server:

LSHOST=<name-of-the-license-server>

If you have a node-locked license:

LSHOST= NO-NET LSEVRRC=/Applications/Thermo-Calc-2018b.app/Contents/Resources/lservrc

In PyCharm, you can add environment variables in the configurations.

Select Run→Edit Configurations to open the Run/Debug Configurations window. Enter the environment variable(s) by clicking the button to the right of the Environment Variables text field.
TC-Python contains classes of these types:

- **TCPython** – which is where you start
- **SystemBuilder** and **System** – where you choose database and elements etc.
- **Calculation** – where you choose and configure the calculation
- **Result** – where you get the results from a calculation you have run

### 3.1 TCPython

This is the starting point for all TC-Python usage.

You can think of this as the start of a “wizard”.

You use it to select databases and elements. That will take you to the next step in the wizard, where you configure the system.

Example

```python
from tc_python import *

with TCPython() as start:
    start.select_database_and_elements(...)  # e.t.c
    # after with clause

# or like this
with TCPython():
    SetUp().select_database_and_elements(...)  # e.t.c
    # after with clause
```

**Note:** This starts a process running a calculation server. Your code will then, via TC-Python, use socket communication to send and receive messages to and from that server. This is for information only.

When your Python script has run as far as to this row

```python
# after with clause
```

the calculation server automatically shuts down, and all temporary files will be deleted. To ensure that this happens, it is important that you structure your Python code using a `with()` clause, as the example above shows.
3.2 SystemBuilder and System

A SystemBuilder is returned when you have selected your database and elements in TCPython.
The SystemBuilder lets you further specify your system, for example with which phases that should be part of your system.

Example:

```python
from tc_python import *

with TCPython() as start:
    start.select_database_and_elements("ALDEMO", ["Al", "Sc"])
    # e.t.c
```

When all configuration is done, you call get_system() which returns an instance of a System class. The System class is fixed and can not be changed. If you later want to change the database, elements or something else, change the SystemBuilder and call get_system() again, or create a new SystemBuilder and call get_system() on that.

From the System you can create one or more calculations, which is the next step in the “wizard”.

**Note:** You can use the same System object to create several calculations.

3.3 Calculation

The best way to see how a calculation can be used, is in the TC-Python examples included with the Thermo-Calc installation.

Some calculations have many settings. Default values are used where it is applicable, and are overridden if you specify something different.

When you have configured your calculation you call calculate() to start the actual calculation. That returns a Result, which is the next step.

3.3.1 Single equilibrium calculations

In single equilibrium calculations you need to specify the correct number of conditions, depending on how many elements your System contains.

You do that by calling set_condition().

An important difference from other calculations is that single equilibrium calculations have two functions to get result values.

The calculate() method, which gives a Result, is used to get actual values. This result is “temporary”, meaning that if you run other calculations or rerun the current one, the resulting object will no longer give values corresponding to the first calculation.

This is different from how other calculations work. If you want a Result that you can use AFTER running other calculations, you need to call calculate_with_state().

**Note:** calculate() has MUCH better performance than calculate_with_state(), and works for almost all situations.
Example:

```python
from tc_python import *

with TCPython() as start:
    gibbs_energy = {
        start.
            select_database_and_elements("FEDEMO", ["Fe", "Cr", "C"]).
            get_system().
            with_single_equilibrium_calculation().
                set_condition(ThermodynamicQuantity.temperature(), 2000.0).
                set_condition(ThermodynamicQuantity.mole_fraction_of_a_component("Cr"), 0.1).
                set_condition(ThermodynamicQuantity.mole_fraction_of_a_component("C"), 0.01).
            calculate().
            get_value_of("G")
    }
```

### 3.3.2 Precipitation calculations

Everything that you can configure in the Precipitation Calculator in Thermo-Calc Graphical Mode, you can configure in this calculation. For the calculation to be possible, you need at least to enter a matrix phase, a precipitate phase, temperature, simulation time and compositions.

Example:

```python
from tc_python import *

with TCPython() as start:
    precipitation_curve = {
        start.
            select_thermodynamic_and_kinetic_databases_with_elements("ALDEMO", "MALDEMO", ["Al", "Sc"]).
            get_system().
            with_isothermal_precipitation_calculation().
                set_composition("Sc", 0.18).
                set_temperature(623.15).
                set_simulation_time(1e5).
                with_matrix_phase(MatrixPhase("FCC_A1").add_precipitate_phase(PrecipitatePhase("AL3SC"))).
                calculate()
    }
```

### 3.3.3 Scheil calculations

You can configure in this calculation everything that can also be configured in the Scheil Calculator in Thermo-Calc Graphical Mode or in the Console Mode. The minimum you need to specify are the elements and their composition. Everything else is set to a default value if you do not specify it explicitly.

Example:

```python
from tc_python import *

with TCPython() as start:
    (continues on next page)
```
temperature_vs_mole_fraction_of_solid = {
    start.
    select_database_and_elements("FEDEMO", ["Fe", "C"]).
    get_system().
    with_scheil_calculation().
    set_composition("C", 0.3).
    calculate().
    get_values_of(ScheilQuantity.temperature(),
                  ScheilQuantity.mole_fraction_of_all_solid_phases())
}

3.3.4 Property diagram calculations

For the property diagram (step) calculation, everything that you can configure in the \textit{Equilibrium Calculator} when choosing \textbf{Property diagram} in Thermo-Calc Graphical Mode can also be configured in this calculation. In Console Mode the property diagram is created using the Step command. The minimum you need to specify are elements, conditions and the calculation axis. Everything else is set to default values, if you don’t specify otherwise.

Example:

\begin{verbatim}
from tc_python import *

with TCPython() as start:
    property_diagram = {
        start.
        select_database_and_elements("FEDEMO", ["Fe", "C"]).
        get_system().
        with_property_diagram_calculation().
        with_axis(CalculationAxis(ThermodynamicQuantity.temperature()).
                  set_min(500).
                  set_max(3000)).
        set_condition(ThermodynamicQuantity.mole_fraction_of_a_component("C"),
                      0.01).
        calculate().
        get_values_grouped_by_stable_phases_of(ThermodynamicQuantity.
                                               temperature(),
                                               ThermodynamicQuantity.volume_
                                               fraction_of_a_phase("ALL"))
    }
\end{verbatim}

3.3.5 Phase diagram calculations

For the phase diagram (map) calculation, everything that you can configure in the \textit{Equilibrium Calculator} when choosing \textbf{Phase diagram} in Thermo-Calc Graphical Mode can also be configured in this calculation. In Console Mode the phase diagram is created using the Map command. The minimum you need to specify are elements, conditions and two calculation axes. Everything else is set to default values, if you don’t specify otherwise.

Example:

\begin{verbatim}
from tc_python import *

with TCPython() as start:
    phase_diagram = {
\end{verbatim}

(continues on next page)
3.4 Result

All calculations have a method called `calculate()` that starts the calculations and when finished, returns a `Result`. The `Result` classes have different methods, depending on the type of calculation. The `Result` is used to get numerical values from a calculation that has run.

Example:

```python
# code above sets up the calculation
r = calculation.calculate()
time, meanRadius = r.get_mean_radius_of("AL3SC")
```

The `Result` objects are completely independent from calculations done before or after they are created. The objects return valid values corresponding to the calculation they were created from, for their lifetime. The only exception is if you call `calculate()` and not `calculate_with_state()` on a single equilibrium calculation.

Example:

```python
# ...
# some code to set up a single equilibrium calculation
# ...

single_eq_result = single_eq_calculation.calculate_with_state()

# ...
# some code to set up a precipitation calculation
# ...

prec_result = precipitation_calculation.calculate()

# ...
# some code to set up a Scheil calculation
# ...
```
scheil_result = scheil_calculations.calculate()

# now it is possible to get results from the single equilibrium calculation,
# without having to re-run it (because it has been calculated with saving of the_
# state)

gibbs = single_eq_result.get_value_of("G")

In other words, you can mix different calculations and results without having to think about which state the calculation server is in.
4.1 All TC-Python objects are non-copyable

Never create a copy of an instance of a class in TC-Python, neither by using the Python builtin function `deepcopy()` nor in any other way. All classes in TC-Python are proxies for classes in the underlying calculation server and normally hold references to result files. A copied class object in Python would consequently point to the same classes and result files in the calculation server.

Instead of making a copy, always create a new instance:

```python
from tc_python import *

with TCPython() as start:
    system = start.select_database_and_elements("FEDEMO", ["Fe", "Cr"]).get_system()
    calculator = system.with_single_equilibrium_calculation()

    # *do not* copy the `calculator` object, create another one instead
    calculator_2 = system.with_single_equilibrium_calculation()

    # now you can use both calculators for different calculations ...
```

4.2 Python Virtual Environments

A Python installation can have several virtual environments. You can think of a virtual environment as a collection of third party packages that you have access to in your Python scripts. `tc_python` is such a package.

To run TC-Python, you need to install it into the same virtual environment as your Python scripts are running in. If your scripts fail on `import tc_python`, you need to execute the following command in the terminal of the same Python environment as your script is running in:

```
pip install TC_Python-<version>-py3-none-any.whl
```

If you use the PyCharm IDE, you should do that within the `Terminal` built into the IDE. This `Terminal` runs automatically within your actual (virtual) environment.

In order to prevent confusion, we recommend in most cases to install TC-Python within your global interpreter, for example by running the `pip install` command within your default Anaconda prompt.
4.3 *with TCPython()* should not be used within a loop

You should call *with TCPython()* only once within each process. When leaving the *with*-clause, all temporary data will be deleted and when entering the next *with*-clause a new Java backend engine process will be started. Currently the used Python-Java bridge *py4j* does not stop the old Java process. This will cause problems in case of many loop iterations.

In most use cases it is anyway considered as bad practice to call ‘with TCPython()’ more than once within a process. This is due to the high overhead caused by managing the resources (each time a process will be started, stopped and possibly a large amount of temporary data will be deleted).

To prevent calling *with TCPython()* multiple times and cleaning up temporary data anyway, you could use the following pattern:

```python
from tc_python import *

# ...
def calculation(calculator):
    # you could also pass the `session` or `system` object if more appropriate
    calculator.set_condition("W(Cr)*", 0.1)
    # further configuration ...

    result = calculator.calculate()
    # ...
    result.invalidate()  # if the temporary data needs to be cleaned up immediately

if __name__ == '__main__':
    with TCPython() as session:
        system = session.select_database_and_elements("FEDEMO", ["Fe", "Cr"]).get_  
        →system()
        calculator = system.with_single_equilibrium_calculation()

        for i in range(50):
            calculation(calculator)
```

The current behaviour will probably be changed in a future release to remove that limitation. It does not affect multi-processing applications (for example for parallelization of calculations).

4.4 Parallel calculations

It is possible to perform parallel calculations with TC-Python using multi-processing. Please note that multi-threading is not suitable for parallelization of computationally intensive tasks in Python. Additionally the Thermo-Calc core is not thread-safe. Using suitable Python-frameworks it is also possible to dispatch the calculations on different computers of a cluster.

A general pattern that can be applied is shown below. This code snippet shows how to perform single equilibrium calculations for different compositions in parallel. In the same way all other calculators of Thermo-Calc can be used or combined. For a real application, probably *numpy* arrays instead of Python arrays should be used for performance reasons.

```python
import concurrent.futures

from tc_python import *
```

(continues on next page)
def do_perform(parameters):
    # this function runs within an own process
    with TCPython() as start:
        elements = ["Fe", "Cr", "Ni", "C"]
        calculation = (start.select_database_and_elements("FEDEMO", elements).
                        get_system().
                        with_single_equilibrium_calculation().
                        set_condition("T", 1100).
                        set_condition("W(C)", 0.1 / 100).
                        set_condition("W(Ni)", 2.0 / 100))

        phase_fractions = []
        cr_contents = range(parameters["cr_min"],
                             parameters["cr_max"],
                             parameters["delta_cr"])
        for cr in cr_contents:
            result = (calculation.
                      set_condition("W(Cr)", cr / 100).
                      calculate())

            phase_fractions.append(result.get_value_of("NPM(BCC_A2)"))

    return phase_fractions

if __name__ == "__main__":
    parameters = [
    {"index": 0, "cr_min": 10, "cr_max": 15, "delta_cr": 1},
    {"index": 1, "cr_min": 15, "cr_max": 20, "delta_cr": 1}]

    bcc_phase_fraction = []
    num_processes = 2

    with concurrent.futures.ProcessPoolExecutor(num_processes) as executor:
        for result_from_process in zip(parameters, executor.map(do_perform, parameters)):
            # params can be used to identify the process and its parameters
            params, phase_fractions_from_process = result_from_process
            bcc_phase_fraction.extend(phase_fractions_from_process)

        # use the result in 'bcc_phase_fraction', for example for plotting
API REFERENCE

5.1 Calculations

5.1.1 Module “single_equilibrium”

```python
class tc_python.single_equilibrium.SingleEquilibriumCalculation(calculator)
Bases: tc_python.abstract_base.AbstractCalculation
```

Configuration for a single equilibrium calculation.

**Note:** Specify the conditions and possibly other settings, the calculation is performed with `calculate()`.

`calculate()` → tc_python.single_equilibrium.SingleEquilibriumTempResult
Performs the calculation and provides a temporary result object that is only valid until something gets changed in the calculation state. The method `calculate()` is the default approach and should be used in most cases.

**Returns** A new `SingleEquilibriumTempResult` object which can be used to get specific values from the calculated result. It is undefined behaviour to use that object after the state of the calculation has been changed.

**Warning:** If the result object should be valid for the whole program lifetime, use `calculate_with_state()` instead.

`calculate_with_state()` → tc_python.single_equilibrium.SingleEquilibriumResult
Performs the calculation and provides a result object that will reflect the present state of the calculation during the whole lifetime of the object. This method comes with a performance and temporary disk space overhead. It should only be used if it is necessary to access the result object again later after the state has been changed. In most cases you should use the method `calculate()`.

**Returns** A new `SingleEquilibriumResult` object which can be used later at any time to get specific values from the calculated result.

`disable_global_minimization()`
Turns the global minimization completely off.

**Returns** This `SingleEquilibriumCalculation` object

`enable_global_minimization()`
Turns the global minimization on (using the default settings).

**Returns** This `SingleEquilibriumCalculation` object
**get_components()** → List[str]

Returns a list of components in the system.

**remove_all_conditions()**

Removes all set conditions.

**Parameters**

- None

**Returns**

This SingleEquilibriumCalculation object

**remove_condition(quantity: Union[tc_python.quantity_factory.ThermodynamicQuantity, str])**

Removes the specified condition.

**Parameters**

- **quantity** – the ThermodynamicQuantity to set as condition, a console syntax string can be used as an alternative (for example “X(Cr)”)  

**Returns**

This SingleEquilibriumCalculation object

**run_poly_command(command: str)**

Runs a Thermo-Calc command from the console POLY-module immediately in the engine.

**Parameters**

- **command** – The Thermo-Calc console command

**Returns**

This SingleEquilibriumCalculation object

**Note:** It should not be necessary for most users to use this method, try to use the corresponding method implemented in the API instead.

**Warning:** As this method runs raw Thermo-Calc commands directly in the engine, it may hang the program in case of spelling mistakes (e.g. forgotten equals sign).

**set_component_to_entered(component: str)**

Sets the specified component to the status ENTERED, that is the default state.

**Parameters**

- **component** – The component name

**Returns**

This SingleEquilibriumCalculation object

**set_component_to_suspended(component: str)**

Sets the specified component to the status SUSPENDED, i.e. it is ignored in the calculation.

**Parameters**

- **component** – The component name

**Returns**

This SingleEquilibriumCalculation object

**set_condition(quantity: Union[tc_python.quantity_factory.ThermodynamicQuantity, str], value: float)**

Sets the specified condition.

**Parameters**

- **quantity** – The ThermodynamicQuantity to set as condition, a console syntax string can be used as an alternative (for example “X(Cr)”)
- **value** – The value of the condition

**Returns**

This SingleEquilibriumCalculation object

**set_phase_to_dormant(phase: str)**

Sets the phase to the status DORMANT, necessary for calculating the driving force to form the specified phase.

**Parameters**

- **phase** – The phase name
Returns This `SingleEquilibriumCalculation` object

**set_phase_to_entered**(phase: str, amount: float)
Sets the phase to the status ENTERED, that is the default state.

Parameters

• **phase** – The phase name
• **amount** – The phase fraction (between 0.0 and 1.0)

Returns This `SingleEquilibriumCalculation` object

**set_phase_to_fixed**(phase: str, amount: float)
Sets the phase to the status FIXED, i.e. it is guaranteed to have the specified phase fraction after the calculation.

Parameters

• **phase** – The phase name
• **amount** – The fixed phase fraction (between 0.0 and 1.0)

Returns This `SingleEquilibriumCalculation` object

**set_phase_to_suspended**(phase: str)
Sets the phase to the status SUSPENDED, i.e. it is ignored in the calculation.

Parameters **phase** – The phase name

Returns This `SingleEquilibriumCalculation` object

**class tc_python.single_equilibrium.SingleEquilibriumResult**(result)
**Bases:** `tc_python.abstract_base.AbstractResult`

Result of a single equilibrium calculation, it can be evaluated using a Quantity or Console Mode syntax.

**get_components**() → List[str]
Returns the components selected in the system (including any components auto-selected by the database(s)).

Returns The selected components

**get_phases**() → List[str]
Returns the phases present in the system due to its configuration. It also contains all phases that have been automatically added during the calculation, this is the difference to the method `System.get_phases_in_system()`.

Returns The phases in the system including automatically added phases

**get_stable_phases**() → List[str]
Returns the stable phases (i.e. the phases present in the current equilibrium).

Returns The stable phases

**get_value_of**(quantity: Union[tc_python.quantity_factory.ThermodynamicQuantity, str]) → float
Returns a value from a single equilibrium calculation.

Parameters **quantity** – the ThermodynamicQuantity to get the value of, a console syntax strings can be used as an alternative (for example “NPM(FCC_A1)”)  

Returns The requested value

**class tc_python.single_equilibrium.SingleEquilibriumTempResult**(result)
**Bases:** `tc_python.abstract_base.AbstractResult`

5.1. Calculations
Result of a single equilibrium calculation that is only valid until something gets changed in the calculation state. It can be evaluated using a Quantity or Console Mode syntax.

**Warning:** Note that it is undefined behaviour to use that object after something has been changed in the state of the calculation, this will result in an `InvalidResultStateException` exception being raised.

### get_components() → List[str]
Returns the components selected in the system (including any components auto-selected by the database(s)).

- **Returns** The selected components
- **Raises** `InvalidResultStateException` – If something has been changed in the state of the calculation since that result object has been created

### get_phases() → List[str]
Returns the phases present in the system due to its configuration. It also contains all phases that have been automatically added during the calculation, this is the difference to the method `System.get_phases_in_system()`.

- **Returns** The phases in the system including automatically added phases
- **Raises** `InvalidResultStateException` – If something has been changed in the state of the calculation since that result object has been created

### get_stable_phases() → List[str]
Returns the stable phases (i.e. the phases present in the current equilibrium).

- **Returns** The stable phases
- **Raises** `InvalidResultStateException` – If something has been changed in the state of the calculation since that result object has been created

### get_value_of(quantity: Union[tc_python.quantity_factory.ThermodynamicQuantity, str]) → float
Returns a value from a single equilibrium calculation.

- **Parameters** `quantity` – the ThermodynamicQuantity to get the value of, a console syntax strings can be used as an alternative (for example “NPM(FCC_A1)”)  
- **Returns** The requested value
- **Raises** `InvalidResultStateException` – If something has been changed in the state of the calculation since that result object has been created

### Module “precipitation”

**class** `tc_python.precipitation.GrowthRateModel`

**Bases:** `enum.Enum`

Choice of the used growth rate model for a precipitate.

- **ADVANCED** = 3

The ADVANCED MODEL was been proposed by Chen, Jeppsson, and Ågren (CJA) (2008) and calculates the velocity of a moving phase interface in multicomponent systems by identifying the operating tie-line from the solution of the flux-balance equations. This model can treat both high supersaturation and cross diffusion rigorously. Spontaneous transitions between different modes (LE and NPLE) of phase transformation can be captured without any ad-hoc treatment.
Note: Since it is not always possible to solve the flux-balance equations and it takes time, and where possible, use a less rigorous but simple and efficient model is preferred.

SIMPLIFIED = 2
The SIMPLIFIED MODEL is based on the advanced model but avoids the difficulty to find the operating tie-line and uses the tie-line across the bulk composition. This is the default growth rate model.

class tc_python.precipitation.MatrixPhase(matrix_phase_name: str)
Bases: object

The matrix phase in a precipitation calculation

    add_precipitate_phase(precipitate_phase: tc_python.precipitation.PrecipitatePhase)
    Adds a precipitate phase.

    Parameters precipitate_phase – The precipitate phase

    set_dislocation_density(dislocation_density: float = 5000000000000.0)
    Enter a numerical value. Default: 5.0E12 m^-2.

    Parameters dislocation_density – The dislocation density [m^-2]

    set_grain_aspect_ratio(grain_aspect_ratio: float = 1.0)
    Enter a numerical value. Default: 1.0.

    Parameters grain_aspect_ratio – The grain aspect ratio [-]

    set_grain_radius(grain_radius: float = 0.0001)
    Sets grain radius / size. Default: 1.0E-4 m

    Parameters grain_radius – The grain radius / size [m]

    set_mobility_enhancement_activation_energy(mobility_enhancement_activation_energy: float = 0.0)
    A value that adds to the activation energy of mobility data from the database. Default: 0.0 J/mol

    Parameters mobility_enhancement_activation_energy – The value that adds to the activation energy of mobility data from the database [J/mol].

    set_mobility_enhancement_prefactor(mobility_enhancement_prefactor: float = 1.0)
    A parameter that multiplies to the mobility data from database. Default: 1.0

    Parameters mobility_enhancement_prefactor – The mobility enhancement factor [-]

    set_molar_volume(volume: float)
    Sets the molar volume of the phase.

    Default: If not set, the molar volume is taken from the thermodynamic database (or set to 7.0e-6 m^3/mol if the database contains no molar volume information).

    Parameters volume – The molar volume [m^3/mol]

    with_elastic_properties_cubic(c11: float, c12: float, c44: float)
    Sets the elastic properties to “cubic” and specifies the elastic stiffness tensor components. Default: if not chosen, the default is DISREGARD

    Parameters
      • c11 – The stiffness tensor component c11 [GPa]
      • c12 – The stiffness tensor component c12 [GPa]
      • c44 – The stiffness tensor component c44 [GPa]
with_elastic_properties_disregard()
    Set to disregard to ignore the elastic properties. Default: This is the default option

with_elastic_properties_isotropic(shear_modulus: float, poisson_ratio: float)
    Sets elastic properties to isotropic. Default: if not chosen, the default is DISREGARD

    Parameters
        • shear_modulus – The shear modulus [GPa]
        • poisson_ratio – The Poisson’s ratio [-]

class tc_python.precipitation.NumericalParameters
    Bases: object

    Numerical parameters

    set_max_overall_volume_change(max_overall_volume_change: float = 0.001)
        This defines the maximum absolute (not ratio) change of the volume fraction allowed during one time step.
        Default: 0.001

        Parameters max_overall_volume_change – The maximum absolute (not ratio) change of the volume fraction allowed during one time step [-]

    set_max_radius_points_per_magnitude(max_radius_points_per_magnitude: float = 200.0)
        Sets the maximum number of grid points over one order of magnitude in radius. Default: 200.0

        Parameters max_radius_points_per_magnitude – The maximum number of grid points over one order of magnitude in radius [-]

    set_max_rel_change_critical_radius(max_rel_change_critical_radius: float = 0.1)
        Used to place a constraint on how fast the critical radius can vary, and thus put a limit on time step.
        Default: 0.1

        Parameters max_rel_change_critical_radius – The maximum relative change of the critical radius [-]

    set_max_rel_change_nucleation_rate_log(max_rel_change_nucleation_rate_log: float = 0.5)
        This parameter ensures accuracy for the evolution of effective nucleation rate. Default: 0.5

        Parameters max_rel_change_nucleation_rate_log – The maximum logarithmic relative change of the nucleation rate [-]

    set_max_rel_radius_change(max_rel_radius_change: float = 0.01)
        The maximum value allowed for relative radius change in one time step. Default: 0.01

        Parameters max_rel_radius_change – The maximum relative radius change in one time step [-]

    set_max_rel_solute_composition_change(max_rel_solute_composition_change: float = 0.01)
        Set a limit on the time step by controlling solute depletion or saturation, especially at isothermal stage.
        Default: 0.01

        Parameters max_rel_solute_composition_change – The limit for the relative solute composition change [-]

    set_max_time_step(max_time_step: float = 0.1)
        The maximum time step allowed for time integration as fraction of the simulation time. Default: 0.1

        Parameters max_time_step – The maximum time step as fraction of the simulation time [-]

    set_max_time_step_during_heating(max_time_step_during_heating: float = 1.0)
        The upper limit of the time step that has been enforced in the heating stages. Default: 1.0 s
Parameters **max_time_step_during_heating** – The maximum time step during heating [s]

**set_max_volume_fraction_dissolve_time_step**

```python
(max_volume_fraction_dissolve_time_step: float = 0.01)
```

Sets the maximum volume fraction of subcritical particles allowed to dissolve in one time step. **Default:** 0.01

Parameters **max_volume_fraction_dissolve_time_step** – The maximum volume fraction of subcritical particles allowed to dissolve in one time step [-]

**set_min_radius_nucleus_as_particle**

```python
(min_radius_nucleus_as_particle: float = 5e-10)
```

The cut-off lower limit of precipitate radius. **Default:** 5.0E-10 m

Parameters **min_radius_nucleus_as_particle** – The minimum radius of a nucleus to be considered as a particle [m]

**set_min_radius_points_per_magnitude**

```python
(min_radius_points_per_magnitude: float = 100.0)
```

Sets the minimum number of grid points over one order of magnitude in radius. **Default:** 100.0

Parameters **min_radius_points_per_magnitude** – The minimum number of grid points over one order of magnitude in radius [-]

**set_radius_points_per_magnitude**

```python
(radius_points_per_magnitude: float = 150.0)
```

Sets the number of grid points over one order of magnitude in radius. **Default:** 150.0

Parameters **radius_points_per_magnitude** – The number of grid points over one order of magnitude in radius [-]

**set_rel_radius_change_class_collision**

```python
(rel_radius_change_class_collision: float = 0.5)
```

Sets the relative radius change for avoiding class collision. **Default:** 0.5

Parameters **rel_radius_change_class_collision** – The relative radius change for avoiding class collision [-]

---

```python
class tc_python.precipitation.ParticleSizeDistribution
```

**Bases:** object

Represents the state of a microstructure evolution at a certain time including its particle size distribution, composition and overall phase fraction.

**add_radius_and_number_density**

```python
(radius: float, number_density: float)
```

Adds a radius and number density pair to the particle size distribution.

**Parameters**

- **radius** – The radius [m]
- **number_density** – The number of particles per unit volume per unit length [m^-4]

**Returns** This `ParticleSizeDistribution` object

**set_initial_composition**

```python
(element: str, composition_value: float)
```

Sets the initial precipitate composition.

**Parameters**

- **element** – The element
- **composition_value** – The composition value [composition unit defined for the calculation]

**Returns** This `ParticleSizeDistribution` object

---

5.1. Calculations
**set_volume_fraction_of_phase_type**(*volume_fraction_of_phase_type_enum: tc_python.precipitation.VolumeFractionOfPhaseType*)

Sets the type of the phase fraction or percentage. **Default**: By default volume fraction is used.

**Parameters**
- **volume_fraction_of_phase_type_enum** – Specifies if volume percent or fraction is used

**Returns**
This *ParticleSizeDistribution* object

**set_volume_fraction_of_phase_value**(*value: float*)

Sets the overall volume fraction of the phase (unit based on the setting of *set_volume_fraction_of_phase_type()*).

**Parameters**
- **value** – The volume fraction 0.0 - 1.0 or percent value 0 - 100

**Returns**
This *ParticleSizeDistribution* object

**class** *tc_python.precipitation.PrecipitateElasticProperties*

**Bases**: object

Represents the elastic transformation strain of a certain precipitate class.

**Note**: This class is only relevant if the option *TransformationStrainCalculationOption.USER_DEFINED* has been chosen using *PrecipitatePhase.set_transformation_strain_calculation_option()*.

The elastic strain can only be considered for non-spherical precipitates.

**set_e11**(*e11: float*)

Sets the elastic strain tensor component e11. **Default**: 0.0

**Parameters**
- **e11** – The elastic strain tensor component e11

**Returns**
This *PrecipitateElasticProperties* object

**set_e12**(*e12: float*)

Sets the strain tensor component e12. **Default**: 0.0

**Parameters**
- **e12** – The elastic strain tensor component e12

**Returns**
This *PrecipitateElasticProperties* object

**set_e13**(*e13: float*)

Sets the elastic strain tensor component e13. **Default**: 0.0

**Parameters**
- **e13** – The elastic strain tensor component e13

**Returns**
This *PrecipitateElasticProperties* object

**set_e22**(*e22: float*)

Sets the elastic strain tensor component e22. **Default**: 0.0

**Parameters**
- **e22** – The elastic strain tensor component e22

**Returns**
This *PrecipitateElasticProperties* object

**set_e23**(*e23: float*)

Sets the elastic strain tensor component e23. **Default**: 0.0

**Parameters**
- **e23** – The elastic strain tensor component e23

**Returns**
This *PrecipitateElasticProperties* object

**set_e33**(*e33: float*)

Sets the elastic strain tensor component e33. **Default**: 0.0

**Parameters**
- **e33** – The elastic strain tensor component e33
Returns This PrecipitateElasticProperties object

class tc_python.precipitation.PrecipitateMorphology
    Bases: enum.Enum
    Available precipitate morphologies.
    
    CUBOID = 3
    Cuboidal precipitates, only available for bulk nucleation.
    
    NEEDLE = 1
    Needle-like precipitates, only available for bulk nucleation.
    
    PLATE = 2
    Plate-like precipitates, only available for bulk nucleation.
    
    SPHERE = 0
    Spherical precipitates, this is the default morphology.

class tc_python.precipitation.PrecipitatePhase(precipitate_phase_name: str)
    Bases: object
    Represents a certain precipitate class (i.e. a group of precipitates with the same phase and settings).
    
    disable_calculate_aspect_ratio_from_elastic_energy()
    Disables the automatic calculation of the aspect ratio from the elastic energy of the phase.
    
    Returns This PrecipitatePhase object

    Note: If you use this method, you are required to set the aspect ratio explicitly using the method
    set_aspect_ratio_value().

    Default: This is the default setting (with an aspect ratio of 1.0).

    disable_driving_force_approximation()
    Will disable driving force approximation for this precipitate class. Default: Driving force approximation
    is disabled.
    
    Returns This PrecipitatePhase object

    enable_calculate_aspect_ratio_from_elastic_energy()
    Enables the automatic calculation of the aspect ratio from the elastic energy of the phase. Default: The
    aspect ratio is set to a value of 1.0.
    
    Returns This PrecipitatePhase object

    enable_driving_force_approximation()
    Will enable driving force approximation for this precipitate class. This approximation is often required
    when simulating precipitation of multiple particles that use the same phase description. E.g. simultaneous
    precipitation of a Metal-Carbide(MC) and Metal-Nitride(MN) if configured as different composition sets
    of the same phase FCC_A1. Default: Driving force approximation is disabled.
    
    Returns This PrecipitatePhase object

    Tip: Use this if simulations with several compositions sets of the same phase cause problems.

set_alias(alias: str)
Sets an alias string that can later be used to get values from a calculated result. Typically used when having
the same phase for several precipitates, but with different nucleation sites. For example two precipitates of
the phase M7C3 with nucleation sites in ‘Bulk’ and at ‘Dislocations’. The alias can be used instead of the phase name when retrieving simulated results.

**Parameters**

- **alias**
  - The alias string for this class of precipitates

**Returns**

This PrecipitatePhase object

**Note:** Typically used when having using the same precipitate phase, but with different settings in the same calculation.

---

**set_aspect_ratio_value**

Sets the aspect ratio of the phase. **Default:** An aspect ratio of 1.0.

**Parameters**

- **aspect_ratio_value**
  - The aspect ratio value

**Returns**

This PrecipitatePhase object

**Note:** Only relevant if `disable_calculate_aspect_ratio_from_elastic_energy()` is used (which is the default).

---

**set_gibbs_energy_addition**

Sets a Gibbs energy addition to the Gibbs energy of the phase. **Default:** 0.0 J/mol

**Parameters**

- **gibbs_energy_addition**
  - The Gibbs energy addition [J/mol]

**Returns**

This PrecipitatePhase object

---

**set_interfacial_energy**

Sets the interfacial energy. **Default:** If the interfacial energy is not set, it gets automatically calculated using a broken-bond model.

**Parameters**

- **interfacial_energy**
  - The interfacial energy [J/m²]

**Returns**

This PrecipitatePhase object

**Note:** The calculation of the interfacial energy using a broken-bond model is based on the assumption of an interface between a bcc- and a fcc-crystal structure with (110) and (111) lattice planes regardless of the actual phases.

---

**set_interfacial_energy_estimation_prefactor**

Sets the interfacial energy prefactor. **Default:** Prefactor of 1.0 (only relevant if the interfacial energy is automatically calculated).

**Parameters**

- **interfacial_energy_estimation_prefactor**
  - The prefactor for the calculated interfacial energy

**Returns**

This PrecipitatePhase object

**Note:** The interfacial energy prefactor is an amplification factor for the automatically calculated interfacial energy. Example: `interfacial_energy_estimation_prefactor = 2.5 => 2.5 * calculated interfacial energy`

---

**set_molar_volume**

Sets the molar volume of the precipitate phase. **Default:** The molar volume obtained from the database. If no molar volume information is present in the database, a value of 7.0e-6 m³/mol is used.
Parameters `volume` – The molar volume [m^3/mol]

Returns This `PrecipitatePhase` object

`set_nucleation_at_dislocations` *(number_density=-1)*
Activates nucleation at dislocations for this class of precipitates. Calling the method overrides any other nucleation setting for this class of precipitates. Default: If not set, by default bulk nucleation is chosen.

Parameters `number_density` – Number density of nucleation sites. If not set, the value is calculated based on the matrix settings (grain size, dislocation density) [m^-3].

Returns This `PrecipitatePhase` object

`set_nucleation_at_grain_boundaries` *(wetting_angle: float = 90.0, number_density: float = -1)*
Activates nucleation at grain boundaries for this class of precipitates. Calling the method overrides any other nucleation setting for this class of precipitates. Default: If not set, by default bulk nucleation is chosen.

Parameters

- `wetting_angle` – If not set, a default value of 90 degrees is used [degrees]
- `number_density` – Number density of nucleation sites. If not set, the value is calculated based on the matrix settings (grain size) [m^-3].

Returns This `PrecipitatePhase` object

`set_nucleation_at_grain_corners` *(wetting_angle: float = 90, number_density: float = -1)*
Activates nucleation at grain corners for this class of precipitates. Calling the method overrides any other nucleation setting for this class of precipitates. Default: If not set, by default bulk nucleation is chosen.

Parameters

- `wetting_angle` – If not set, a default value of 90 degrees is used [degrees]
- `number_density` – Number density of nucleation sites. If not set, the value is calculated based on the matrix settings (grain size) [m^-3].

Returns This `PrecipitatePhase` object

`set_nucleation_at_grain_edges` *(wetting_angle=90, number_density=-1)*
Activates nucleation at the grain edges for this class of precipitates. Calling the method overrides any other nucleation setting for this class of precipitates. Default: If not set, by default bulk nucleation is chosen.

Parameters

- `wetting_angle` – If not set, a default value of 90 degrees is used [degrees]
- `number_density` – Number density of nucleation sites. If not set, the value is calculated based on the matrix settings (grain size) [m^-3].

Returns This `PrecipitatePhase` object

`set_nucleation_in_bulk` *(number_density: float = -1)*
Activates nucleation in the bulk for this class of precipitates. Calling the method overrides any other nucleation setting for this class of precipitates. Default: This is the default setting (with an automatically calculated number density).

Parameters `number_density` – Number density of nucleation sites. If not set, the value is calculated based on the matrix settings (molar volume) [m^-3]

Returns This `PrecipitatePhase` object

`set_phase_boundary_mobility` *(phase_boundary_mobility: float)*
Sets the phase boundary mobility. Default: 10.0 m^4/(Js).
Parameters phase_boundary_mobility – The phase boundary mobility \[ \text{m}^4/(\text{Js}) \]

Returns This PrecipitatePhase object

set_precipitate_morphology (precipitate_morphology_enum: tc_python.precipitation.PrecipitateMorphology)

Sets the precipitate morphology. Default: PrecipitateMorphology.SPHERE

Parameters precipitate_morphology_enum – The precipitate morphology

Returns This PrecipitatePhase object

set_transformation_strain_calculation_option (transformation_strain_calculation_option_enum: tc_python.precipitation.TransformationStrainCalculationOption)

Sets the transformation strain calculation option. Default: TransformationStrainCalculationOption.DISREGARD.

Parameters transformation_strain_calculation_option_enum – The chosen option

Returns This PrecipitatePhase object

set_wetting_angle (wetting_angle: float)

Sets the wetting angle. Only relevant if the activated nucleation sites use that setting. Default: A wetting angle of 90 degrees.

Parameters wetting_angle – The wetting angle [degrees]

Returns This PrecipitatePhase object

with_elastic_properties (elastic_properties: tc_python.precipitation.PrecipitateElasticProperties)

Sets the elastic properties. Default: The elastic transformation strain is disregarded by default.

Parameters elastic_properties – The elastic properties object

Returns This PrecipitatePhase object

Note: This method has only an effect if the option TransformationStrainCalculationOption.USER_DEFINED has been chosen using the method set_transformation_strain_calculation_option().

with_growth_rate_model (growth_rate_model_enum: tc_python.precipitation.GrowthRateModel)

Sets the growth rate model for the class of precipitates. Default: GrowthRateModel.SIMPLIFIED

Parameters growth_rate_model_enum – The growth rate model

Returns This PrecipitatePhase object

with_particle_size_distribution (particle_size_distribution: tc_python.precipitation.ParticleSizeDistribution)

Sets the initial particle size distribution for this class of precipitates. Default: If the initial particle size distribution is not explicitly provided, the simulation will start from a supersaturated matrix.

Parameters particle_size_distribution – The initial particle size distribution object

Returns This PrecipitatePhase object

Tip: Use this option if you want to study the further evolution of an existing microstructure.

class tc_python.precipitation.PrecipitationCCTCalculation (calculation)

Bases: tc_python.precipitation.PrecipitationCalculation

Configuration for a Continuous-Cooling-Time (CCT) precipitation calculation.
calculate() → tc_python.precipitation.PrecipitationCalculationTTTorCCTResult

Runs the CCT-diagram calculation.

Returns A PrecipitationCalculationTTTorCCTResult which later can be used to get specific values from the calculated result

set_cooling_rates(cooling_rates: List[float])

Sets all cooling rates for which the CCT-diagram should be calculated.

Parameters cooling_rates – A list of cooling rates [K/s]

Returns This PrecipitationCCTCalculation object

set_max_temperature(max_temperature: float)

Sets maximum temperature of the CCT-diagram.

Parameters max_temperature – the maximum temperature [K]

Returns This PrecipitationCCTCalculation object

set_min_temperature(min_temperature: float)

Sets the minimum temperature of the CCT-diagram.

Parameters min_temperature – the minimum temperature [K]

Returns This PrecipitationCCTCalculation object

stop_at_volume_fraction_of_phase(stop_criterion_value: float)

Sets the stop criterion as a volume fraction of the phase. This setting is applied to all phases.

Parameters stop_criterion_value – the volume fraction of the phase (a value between 0 and 1)

Returns This PrecipitationCCTCalculation object

class tc_python.precipitation.PrecipitationCalculation(calculation)

Bases: tc_python.abstract_base.AbstractCalculation

Abstract base class for all precipitation calculations. Cannot be instantiated, use one of its subclasses instead.

set_composition(element_name: str, value: float)

Sets the composition of the elements. The unit for the composition can be changed using set_composition_unit(). Default: Mole percent (CompositionUnit.MOLE_PERCENT)

Parameters

• element_name – The element
• value – The composition (fraction or percent depending on the composition unit)

Returns This PrecipitationCalculation object

set_composition_unit(unit_enum: tc_python.utils.CompositionUnit)

Sets the composition unit. Default: Mole percent (CompositionUnit.MOLE_PERCENT).

Parameters unit_enum – The new composition unit

Returns This PrecipitationCalculation object

with_matrix_phase(matrix_phase: tc_python.precipitation.MatrixPhase)

Sets the matrix phase.

Parameters matrix_phase – The matrix phase

Returns This PrecipitationCalculation object
with_numerical_parameters(numerical_parameters: tc_python.precipitation.NumericalParameters)

Sets the numerical parameters. If not specified, reasonable defaults will be used.

Parameters numerical_parameters -- The parameters

Returns This PrecipitationCalculation object

class tc_python.precipitation.PrecipitationCalculationResult(result)
Bases: tc_python.abstract_base.AbstractResult

Result of a precipitation calculation. This can be used to query for specific values.

class tc_python.precipitation.PrecipitationCalculationSingleResult(result)
Bases: tc_python.precipitation.PrecipitationCalculationResult

Result of a isothermal or non-isothermal precipitation calculation. This can be used to query for specific values.
A detailed definition of the axis variables can be found in the Help.

get_aspect_ratio_distribution_for_particle_length_of(precipitate_id: str, time: float) → [typing.List[float], typing.List[float]]

Returns the aspect ratio distribution of a precipitate in dependency of its mean particle length at a certain time. Only available if the morphology is set to PrecipitateMorphology.NEEDLE or PrecipitateMorphology.PLATE.

Parameters

• time -- The time [s]

• precipitate_id -- The id of a precipitate can either be the phase name or an alias

Returns A tuple of two lists of floats (mean particle length [m], aspect ratio)

get_aspect_ratio_distribution_for_radius_of(precipitate_id: str, time: float) → [typing.List[float], typing.List[float]]

Returns the aspect ratio distribution of a precipitate in dependency of its mean radius at a certain time. Only available if the morphology is set to PrecipitateMorphology.NEEDLE or PrecipitateMorphology.PLATE.

Parameters

• time -- The time [s]

• precipitate_id -- The id of a precipitate can either be the phase name or an alias

Returns A tuple of two lists of floats (mean radius [m], aspect ratio)

get_critical_radius_of(precipitate_id: str) → [typing.List[float], typing.List[float]]

Returns the critical radius of a precipitate in dependency of the time.

Parameters precipitate_id -- The id of a precipitate can either be phase name or alias

Returns A tuple of two lists of floats (time [s], critical radius [m])

get_cubic_factor_distribution_for_particle_length_of(precipitate_id: str, time: float) → [typing.List[float], typing.List[float]]

Returns the cubic factor distribution of a precipitate in dependency of its mean particle length at a certain time. Only available if the morphology is set to PrecipitateMorphology.CUBOID.

Parameters

• time -- The time in seconds

• precipitate_id -- The id of a precipitate can either be the phase name or an alias
Returns A tuple of two lists of floats (particle length [m], cubic factor)

get_cubic_factor_distribution_for_radius_of(precipitate_id: str, time: float) → [typing.List[float], typing.List[float]]

Returns the cubic factor distribution of a precipitate in dependency of its mean radius at a certain time.
Only available if the morphology is set to PrecipitateMorphology.CUBOID.

Parameters

* time – The time [s]
* precipitate_id – The id of a precipitate can either be the phase name or an alias

Returns A tuple of two lists of floats (radius [m], cubic factor)

get_driving_force_of(precipitate_id: str) → [typing.List[float], typing.List[float]]

Returns the (by R * T) normalized driving force of a precipitate in dependency of the time.

Parameters precipitate_id – The id of a precipitate can either be the phase name or an alias

Returns A tuple of two lists of floats (time [s], normalized driving force)

get_matrix_composition_in_mole_fraction_of(element_name: str) → [typing.List[float], typing.List[float]]

Returns the matrix composition (as mole fractions) of a certain element in dependency of the time.

Parameters element_name – The element

Returns A tuple of two lists of floats (time [s], mole fraction)

get_matrix_composition_in_weight_fraction_of(element_name: str) → [typing.List[float], typing.List[float]]

Returns the matrix composition (as weight fraction) of a certain element in dependency of the time.

Parameters element_name – The element

Returns A tuple of two lists of floats (time [s], weight fraction)

get_mean_aspect_ratio_of(precipitate_id: str) → [typing.List[float], typing.List[float]]

Returns the mean aspect ratio of a precipitate in dependency of the time. Only available if the morphology
is set to PrecipitateMorphology.NEEDLE or PrecipitateMorphology.PLATE.

Parameters precipitate_id – The id of a precipitate can either be the phase name or an alias

Returns A tuple of two lists of floats (time [s], mean aspect ratio)

get_mean_cubic_factor_of(precipitate_id: str) → [typing.List[float], typing.List[float]]

Returns the mean cubic factor of a precipitate in dependency of the time. Only available if the morphology
is set to PrecipitateMorphology.CUBOID.

Parameters precipitate_id – The id of a precipitate can either be the phase name or an alias

Returns A tuple of two lists of floats (time [s], mean cubic factor)

get_mean_particle_length_of(precipitate_id: str) → [typing.List[float], typing.List[float]]

Returns the mean particle length of a precipitate in dependency of the time. Only available if the morphology
is set to PrecipitateMorphology.NEEDLE or PrecipitateMorphology.PLATE.

Parameters precipitate_id – The id of a precipitate can either be the phase name or an alias

Returns A tuple of two lists of floats (time [s], mean particle length [m])
get_mean_radius_of (precipitate_id: str) → [typing.List[float], typing.List[float]]

Returns the mean radius of a precipitate in dependency of the time.

Parameters precipitate_id – The id of a precipitate can either be phase name or alias

Returns A tuple of two lists of floats (time [s], mean radius [m])

get_nucleation_rate_of (precipitate_id: str) → [typing.List[float], typing.List[float]]

Returns the nucleation rate of a precipitate in dependency of the time.

Parameters precipitate_id – The id of a precipitate can either be the phase name or an alias

Returns A tuple of two lists of floats (time [s], nucleation rate [m^-3 s^-1])

get_number_density_of (precipitate_id: str) → [typing.List[float], typing.List[float]]

Returns the particle number density of a precipitate in dependency of the time.

Parameters precipitate_id – The id of a precipitate can either be the phase name or alias

Returns A tuple of two lists of floats (time [s], particle number density [m^-3])

get_size_distribution_for_particle_length_of (precipitate_id: str, time: float) → [typing.List[float], typing.List[float]]

Returns the size distribution of a precipitate in dependency of its mean particle length at a certain time.

Parameters

• time – The time [s]

• precipitate_id – The id of a precipitate can either be the phase name or an alias

Returns A tuple of two lists of floats (particle length[m], number of particles per unit length [m^-4])

get_size_distribution_for_radius_of (precipitate_id: str, time: float) → [typing.List[float], typing.List[float]]

Returns the size distribution of a precipitate in dependency of its mean radius at a certain time.

Parameters

• time – The time [s]

• precipitate_id – The id of a precipitate can either be the phase name or an alias

Returns A tuple of two lists of floats (radius [m], number of particles per unit volume per unit length [m^-4])

get_volume_fraction_of (precipitate_id: str) → [typing.List[float], typing.List[float]]

Returns the volume fraction of a precipitate in dependency of the time.

Parameters precipitate_id – The id of a precipitate can either be the phase name or alias

Returns A tuple of two lists of floats (time [s], volume fraction)

class tc_python.precipitation.PrecipitationCalculationTTTOrCCTResult (result)

Bases: tc_python.precipitation.PrecipitationCalculationResult

Result of a TTT or CCT precipitation calculation.

get_result_for_precipitate (precipitate_id: str) → [typing.List[float], typing.List[float]]

Returns the calculated data of a TTT or CCT diagram for a certain precipitate.

Parameters precipitate_id – The id of a precipitate can either be the phase name or an alias
Returns A tuple of two lists of floats (time [s], temp [K])

class tc_python.precipitation.PrecipitationIsoThermalCalculation(calculation)
   Bases: tc_python.precipitation.PrecipitationCalculation

Configuration for an isothermal precipitation calculation.

calculate() → tc_python.precipitation.PrecipitationCalculationSingleResult
   Runs the isothermal precipitation calculation.

   Returns A PrecipitationCalculationSingleResult which later can be used to get specific values
   from the calculated result

set_simulation_time(simulation_time: float)
   Sets the simulation time.

   Parameters simulation_time – The simulation time [s]

   Returns This PrecipitationIsoThermalCalculation object

set_temperature(temperature: float)
   Sets the temperature for the isothermal simulation.

   Parameters temperature – the temperature [K]

   Returns This PrecipitationIsoThermalCalculation object

class tc_python.precipitation.PrecipitationNonIsoThermalCalculation(calculation)
   Bases: tc_python.precipitation.PrecipitationCalculation

Configuration for a non-isothermal precipitation calculation.

calculate() → tc_python.precipitation.PrecipitationCalculationSingleResult
   Runs the non-isothermal precipitation calculation.

   Returns A PrecipitationCalculationSingleResult which later can be used to get specific values
   from the calculated result

set_simulation_time(simulation_time: float)
   Sets the simulation time.

   Parameters simulation_time – The simulation time [s]

   Returns This PrecipitationNonThermalCalculation object

with_temperature_profile(temperature_profile: tc_python.utils.TemperatureProfile)
   Sets the temperature profile to use with this calculation.

   Parameters temperature_profile – the temperature profile object (specifying time / tem-
   perature points)

   Returns This PrecipitationNonThermalCalculation object

class tc_python.precipitation.PrecipitationTTTCalculation(calculation)
   Bases: tc_python.precipitation.PrecipitationCalculation

Configuration for a TTT (Time-Temperature-Transformation) precipitation calculation.

calculate() → tc_python.precipitation.PrecipitationCalculationTTTorCCTResult
   Runs the TTT-diagram calculation.

   Returns A PrecipitationCalculationTTTorCCTResult which later can be used to get specific val-
   ues from the calculated result.

5.1. Calculations
set_max_annealing_time(max_annealing_time: float)
Sets the maximum annealing time, i.e. the maximum time of the simulation if the stopping criterion is not reached.

Parameters max_annealing_time – the maximum annealing time [s]

Returns This PrecipitationTTTCalculation object

set_max_temperature(max_temperature: float)
Sets the maximum temperature for the TTT-diagram.

Parameters max_temperature – the maximum temperature [K]

Returns This PrecipitationTTTCalculation object

set_min_temperature(min_temperature: float)
Sets the minimum temperature for the TTT-diagram.

Parameters min_temperature – the minimum temperature [K]

Returns This PrecipitationTTTCalculation object

set_temperature_step(temperature_step: float)
Sets the temperature step for the TTT-diagram, if unset the default value is 10 K.

Parameters temperature_step – the temperature step [K]

Returns This PrecipitationTTTCalculation object

stop_at_percent_of_equilibrium_fraction(percentage: float)
Sets the stop criterion to a percentage of the overall equilibrium phase fraction, alternatively a required volume fraction can be specified (using stop_at_volume_fraction_of_phase()).

Parameters percentage – the percentage to stop at (value between 0 and 100)

Returns This PrecipitationTTTCalculation object

stop_at_volume_fraction_of_phase(volume_fraction: float)
Sets the stop criterion as a volume fraction of the phase, alternatively a required percentage of the equilibrium phase fraction can be specified (using stop_at_percent_of_equilibrium_fraction()). Stopping at a specified volume fraction is the default setting.

This setting is applied to all phases.

Parameters volume_fraction – the volume fraction to stop at (a value between 0 and 1)

Returns This PrecipitationTTTCalculation object

class tc_python.precipitation.TransformationStrainCalculationOption
Bases: enum.Enum
Options for calculating the transformation strain.

CALCULATE_FROM_MOLAR_VOLUME = 2
Calculates the transformation strain from the molar volume, obtains a purely dilatational strain.

DISREGARD = 1
Ignores the transformation strain, this is the default setting.

USER_DEFINED = 3
Transformation strain to be specified by the user.

class tc_python.precipitation.VolumeFractionOfPhaseType
Bases: enum.Enum
Unit of the volume fraction of a phase.
VOLUME_FRACTION = 6
   Volume fraction (0 - 1), this is the default.

VOLUME_PERCENT = 5
   Volume percent (0% - 100%).

5.1.3 Module “scheil”

class tc_python.scheil.ScheilCalculation (calculator)
   Bases: tc_python.abstract_base.AbstractCalculation

   Configuration for a Scheil solidification calculation.

   Note: Specify the settings, the calculation is performed with calculate().

   calculate() → tc_python.scheil.ScheilCalculationResult
      Runs the Scheil calculation.

      Warning: Scheil calculations do not support the GAS phase being selected, this means the GAS phase must always be deselected in the system if it is present in the database

      Returns A ScheilCalculationResult which later can be used to get specific values from the simulation.

   disable_approximate_driving_force_for_metastable_phases ()
      Disables the approximation of the driving force for metastable phases.

      Default: Enabled

      Note: When enabled, the metastable phases are included in all iterations. However, these may not have reached their most favourable composition and thus their driving forces may be only approximate.

      If it is important that these driving forces are correct, use disable_approximate_driving_force_for_metastable_phases() to force the calculation to converge for the metastable phases.

      Returns This ScheilCalculation object

   disable_global_minimization ()
      Disables global minimization.

      Default: Disabled

      Note: When enabled, a global minimization test is performed when an equilibrium is reached. This costs more computer time but the calculations are more robust.

      Returns This ScheilCalculation object
**enable_approximate_driving_force_for_metastable_phases()**

Enables the approximation of the driving force for metastable phases.

**Default:** Enabled

**Note:** When enabled, the metastable phases are included in all iterations. However, these may not have reached their most favourable composition and thus their driving forces may be only approximate.

If it is important that these driving forces are correct, use `disable_approximate_driving_force_for_metastable_phases()` to force the calculation to converge for the metastable phases.

**Returns** This `ScheilCalculation` object

**enable_global_minimization()**

Enables global minimization.

**Default:** Disabled

**Note:** When enabled, a global minimization test is performed when an equilibrium is reached. This costs more computer time but the calculations are more robust.

**Returns** This `ScheilCalculation` object

**set_composition** *(component_name: str, value: float)*

Sets the composition of a component. The unit for the composition can be changed using `set_composition_unit()`.

**Default:** Mole percent (`CompositionUnit.MOLE_PERCENT`)

**Parameters**

• `component_name` – The component

• `value` – The composition value [composition unit defined for the calculation]

**Returns** This `ScheilCalculation` object

**set_composition_unit** *(unit_enum: tc_python.utils.CompositionUnit = <CompositionUnit.MOLE_PERCENT: 1>)*

Sets the composition unit.

**Default:** Mole percent (`CompositionUnit.MOLE_PERCENT`).

**Parameters**

• `unit_enum` – The new composition unit

**Returns** This `ScheilCalculation` object

**set_fast_diffusing_elements** *(elements: List[str]*)

Sets elements as fast diffusing. This allows redistribution of these elements in both the solid and liquid parts of the alloy.

**Default:** No fast-diffusing elements.

**Parameters**

• `elements` – The elements

**Returns** This `ScheilCalculation` object
**set_liquid_phase** *(phase_name: str = 'LIQUID')*

Sets the phase used as the liquid phase.

**Default:** The phase “LIQUID”.

**Parameters** phase_name – The phase name

**Returns** This *ScheilCalculation* object

**set_max_no_of_iterations** *(max_no_of_iterations: int = 500)*

Set the maximum number of iterations.

**Default:** max. 500 iterations

**Note:** As some models give computation times of more than 1 CPU second/iteration, this number is also used to check the CPU time and the calculation stops if 500 CPU seconds/iterations are used.

**Parameters** max_no_of_iterations – The max. number of iterations

**Returns** This *ScheilCalculation* object

**set_required_accuracy** *(accuracy: float = 1e-06)*

Sets the required relative accuracy.

**Default:** 1.0E-6

**Note:** This is a relative accuracy, and the program requires that the relative difference in each variable must be lower than this value before it has converged. A larger value normally means fewer iterations but less accurate solutions. The value should be at least one order of magnitude larger than the machine precision.

**Parameters** accuracy – The required relative accuracy

**Returns** This *ScheilCalculation* object

**set_smallest_fraction** *(smallest_fraction: float = 1e-12)*

Sets the smallest fraction for constituents that are unstable.

It is normally only in the gas phase that you can find such low fractions.

The **default value** for the smallest site-fractions is 1E-12 for all phases except for IDEAL phase with one sublattice site (such as the GAS mixture phase in many databases) for which the default value is always as 1E-30.

**Parameters** smallest_fraction – The smallest fraction for constituents that are unstable

**Returns** This *ScheilCalculation* object

**set_start_temperature** *(temperature_in_kelvin: float = 2500.0)*

Sets the start temperature.

**Warning:** The start temperature needs to be higher than the liquidus temperature of the alloy.

**Default:** 2500.0 K

**Parameters** temperature_in_kelvin – The temperature [K]
Returns This `ScheilCalculation` object

```plaintext
set_temperature_step (temperature_step_in_kelvin: float = 1.0)
```
Sets the temperature step. Decreasing the temperature step increases the accuracy, but the default value is usually adequate.

**Default step**: 1.0 K

**Parameters**
- `temperature_step_in_kelvin` – The temperature step [K]

**Returns** This `ScheilCalculation` object

```plaintext
terminate_on_fraction_of_liquid_phase (fraction_to_terminate_at: float = 0.01)
```
Sets the termination condition to a specified remaining fraction of liquid phase.

**Default**: Terminates at 0.01 fraction of liquid phase.

**Note**: Either the termination criterion is set to a temperature or fraction of liquid limit, both together are not possible.

**Parameters**
- `fraction_to_terminate_at` – the termination fraction of liquid phase (value between 0 and 1)

**Returns** This `ScheilCalculation` object

```plaintext
terminate_on_temperature (temperature_in_kelvin: float)
```
Sets the termination condition to a specified temperature.

**Default**: Terminates at 0.01 fraction of liquid phase, i.e. not at a specified temperature.

**Note**: Either the termination criterion is set to a temperature or fraction of liquid limit, both together are not possible.

**Parameters**
- `temperature_in_kelvin` – the termination temperature [K]

**Returns** This `ScheilCalculation` object

```plaintext
class tc_python.scheil.ScheilCalculationResult (result)
```
**Bases**: `tc_python.abstract_base.AbstractResult`

Result of a Scheil calculation.

```plaintext
get_values_grouped_by_quantity_of (x_quantity: Union[tc_python.quantity_factory.ScheilQuantity,
str], y_quantity: Union[tc_python.quantity_factory.ScheilQuantity,
str], sort_and_merge: bool = True) → Dict[str, tc_python.utils.ResultValueGroup]
```
Returns x-y-line data grouped by the multiple datasets of the specified quantities (for example in dependency of phases or components). Use `get_values_of()` instead if you need no separation. The available quantities can be found in the documentation of the factory class `ScheilQuantity`.

**Note**: The different datasets might contain `NaN`-values between different subsections and might not be sorted **even if the flag ‘sort_and_merge‘ has been set** (because they might be unsortable due to their nature).

**Parameters**
• **x_quantity** – The first Scheil quantity (“x-axis”), console syntax strings can be used as an alternative (for example “T”)

• **y_quantity** – The second Scheil quantity (“y-axis”), console syntax strings can be used as an alternative (for example “NV”)

• **sort_and_merge** – If True, the data will be sorted and merged into as few subsections as possible (divided by NaN)

**Returns** Dict containing the `ResultValueGroup` dataset objects with their quantity labels as keys

```python
get_values_grouped_by_stable_phases_of(x_quantity: Union[tc_python.quantity_factory.ScheilQuantity, str],
                                y_quantity: Union[tc_python.quantity_factory.ScheilQuantity, str],
                                sort_and_merge: bool = True) → Dict[str, tc_python.utils.ResultValueGroup]
```

Returns x-y-line data grouped by the sets of “stable phases” (for example “LIQUID” or “LIQUID + FCC_A1”). Use `get_values_of()` instead if you need no separation. The available quantities can be found in the documentation of the factory class `ScheilQuantity`.

**Note:** The different datasets might contain NaN-values between different subsections and might not be sorted even if the flag ‘sort_and_merge’ has been set (because they might be unsortable due to their nature).

**Parameters**

• **x_quantity** – The first Scheil quantity (“x-axis”), console syntax strings can be used as an alternative (for example “T”)

• **y_quantity** – The second Scheil quantity (“y-axis”), console syntax strings can be used as an alternative (for example “NV”)

• **sort_and_merge** – If True, the data will be sorted and merged into as few subsections as possible (divided by NaN)

**Returns** Dict containing the `ResultValueGroup` dataset objects with their “stable phases” labels as keys

```python
get_values_of(x_quantity: Union[tc_python.quantity_factory.ScheilQuantity, str],
             y_quantity: Union[tc_python.quantity_factory.ScheilQuantity, str]) → [typing.List[float], typing.List[float]]
```

Returns sorted x-y-line data without any separation. Use `get_values_grouped_by_quantity_of()` or `get_values_grouped_by_stable_phases_of()` instead if you need such a separation. The available quantities can be found in the documentation of the factory class `ScheilQuantity`.

**Note:** This method will always return sorted data without any NaN-values. In case of ambiguous quantities (for example: `CompositionOfPhaseAsWeightFraction("FCC_A1", "All")`) that can give data that is hard to interpret. In such a case you need to choose the quantity in another way or use one of the other methods.

**Parameters**

• **x_quantity** – The first Scheil quantity (“x-axis”), console syntax strings can be used as an alternative (for example “T”)

---

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• **y_quantity** – The second Scheil quantity (“y-axis”), console syntax strings can be used as an alternative (for example “NV”)

**Returns**  A tuple containing the x- and y-data in lists

### 5.1.4 Module “step_or_map_diagrams”

```python
class tc_python.step_or_map_diagrams.AbstractAxisType
    Bases: object
    The abstract base class for all axis types.
    
    get_type() → str
        Convenience method for getting the axis type.
        
        **Returns** The type

class tc_python.step_or_map_diagrams.AxisType
    Bases: tc_python.step_or_map_diagrams.AbstractAxisType
    Factory class providing objects for configuring a logarithmic or linear axis by using `AxisType.linear()` or `AxisType.logarithmic()`.

    @classmethod
    linear()
        Creates an object for configuring a linear calculation axis.
        
        **Default:** A minimum number of 40 steps.

    **Note:** The returned object can be configured regarding the maximum step size or the minimum number of steps on the axis.

    **Returns** A new `Linear` object

    @classmethod
    logarithmic()
        Creates an object for configuring a logarithmic calculation axis.
        
        **Default:** A scale factor of 1.1

    **Note:** The returned object can be configured regarding the scale factor.

    **Returns** A new `Logarithmic` object
```

```python
class tc_python.step_or_map_diagrams.CalculationAxis(quantity: Union[tc_python.quantity_factory.ThermodynamicQuantity, str])
    Bases: object
    A calculation axis used for property and phase diagram calculations.

    **Note:** A calculation axis is defining the varied condition and the range of variation. It is the same concept as in Thermo-Calc Graphical Mode or Console Mode.

    **Default:** A `Linear` axis with a minimum number of 40 steps
```
**set_max**(max: float)
Sets the maximum quantity value of the calculation axis.

There is no default value set, it always needs to be defined.

Parameters max – The maximum quantity value of the axis [unit according to the axis quantity]

Returns This CalculationAxis object

**set_min**(min: float)
Sets the minimum quantity value of the calculation axis.

There is no default value set, it always needs to be defined.

Parameters min – The minimum quantity value of the axis [unit according to the axis quantity]

Returns This CalculationAxis object

**set_start_at**(at: float)
Sets the starting point of the calculation on the axis.

Default: The default starting point is the center between the minimum and maximum quantity value

Parameters at – The starting point on the axis [unit according to the axis quantity]

Returns This CalculationAxis object

**with_axis_type**(axis_type: tc_python.step_or_map_diagrams.AxisType)
Sets the type of the axis.

Default: A Linear axis with a minimum number of 40 steps

Parameters axis_type – The axis type (linear or logarithmic)

Returns This CalculationAxis object

---

**Linear**

Bases: tc_python.step_or_map_diagrams.AbstractAxisType

Represents a linear axis.

**get_type**() ➔ str
Convenience method for getting axis type.

Returns The type

**set_max_step_size**(max_step_size: float)
Sets the axis to use the maximum step size configuration.

Default: This is not the default which is minimum number of steps

Note: Either maximum step size or minimum number of steps can be used but not both at the same time.

Parameters max_step_size – The maximum step size [unit according to the axis quantity]

Returns This Linear object

**set_min_nr_of_steps**(min_nr_of_steps: float = 40)
Sets the axis to use the minimum number of steps configuration.

Default: This is the default option (with a minimum number of steps of 40)

---

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Note: Either maximum step size or minimum number of steps can be used but not both at the same time.

Parameters `min_nr_of_steps` – The minimum number of steps

Returns This Linear object

class tc_python.step_or_map_diagrams.Logarithmic(scale_factor: float = 1.1)
Bases: tc_python.step_or_map_diagrams.AbstractAxisType

Represents a logarithmic axis.

Note: A logarithmic axis is useful for low fractions like in a gas phase where 1E-7 to 1E-2 might be an interesting range. For the pressure a logarithmic axis is often also useful.

def get_type() → str
    Convenience method for getting axis type.
    Returns The type

def set_scale_factor(scale_factor: float = 1.1)
    Sets the scale factor.
    Default: 1.1
    Parameters scale_factor – The scale factor setting the maximum factor between two calculated values, must be larger than 1.0
    Returns This Logarithmic object

class tc_python.step_or_map_diagrams.PhaseDiagramCalculation(calculator)
Bases: tc_python.step_or_map_diagrams.ThermodynamicCalculation

Configuration for a phase diagram calculation.

Note: Specify the conditions, the calculation is performed with `calculate()`.

def calculate(keep_previous_results: bool = False) → tc_python.step_or_map_diagrams.PhaseDiagramResult
    Performs the phase diagram calculation.

    Warning: If you use `keep_previous_results=True`, you must not use another calculator or even get results in between the calculations using `calculate()`. Then the previous results will actually be lost.

    Parameters keep_previous_results – If True, results from any previous call to this method are appended. This can be used to combine calculations with multiple start points if the mapping fails at a certain condition.
    Returns A new PhaseDiagramResult object which later can be used to get specific values from the calculated result.

with_first_axis(axis: tc_python.step_or_map_diagrams.CalculationAxis)
    Sets the first calculation axis.

    Parameters axis – The axis
Returns This \texttt{PhaseDiagramCalculation} object

\textbf{with\_second\_axis} (axis: \texttt{tc\_python.step\_or\_map\_diagrams.CalculationAxis})

Sets the second calculation axis.

\textbf{Parameters} \texttt{axis} – The axis

\textbf{Returns} This \texttt{PhaseDiagramCalculation} object

\texttt{class tc\_python.step\_or\_map\_diagrams.PhaseDiagramResult} (result)

Bases: \texttt{tc\_python.abstract\_base.AbstractResult}

Result of a phase diagram calculation, it can be evaluated using quantities or Console Mode syntax.

\textbf{add\_coordinate\_for\_phase\_label} (x: float, y: float)

Sets a coordinate in the result plot for which the stable phases will be evaluated and provided in the result data object. This can be used to plot the phases of a region into the phase diagram or just to programmatically evaluate the phases in certain regions.

\textbf{Warning:} This method takes coordinates of the \texttt{plot} axes and not of the calculation axis.

\textbf{Parameters}

- \texttt{x} – The coordinate of the first \texttt{plot} axis (“x-axis”) [unit of the \texttt{plot} axis]
- \texttt{y} – The coordinate of the second \texttt{plot} axis (“y-axis”) [unit of the \texttt{plot} axis]

\textbf{Returns} This \texttt{PhaseDiagramResult} object

\textbf{get\_values\_grouped\_by\_quantity\_of} (x\_quantity: \texttt{Union[tc\_python.quantity\_factory.ThermodynamicQuantity, str]}, y\_quantity: \texttt{Union[tc\_python.quantity\_factory.ThermodynamicQuantity, str]}) \rightarrow \texttt{tc\_python.step\_or\_map\_diagrams.PhaseDiagramResultValues}

Returns x-y-line data grouped by the multiple datasets of the specified quantities (for example in dependency of components). The available quantities can be found in the documentation of the factory class \texttt{ThermodynamicQuantity}. Usually the result data represents the phase diagram.

\textbf{Note:} The different datasets will contain NaN-values between different subsections and will not be sorted (because they are unsortable due to their nature).

\textbf{Parameters}

- \texttt{x\_quantity} – The first quantity (“x-axis”), console syntax strings can be used as an alternative (for example “T”)
- \texttt{y\_quantity} – The second quantity (“y-axis”), console syntax strings can be used as an alternative (for example “NV”)

\textbf{Returns} The phase diagram data

\textbf{get\_values\_grouped\_by\_stable\_phases\_of} (x\_quantity: \texttt{Union[tc\_python.quantity\_factory.ThermodynamicQuantity, str]}, y\_quantity: \texttt{Union[tc\_python.quantity\_factory.ThermodynamicQuantity, str]}) \rightarrow \texttt{tc\_python.step\_or\_map\_diagrams.PhaseDiagramResultValues}

Returns x-y-line data grouped by the sets of “stable phases” (for example “LIQUID” or “LIQUID + FCC\_A1”). The available quantities can be found in the documentation of the factory class \texttt{ThermodynamicQuantity}. Usually the result data represents the phase diagram.
Note: The different datasets will contain NaN-values between different subsections and will not be sorted (because they are unsortable due to their nature).

Parameters

- **x_quantity** – The first quantity ("x-axis"), console syntax strings can be used as an alternative (for example "T")
- **y_quantity** – The second quantity ("y-axis"), console syntax strings can be used as an alternative (for example "NV")

Returns The phase diagram data

**remove_phase_labels()**

Erases all added coordinates for phase labels.

Returns This PhaseDiagramResult object

**set_phase_name_style**(phase_name_style_enum: tc_python.step_or_map_diagrams.PhaseNameStyle = <PhaseNameStyle.NONE: 0>)

Sets the style of the phase name labels that will be used in the result data object (constitution description, ordering description, ...).

Default: PhaseNameStyle.NONE

Parameters **phase_name_style_enum** – The phase name style

Returns This PhaseDiagramResult object

class tc_python.step_or_map_diagrams.PhaseDiagramResultValues(phase_diagram_values_java)

Bases: object

Represents the data of a phase diagram.

**get_invariants()** \(\rightarrow\) tc_python.utils.ResultSetValueGroup

Returns the x- and y-datasets of all invariants in the phase diagram.

Note: The datasets will normally contain different sections separated by NaN-values.

Returns The invariants dataset object

**get_lines()** \(\rightarrow\) Dict[str, tc_python.utils.ResultSetValueGroup]

Returns the x- and y-datasets of all phase boundaries in the phase diagram.

Note: The datasets will normally contain different sections separated by NaN-values.

Returns Dict containing the phase boundary datasets with the quantities or stable phases as keys (depending on the used method to get the values)

**get_phase_labels()** \(\rightarrow\) List[tc_python.step_or_map_diagrams.PhaseLabel]

Returns the phase labels added for certain coordinates using PhaseDiagramResult.add_coordinate_for_phase_label().

Returns The list with the phase label data (that contains plot coordinates and stable phases)
get_tie_lines() → tc_python.utils.ResultValueGroup

Returns the x- and y-datasets of all tie-lines in the phase diagram.

Note: The datasets will normally contain different sections separated by NaN-values.

Returns The tie-line dataset object

class tc_python.step_or_map_diagrams.PhaseLabel (x: float, y: float, text: str)
    Bases: object

    Represents a phase label at a plot coordinate, i.e. the stable phases that are present at that plot coordinate.

    Variables
    • x – The coordinate of the first plot axis (“x-axis”) [unit of the plot axis]
    • y – The coordinate of the second plot axis (“y-axis”) [unit of the plot axis]
    • text – The label (i.e. the stable phases at that point in the phase diagram, for example “LIQUID + FCC_A1”)

class tc_python.step_or_map_diagrams.PhaseNameStyle
    Bases: enum.Enum

    The style of the phase names used in the labels.

    ALL = 1
    Adding ordering and constitution description.

    CONSTITUTION_DESCRIPTION = 3
    Adding only constitution description.

    NONE = 0
    Only the phase names.

    ORDERING_DESCRIPTION = 4
    Adding only ordering description.

class tc_python.step_or_map_diagrams.PropertyDiagramCalculation (calculator)
    Bases: tc_python.step_or_map_diagrams.ThermodynamicCalculation

    Configuration for a property diagram calculation.

    Note: Specify the conditions, the calculation is performed with calculate().

    calculate (keep_previous_results: bool = False) → tc_python.step_or_map_diagrams.PropertyDiagramResult
    Performs the property diagram calculation.

    Warning: If you use keep_previous_results=True, you must not use another calculator or even get results in between the calculations using calculate(). Then the previous results will actually be lost.

    Parameters keep_previous_results – If True, results from any previous call to this method are appended. This can be used to combine calculations with multiple start points if the stepping fails at a certain condition.

    Returns A new PropertyDiagramResult object which later can be used to get specific values from the calculated result

5.1. Calculations
disable_step_separate_phases()
Disables step separate phases. This is the default setting.

Returns This PropertyDiagramCalculation object

enable_step_separate_phases()
Enables step separate phases.

Default: By default separate phase stepping is disabled

Note: This is an advanced option, it is used mostly to calculate how the Gibbs energy for a number of phases varies for different compositions. This is particularly useful to calculate Gibbs energies for complex phases with miscibility gaps and for an ordered phase that is never disordered (e.g. SIGMA-phase, G-phase, MU-phase, etc.).

Returns This PropertyDiagramCalculation object

with_axis(axis: tc_python.step_or_map_diagrams.CalculationAxis)
Sets the calculation axis.

Parameters axis – The axis

Returns This PropertyDiagramCalculation object

class tc_python.step_or_map_diagrams.PropertyDiagramResult(result)
Bases: tc_python.abstract_base.AbstractResult
Result of a property diagram. This can be used to query for specific values.

get_values_grouped_by_quantity_of(x_quantity: Union[tc_python.quantity_factory.ThermodynamicQuantity, str], y_quantity: Union[tc_python.quantity_factory.ThermodynamicQuantity, str], sort_and_merge: bool = True) → Dict[str, tc_python.utils.ResultValueGroup]
Returns x-y-line data grouped by the multiple datasets of the specified quantities (typically the phases). The available quantities can be found in the documentation of the factory class ThermodynamicQuantity.

Note: The different datasets might contain NaN-values between different subsections and might not be sorted even if the flag 'sort_and_merge' has been set (because they might be unsortable due to their nature).

Parameters
• x_quantity – The first quantity (“x-axis”), console syntax strings can be used as an alternative (for example “T”)
• y_quantity – The second quantity (“y-axis”), console syntax strings can be used as an alternative (for example “NV”)
• sort_and_merge – If True, the data will be sorted and merged into as few subsections as possible (divided by NaN)

Returns Dict containing the datasets with the quantities as their keys
get_values_grouped_by_stable_phases_of(x_quantity: Union[tc_python.quantity_factory.ThermodynamicQuantity, str], y_quantity: Union[tc_python.quantity_factory.ThermodynamicQuantity, str], sort_and_merge: bool = True) → Dict[str, tc_python.utils.ResultValueGroup]

Returns x-y-line data grouped by the sets of “stable phases” (for example “LIQUID” or “LIQUID + FCC_A1”). The available quantities can be found in the documentation of the factory class ThermodynamicQuantity.

Note: The different datasets might contain NaN-values between different subsections and different lines of an ambiguous dataset. They might not be sorted even if the flag ‘sort_and_merge’ has been set (because they might be unsortable due to their nature).

Parameters

• x_quantity – The first quantity (“x-axis”), console syntax strings can be used as an alternative (for example “T”)

• y_quantity – The second quantity (“y-axis”), console syntax strings can be used as an alternative (for example “NV”)

• sort_and_merge – If True, the data will be sorted and merged into as few subsections as possible (divided by NaN)

Returns Dict containing the datasets with the quantities as their keys

get_values_of(x_quantity: Union[tc_python.quantity_factory.ThermodynamicQuantity, str], y_quantity: Union[tc_python.quantity_factory.ThermodynamicQuantity, str]) → [typing.List[float], typing.List[float]]

Returns sorted x-y-line data without any separation. Use get_values_grouped_by_quantity_of() or get_values_grouped_by_stable_phases_of() instead if you need such a separation. The available quantities can be found in the documentation of the factory class ThermodynamicQuantity.

Note: This method will always return sorted data without any NaN-values. If it is unsortable that might give data that is hard to interpret. In such a case you need to choose the quantity in another way or use one of the other methods. One example of this is to use quantities with All-markers, for example MassFractionOfAComponent(‘All’).

Parameters

• x_quantity – The first Thermodynamic quantity (“x-axis”), console syntax strings can be used as an alternative (for example “T”)

• y_quantity – The second Thermodynamic quantity (“y-axis”), console syntax strings can be used as an alternative (for example “NV”)

Returns A tuple containing the x- and y-data in lists

set_phase_name_style(phase_name_style_enum: tc_python.step_or_map_diagrams.PhaseNameStyle = <PhaseNameStyle.NONE: 0>)

Sets the style of the phase name labels that will be used in the result data object (constitution description, ordering description, . . . ).

Default: PhaseNameStyle.NONE

Parameters phase_name_style_enum – The phase name style
Returns This PropertyDiagramResult object

class tc_python.step_or_map_diagrams.ThermodynamicCalculation(calculator)
Bases: tc_python.abstract_base.AbstractCalculation

Contains functionality that is common for phase diagram as well as property diagram calculations.

disable_global_minimization()
  Disables global minimization.
  Default: Enabled
  Returns This ThermodynamicCalculation object

enable_global_minimization()
  Enables global minimization.
  Default: Enabled
  Returns This ThermodynamicCalculation object

get_components() → List[str]
  Returns a list of the components.
  Returns The components

remove_all_conditions()
  Removes all set conditions.
  Returns This ThermodynamicCalculation object

remove_condition(quantity: Union[tc_python.quantity_factory.ThermodynamicQuantity, str])
  Removes the specified condition.
  Parameters quantity – The ThermodynamicQuantity to set as condition, a console syntax strings can be used as an alternative (for example X(Cr))
  Returns This ThermodynamicCalculation object

run_poly_command(command: str)
  Runs a Thermo-Calc command from the console POLY-module immediately in the engine.
  Parameters command – The Thermo-Calc console command
  Returns This ThermodynamicCalculation object

Note: It should not be necessary for most users to use this method, try to use the corresponding method implemented in the API instead.

Warning: As this method runs raw Thermo-Calc commands directly in the engine, it may hang the program in case of spelling mistakes (e.g. forgotten equals sign).

set_component_to_entered(component: str)
  Sets the specified component to the status ENTERED, that is the default state.
  Parameters component – The component name
  Returns This ThermodynamicCalculation object

set_component_to_suspended(component: str)
  Sets the specified component to the status SUSPENDED, i.e. it is ignored in the calculation.
Parameters `component` – The component name

Returns This `ThermodynamicCalculation` object

### set_condition

```python
set_condition(quantity: Union[tc_python.quantity_factory.ThermodynamicQuantity, str], value: float)
```

Sets the specified condition.

**Parameters**

- `quantity` – The ThermodynamicQuantity to set as condition, a console syntax string can be used as an alternative (for example `X(Cr)`)
- `value` – The value of the condition

**Returns** This `ThermodynamicCalculation` object

### set_phase_to_dormant

```python
set_phase_to_dormant(phase: str)
```

Sets the phase to the status DORMANT, necessary for calculating the driving force to form the specified phase.

**Parameters**

- `phase` – The phase name

**Returns** This `ThermodynamicCalculation` object

### set_phase_to_entered

```python
set_phase_to_entered(phase: str, amount: float)
```

Sets the phase to the status ENTERED, that is the default state.

**Parameters**

- `phase` – The phase name
- `amount` – The phase fraction (between 0.0 and 1.0)

**Returns** This `ThermodynamicCalculation` object

### set_phase_to_fixed

```python
set_phase_to_fixed(phase: str, amount: float)
```

Sets the phase to the status FIXED, i.e. it is guaranteed to have the specified phase fraction after the calculation.

**Parameters**

- `phase` – The phase name
- `amount` – The fixed phase fraction (between 0.0 and 1.0)

**Returns** This `ThermodynamicCalculation` object

### set_phase_to_suspended

```python
set_phase_to_suspended(phase: str)
```

Sets the phase to the status SUSPENDED, i.e. it is ignored in the calculation.

**Parameters**

- `phase` – The phase name

**Returns** This `ThermodynamicCalculation` object

## 5.2 Module “system”

### tc_python.system.MultiDatabaseSystemBuilder

```python
class tc_python.system.MultiDatabaseSystemBuilder(multi_database_system_builder)
```

**Bases:** `object`

Used to select databases, elements, phases etc. and create a System object. The difference to the class SystemBuilder is that the operations are performed on all the previously selected databases. The system is then used to create calculations.

### 5.2. Module “system”
deselect_phase(phase_name_to_deselect: str)
Deselects a phase for both the thermodynamic and the kinetic database.

Parameters phase_name_to_deselect – The phase name

Returns This MultiDatabaseSystemBuilder object

get_system() → tc_python.system.System
Creates a new System object that is the basis for all calculation types. Several calculation types can be
defined later from the object, they will be independent.

Returns A new System object

select_phase(phase_name_to_select: str)
Selects a phase for both the thermodynamic and the kinetic database.

Parameters phase_name_to_select – The phase name

Returns This MultiDatabaseSystemBuilder object

without_default_phases()
Removes all the default phases from both the thermodynamic and the kinetic database, any phase now
needs to be selected manually for the databases.

Returns This MultiDatabaseSystemBuilder object

class tc_python.system.System(system_instance)
Bases: object

A system containing selections for databases, elements, phases etc.

Note: For the defined system, different calculations can be configured and run. Instances of this class should
always be created from a SystemBuilder.

Note: The system object is immutable, i.e. it cannot be changed after is has been created. If you want to
change the system, you must instead create a new one.

get_all_phases_in_databases() → List[str]
Returns all phases present in the selected databases, regardless on selected elements, phases etc.

Returns a list of phases

get_phases_in_system() → List[str]
Returns all phases present in the system due to its configuration (selected elements, phases, etc.).

Returns a list of phases

with_cct_precipitation_calculation() → tc_python.precipitation.PrecipitationCCTCalculation
Creates a CCT-diagram calculation.

Returns A new PrecipitationCCTCalculation object

with_isothermal_precipitation_calculation() → tc_python.precipitation.PrecipitationIsoThermalCalculation
Creates an isothermal precipitation calculation.

Returns A new PrecipitationIsoThermalCalculation object

with_non_isothermal_precipitation_calculation()
Creates a non-isothermal precipitation calculation.

Returns A new PrecipitationNonIsoThermalCalculation object
with_phase_diagram_calculation (default_conditions: bool = True, components: List[str] = []) →
tc_python.step_or_map_diagrams.PhaseDiagramCalculation

Creates a phase diagram (map) calculation.

Parameters

• default_conditions – If True, automatically sets the conditions \( N=1 \) and \( P=100000 \)

• components – Specify here the components of the system (for example: [AL2O3, ...]),
  only necessary if they differ from the elements. If this option is used, all elements of the
  system need to be replaced by a component.

Returns A new PhaseDiagramCalculation object

with_property_diagram_calculation (default_conditions: bool = True, components: List[str] = []) →
tc_python.step_or_map_diagrams.PropertyDiagramCalculation

Creates a property diagram (step) calculation.

Parameters

• default_conditions – If True, automatically sets the conditions \( N=1 \) and \( P=100000 \)

• components – Specify here the components of the system (for example: [AL2O3, ...]),
  only necessary if they differ from the elements. If this option is used, all elements of the
  system need to be replaced by a component.

Returns A new PropertyDiagramCalculation object

with_scheil_calculation () → tc_python.scheil.ScheilCalculation

Creates a Scheil solidification calculation.

Warning: Scheil calculations do not support the GAS phase being selected, this means the ‘GAS’
phase must always be deselected in the system if it is present in the database

Returns A new ScheilCalculation object

with_single_equilibrium_calculation (default_conditions: bool = True, components: List[str] = []) →
tc_python.single_equilibrium.SingleEquilibriumCalculation

Creates a single equilibrium calculation.

Parameters

• default_conditions – If True, automatically sets the conditions \( N=1 \) and \( P=100000 \)

• components – Specify here the components of the system (for example: [AL2O3, ...]),
  only necessary if they differ from the elements. If this option is used, all elements of the
  system need to be replaced by a component.

Returns A new SingleEquilibriumCalculation object

with_ttt_precipitation_calculation () → tc_python.precipitation.PrecipitationTTTCalculation

Creates a TTT-diagram calculation.

Returns A new PrecipitationTTTCalculation object
class tc_python.system.SystemBuilder(system_builder)
    Bases: object

    Used to select databases, elements, phases etc. and create a System object. The system is then used to create calculations.

deselect_phase(phase_name_to_deselect: str)
    Deselects a phase in the last specified database only.

    Parameters
    -----
    phase_name_to_deselect -- The name of the phase

    Returns
    -------
    This SystemBuilder object

get_system() → tc_python.system.System
    Creates a new System object that is the basis for all calculation types. Several calculation types can be defined later from the object, they will be independent.

    Returns
    -------
    A new System object

get_system_for_scheil_calculations() → tc_python.system.System
    Creates a new System object without gas phases being selected, that is the basis for all calculation types, but its particularly useful for Scheil solidification calculations, where the model does not allow that a gas phase is selected in the system. Several calculation types can be defined later from the object, they will be independent.

    Returns
    -------
    A new System object

select_database_and_elements(database_name: str, list_of_element_strings: List[str])
    Selects thermodynamic or kinetic database and its selected elements (that will be appended). After that, phases can be selected or unselected.

    Parameters
    -----
    • database_name -- The database name, for example "FEDEMO"
    • list_of_element_strings -- A list of one or more elements as strings, for example ["Fe", "C"]

    Returns
    -------
    This SystemBuilder object

select_phase(phase_name_to_select: str)
    Selects a phase in the last specified database only.

    Parameters
    -----
    phase_name_to_select -- The name of the phase

    Returns
    -------
    This SystemBuilder object

select_user_database_and_elements(path_to_user_database: str, list_of_element_strings: List[str])
    Selects a thermodynamic database which is a user-defined database and select its elements (that will be appended).

    Parameters
    -----
    • path_to_user_database -- The path to the database file (*.TDB), defaults to the current working directory. Only the filename is required if the database is located in the same folder as the Python script.
    • list_of_element_strings -- A list of one or more elements as strings, for example ["Fe", "C"]

    Returns
    -------
    This SystemBuilder object
without_default_phases()  
Deselects all default phases in the last specified database only, any phase needs now to be selected manually for that database.

Returns This SystemBuilder object

5.3 Module “server”

class tc_python.server.SetUp(debug_logging=False)  
Bases: object  
Starting point for all calculations.  

Note: This class exposes methods that have no precondition, it is used for choosing databases and elements.

select_database_and_elements(database_name: str, list_of_elements: List[str]) → tc_python.system.SystemBuilder  
Selects a first thermodynamic or kinetic database and selects the elements in it.

Parameters  
• database_name – The name of the database, for example “FEDEMO”  
• list_of_elements – The list of the selected elements in that database, for example [“Fe”, “C”]

Returns A new SystemBuilder object

select_thermodynamic_and_kinetic_databases_with_elements(thermodynamic_db_name: str, kinetic_db_name: str, list_of_elements: List[str]) → tc_python.system.MultiDatabaseSystemBuilder  
Selects the thermodynamic and kinetic database at once, guarantees that the databases are added in the correct order. Further rejection or selection of phases applies to both databases.

Parameters  
• thermodynamic_db_name – The thermodynamic database name, for example “FEDEMO”  
• kinetic_db_name – The kinetic database name, for example “MFEDEMO”  
• list_of_elements – The list of the selected elements in that database, for example [“Fe”, “C”]

Returns A new MultiDatabaseSystemBuilder object

select_user_database_and_elements(path_to_user_database: str, list_of_elements: List[str]) → tc_python.system.SystemBuilder  
Selects a user defined database and selects the elements in it.

Parameters  
• path_to_user_database – The path to the database file (*.TDB), defaults to the current working directory. Only filename is required if the database is located in the same folder as the Python script.
• **list_of_elements** – The list of the selected elements in that database, for example
  ```
  ["Fe", "C"]
  ```

  **Returns**  A new `SystemBuilder` object

  **set_log_level_to_debug**()
  Sets log level to DEBUG

  **Returns**  This SetUp object

  **set_log_level_to_info**()
  Sets log level to INFO

  **Returns**  This SetUp object

**class** `tc_python.server.TCPython(debug_mode=False, debug_logging=False)`

**Bases:** `object`

**Starting point of the API. Typical syntax:**

```python
with TCPython() as session:
    session.select_database_and_elements(...)  
```

**Warning:** You should not run `TCPython()` more than once within one process (for example within a loop). Otherwise you will have for each call an open Java backend engine process that will never be closed. This behaviour does not affect multi-processing (for parallelization). That limitation might change in a future release.

Instead you should pass the session or calculator object into the loop and use them there.

**tc_python.server.start_api_server(debug_mode=False, is_unittest=False)**

**Starts a process of the API server and sets up the socket communication with it.**

**Parameters**

• **debug_mode** – If True it is tried to open a connection to an already running API-server. This is only used for debugging the API itself.

• **is_unittest** – Should be True if called by a unit test, only to be used internally for development.

**Warning:** Most users should use `TCPython` using a with-statement for automatic management of the resources (network sockets and temporary files). If you anyway need to use that method, make sure to call `stop_api_server()` in any case using the try-finally-pattern.

**tc_python.server.stop_api_server()**

**Clears all resources used by the session (i.e. shuts down the API server and deletes all temporary files). The disk usage of temporary files might be significant.**

**Warning:** Call this method only if you used `start_api_server()` initially. It should never be called when the API has been initialized in a with-statement using `TCPython`.  

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**Chapter 5. API Reference**

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5.4 Module “quantity_factory”

class tc_python.quantity_factory.ScheilQuantity
   Bases: tc_python.quantity.AbstractQuantity

Factory class providing quantities used for defining a Scheil calculation result (tc_python.scheil.ScheilCalculationResult).

classmethod apparent_heat_capacity_per_gram()
   Creates a quantity representing the apparent heat capacity [J/g/K].
   Returns A new ApparentHeatCapacityPerGram object.

classmethod apparent_heat_capacity_per_mole()
   Creates a quantity representing the apparent heat capacity [J/mol/K].
   Returns A new ApparentHeatCapacityPerMole object.

classmethod apparent_volumetric_thermal_expansion_coefficient()
   Creates a quantity representing the apparent volumetric thermal expansion coefficient of the system [1/K].
   Returns A new ApparentVolumetricThermalExpansionCoefficient object.

classmethod composition_of_phase_as_mole_fraction(phase: str = 'All', component: str = 'All')
   Creates a quantity representing the composition of a phase [mole-fraction].
   Parameters
   • phase – The name of the phase, use All to choose all stable phases
   • component – The name of the component, use All to choose all components
   Returns A new CompositionOfPhaseAsMoleFraction object.

classmethod composition_of_phase_as_weight_fraction(phase: str = 'All', component: str = 'All')
   Creates a quantity representing the composition of a phase [weight-fraction].
   Parameters
   • phase – The name of the phase, use All to choose all stable phases
   • component – The name of the component, use All to choose all components
   Returns A new CompositionOfPhaseAsWeightFraction object.

classmethod density_of_solid_phase(phase: str = 'All')
   Creates a quantity representing the average density of a solid phase [g/cm^3].
   Parameters phase – The name of the phase or All to choose all solid phases
   Returns A new DensityOfSolidPhase object.

classmethod density_of_system()
   Creates a quantity representing the average density of the system [g/cm^3].
   Returns A new DensityOfSystem object.

classmethod distribution_of_component_of_phase(phase: str, component: str)
   Creates a quantity representing the (molar) fraction of the specified component being present in the specified phase compared to the overall system [-]. This corresponds to the degree of segregation to that phase.
   Parameters
   • phase – The name of the phase
• **component** – The name of the component

**Returns** A new `DistributionOfComponentOfPhase` object.

```python
classmethod heat_per_gram()
```

Creates a quantity representing the total heat release from the liquidus temperature down to the current temperature [J/g].

**Note:** The total or apparent heat release during the solidification process consists of two parts: one is the so-called latent heat, i.e. heat due to the liquid -> solid phase transformation (`latent_heat_per_mole()` and `latent_heat_per_gram()`), and the other is the heat related to the specific heat of liquid and solid phases (`heat_per_mole()` and `heat_per_gram()`).

**Returns** A new `HeatPerGram` object.

```python
classmethod heat_per_mole()
```

Creates a quantity representing the total heat release from the liquidus temperature down to the current temperature [J/mol].

**Note:** The total or apparent heat release during the solidification process consists of two parts: one is the so-called latent heat, i.e. heat due to the liquid -> solid phase transformation (`latent_heat_per_mole()` and `latent_heat_per_gram()`), and the other is the heat related to the specific heat of liquid and solid phases (`heat_per_mole()` and `heat_per_gram()`).

**Returns** A new `HeatPerMole` object.

```python
classmethod latent_heat_per_gram()
```

Creates a quantity representing the cumulated latent heat release from the liquidus temperature down to the current temperature [J/g].

**Note:** The total or apparent heat release during the solidification process consists of two parts: one is the so-called latent heat, i.e. heat due to the liquid -> solid phase transformation (`latent_heat_per_mole()` and `latent_heat_per_gram()`), and the other is the heat related to the specific heat of liquid and solid phases (`heat_per_mole()` and `heat_per_gram()`).

**Returns** A new `LatentHeatPerGram` object.

```python
classmethod latent_heat_per_mole()
```

Creates a quantity representing the cumulated latent heat release from the liquidus temperature down to the current temperature [J/mol].

**Note:** The total or apparent heat release during the solidification process consists of two parts: one is the so-called latent heat, i.e. heat due to the liquid -> solid phase transformation (`latent_heat_per_mole()` and `latent_heat_per_gram()`), and the other is the heat related to the specific heat of liquid and solid phases (`heat_per_mole()` and `heat_per_gram()`).

**Returns** A new `LatentHeatPerMole` object.
classmethod mass_fraction_of_a_solid_phase(phase: str = 'All')
Creates a quantity representing the mass fraction of a solid phase.

Parameters phase – The name of the phase or All to choose all solid phases
Returns A new MassFractionOfASolidPhase object.

classmethod mass_fraction_of_all_liquid()
Creates a quantity representing the total mass fraction of all the liquid phase.
Returns A new MassFractionOfAllLiquid object.

classmethod mass_fraction_of_all_solid_phases()
Creates a quantity representing the total mass fraction of all solid phases.
Returns A new MassFractionOfAllSolidPhase object.

classmethod molar_volume_of_phase(phase: str = 'All')
Creates a quantity representing the molar volume of a phase [m^3/mol].

Parameters phase – The name of the phase or All to choose all phases
Returns A new MolarVolumeOfPhase object.

classmethod molar_volume_of_system()
Creates a quantity representing the molar volume of the system [m^3/mol].
Returns A new MolarVolumeOfSystem object.

classmethod mole_fraction_of_a_solid_phase(phase: str = 'All')
Creates a quantity representing the molar fraction of a solid phase.

Parameters phase – The name of the phase or All to choose all solid phases
Returns A new MoleFractionOfASolidPhase object.

classmethod mole_fraction_of_all_liquid()
Creates a quantity representing the total molar fraction of all the liquid phase.
Returns A new MoleFractionOfAllLiquid object.

classmethod mole_fraction_of_all_solid_phases()
Creates a quantity representing the total molar fraction of all solid phases.
Returns A new MoleFractionOfAllSolidPhases object.

classmethod site_fraction_of_component_in_phase(phase: str = 'All', component: str = 'All', sub_lattice_ordinal_no: int = None)
Creates a quantity representing the site fractions [-].

Parameters

• phase – The name of the phase, use All to choose all stable phases

• component – The name of the component, use All to choose all components

• sub_lattice_ordinal_no – The ordinal number (i.e. 0, 1, 2, ...) of the sublattice
  of interest, use None to choose all sublattices

Note: Consult the tab “Phases and Phase Constitution” in the System Definer activity of Thermo-Calc and
click on the respective phase in the list to obtain more information about the sublattices. The sublattices
are shown there in the same order as it needs to be specified here.
Returns A new SiteFractionOfComponentInPhase object.

classmethod temperature()  
Creates a quantity representing the temperature [K].

Returns A new Temperature object.

class tc_python.quantity_factory.ThermodynamicQuantity  
Bases: tc_python.quantity.AbstractQuantity  

Factory class providing quantities used for defining equilibrium calculations (single equilibrium, property and phase diagrams, ...) and their results.

Note: In this factory class only the most common quantities are defined, you can always use the Console Mode syntax strings in the respective methods as an alternative (for example: “NPM(*)”).

classmethod activity_of_component(component: str = 'All')  
Creates a quantity representing the activity of a component [-].

Parameters component – The name of the component, use “All” to choose all components

Returns A new ActivityOfComponent object.

Creates a quantity representing the chemical diffusion coefficient of a phase [m^2/s].

Parameters
  • phase – The name of the phase
  • diffusing_element – The diffusing element
  • gradient_element – The gradient element
  • reference_element – The reference element (for example “Fe” in a steel)

Returns A new ChemicalDiffusionCoefficient object.

classmethod composition_of_phase_as_mole_fraction(phase: str = 'All', component: str = 'All')  
Creates a quantity representing the composition of a phase [mole-fraction].

Parameters
  • phase – The name of the phase, use All to choose all stable phases
  • component – The name of the component, use All to choose all components

Returns A new CompositionOfPhaseAsMoleFraction object.

classmethod composition_of_phase_as_weight_fraction(phase: str = 'All', component: str = 'All')  
Creates a quantity representing the composition of a phase [weight-fraction].

Parameters
  • phase – The name of the phase, use All to choose all stable phases
  • component – The name of the component, use All to choose all components

Returns A new CompositionOfPhaseAsWeightFraction object.
classmethod gibbs_energy_of_a_phase (phase: str = 'All')
Creates a quantity representing the Gibbs energy of a phase \([J]\).

Parameters phase – The name of the phase or All to choose all phases

Returns A new GibbsEnergyOfAPhase object.

classmethod mass_fraction_of_a_component (component: str = 'All')
Creates a quantity representing the mass fraction of a component.

Parameters component – The name of the component or All to choose all components

Returns A new MassFractionOfAComponent object.

classmethod mass_fraction_of_a_phase (phase: str = 'All')
Creates a quantity representing the mass fraction of a phase.

Parameters phase – The name of the phase or All to choose all phases.

Returns A new MassFractionOfAPhase object.

classmethod mole_fraction_of_a_component (component: str = 'All')
Creates a quantity representing the mole fraction of a component.

Parameters component – The name of the component or All to choose all components

Returns A new MoleFractionOfAComponent object.

classmethod mole_fraction_of_a_phase (phase: str = 'All')
Creates a quantity representing the mole fraction of a phase.

Parameters phase – The name of the phase or All to choose all phases.

Returns A new MoleFractionOfAPhase object.

classmethod normalized_driving_force_of_a_phase (phase: str = 'All')
Creates a quantity representing normalized driving force of a phase [-].

Parameters phase – The name of the phase or All to choose all phases.

Returns A new DrivingForceOfAPhase object.

classmethod pressure ()
Creates a quantity representing the pressure \([\text{Pa}]\).

Returns A new Pressure object.

classmethod system_size ()
Creates a quantity representing the system size \([\text{mol}]\).

Returns A new SystemSize object.

classmethod temperature ()
Creates a quantity representing the temperature \([\text{K}]\).

Returns A new Temperature object.

classmethod tracer_diffusion_coefficient (phase: str, diffusing_element: str)
Creates a quantity representing tracer diffusion coefficient of a phase \([\text{m}^2/\text{s}]\).

Parameters

Warning: A driving force calculation requires that the respective phase has been set to the state DORMANT. The parameter All is only reasonable if all phases have been set to that state.

5.4. Module “quantity_factory”
• **phase** – The name of the phase

• **diffusing_element** – The diffusing element

    Returns  A new `TracerDiffusionCoefficient` object.

**classmethod volume_fraction_of_a_phase**(phase: str = ‘All’)

    Creates a quantity representing the volume fraction of a phase.

    **Parameters**
    - **phase** – The name of the phase or `All` to choose all phases

    **Returns**  A new `VolumeFractionOfAPhase` object.

### 5.5 Module “utils”

**class tc_python.utils.CompositionUnit**

    Bases: `enum.Enum`

    The composition unit.

    **MASS_FRACTION** = 2

        Mass fraction.

    **MASS_PERCENT** = 3

        Mass percent.

    **MOLE_FRACTION** = 0

        Mole fraction.

    **MOLE_PERCENT** = 1

        Mole percent.

**class tc_python.utils.Condition**(console_mode_syntax: str, value: float)

    Bases: `object`

    A condition expressed in Console mode syntax (e.g. “X(Cr)”).

**class tc_python.utils.ResultValueGroup** *(result_line_group_java)*

    Bases: `object`

    A x-y-dataset representing a line data calculation result (i.e. a Thermo-Calc quantity 1 vs. quantity 2).

    **Warning:** Depending on the calculator, the dataset might contain NaN-values to separate the data between different subsets.

    **Variables**

    - **x** – list of floats representing the first quantity (“x-axis”)

    - **y** – list of floats representing the second quantity (“y-axis”)

**class tc_python.utils.TemperatureProfile**

    Bases: `object`

    Represents a time-temperature profile used by non-isothermal precipitation calculations.

    **add_time_temperature**(time: float, temperature: float)

        Adds a time-temperature point to the non-isothermal temperature profile.

    **Parameters**
- **time** – The time [s]
- **temperature** – The temperature [K]

**Returns**
This `TemperatureProfile` object

### 5.6 Module “exceptions”

**exception** `tc_python.exceptions.APIServerException`
---
**Bases:** `tc_python.exceptions.GeneralException`
- An exception that occurred during the communication with the API-server. It is normally not related to an error in the user program.

**exception** `tc_python.exceptions.CalculationException`
---
**Bases:** `tc_python.exceptions.TCException`
- An exception that occurred during a calculation.

**exception** `tc_python.exceptions.ComponentNotExistingException`
---
**Bases:** `tc_python.exceptions.GeneralException`
- The selected component is not existing.

**exception** `tc_python.exceptions.DatabaseException`
---
**Bases:** `tc_python.exceptions.CalculationException`
- Error loading a thermodynamic or kinetic database, typically due to a misspelled database name or a database missing in the system.

**exception** `tc_python.exceptions.DegreesOfFreedomNotZeroException`
---
**Bases:** `tc_python.exceptions.CalculationException`
- The degrees of freedom in the system are not zero, i.e. not all required conditions have been defined. Please check the conditions given in the exception message.

**exception** `tc_python.exceptions.EquilibriumException`
---
**Bases:** `tc_python.exceptions.CalculationException`
- An equilibrium calculation has failed, this might happen due to inappropriate conditions or a very difficult problem that can not be solved.

**exception** `tc_python.exceptions.GeneralException`
---
**Bases:** `tc_python.exceptions.TCException`
- A general exception that might occur in different situations.

**exception** `tc_python.exceptions.InvalidCalculationConfigurationException`
---
**Bases:** `tc_python.exceptions.CalculationException`
- Thrown when errors are detected in the way calculations are set up.

**exception** `tc_python.exceptions.InvalidNumberOfResultGroupsException`
---
**Bases:** `tc_python.exceptions.ResultException`
- A calculation result contains several result groups, which is not supported for the used method.

**exception** `tc_python.exceptions.InvalidResultStateException`
---
**Bases:** `tc_python.exceptions.CalculationException`
- Trying to access an invalid result (for example a `SingleEquilibriumTempResult` object that got already invalidated by condition changes or a result that was invalidated by calling `invalidate()` on it).
exception tc_python.exceptions.LicenseException
    Bases: tc_python.exceptions.GeneralException
    No valid license for the API or any Thermo-Calc product used by it found.

exception tc_python.exceptions.NoDataForPhaseException
    Bases: tc_python.exceptions.ResultException
    There is no result data available for a selected phase.

exception tc_python.exceptions.PhaseNotExistingException
    Bases: tc_python.exceptions.GeneralException
    The selected phase is not existing, so no data can be provided for it.

exception tc_python.exceptions.ResultException
    Bases: tc_python.exceptions.TCException
    An exception that occurred during the configuration of a calculation result.

exception tc_python.exceptions_SyntaxException
    Bases: tc_python.exceptions.CalculationException
    Syntax error in a Console Mode expression.

exception tc_python.exceptions.TCException
    Bases: Exception
    The root exception of TC-Python.

tc_python.exceptions.raise_python_exceptions(func)
    Internal method of the API: Usage of that decorator maps all relevant Java exceptions in the API to the appropriate Python exception.

5.7 Module “abstract_base”

class tc_python.abstract_base.AbstractCalculation(calculator)
    Bases: object
    Abstract base class for calculations.
    get_configuration_as_string() → str
    Returns detailed information about the current state of the calculation object.

    Warning: The structure of the calculator objects is an implementation detail and might change between releases without notice. Therefore do not rely on the internal object structure.

class tc_python.abstract_base.AbstractResult(result)
    Bases: object
    Abstract base class for results. This can be used to query for specific values.
    invalidate()
    Invalidates the object and frees the disk space used by it. This is only required if the disk space occupied by the object needs to be released during the calculation. No data can be retrieved from the object afterwards.
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