teqp Release 0.19.1

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CONTENTS:

1	Getti 1.1 1.2	ng Started 1 Introduction 1 Installation 1
2	C++ 2.1 2.2 2.3	interface Introduction
3	Deriv 3.1 3.2	Thermodynamic Derivatives
4	Mode 4.1 4.2 4.3 4.4 4.5 4.6 4.7 4.8 4.9 4.10 4.11 4.12 4.13 4.14 4.15 4.16 4.17 4.18	els 15 Constructing Models 15 General cubics 17 Quantum PR 20 Advanced cubic mixing rules 23 RK-PR 26 Cubic Plus Association (CPA) 36 LKP (Lee-Kesler-Plöcker) 31 Model Potentials 33 PC-SAFT 35 SAFT-VR-Mie 41 SAFT-VR-Mie with polar contributions 47 Association 56 Multi-fluid EOS 58 Multfluid mutant 61 REFPROP < 10.0 conversion
5	Algor 5.1 5.2 5.3 5.4 5.5 5.6	rithms 73 Phase equilibria 73 Tracing (isobars and isotherms) 75 VLLE 78 VLLE @ constant pressure 81 Critical curves & points 83 Information 90

	Examples 6.1 The teqp paper in I&ECR	91 91
	Fitting 7.1 Multi-fluid Parameter Fitting	97 97
	teqp8.1teqp package	101 101
9	Indices and tables	
Рy	thon Module Index	
Inc	dex	117

CHAPTER

ONE

GETTING STARTED

1.1 Introduction

teqp (phonetically: tɛk pi) is a C++-based library with wrappers. It was written because implementing EOS (particularly working out the derivatives) is a painful, error-prone, and slow process. The advent of open-source automatic differentiation libraries makes the implementation of EOS as fast as hand-written derivatives, and much easier to implement without errors.

There is a paper about teqp: https://doi.org/10.1021/acs.iecr.2c00237

The documentation is based on the Python wrapper because it can be readily integrated with the documentation tools (sphinx in this case) and can be auto-generated at documentation build time.

1.2 Installation

1.2.1 Python

The library can be installed with:

```
pip install teqp
```

because the binary wheels for all major platforms are provided on pypi.

If you desire to build teap yourself, it is recommended to pull from github and build a binary wheel, and then subsequently install that wheel:

```
git clone --recursive https://github.com/usnistgov/teqp
cd teqp
python setup.py bdist_wheel
pip install dist/*.whl # or replace with the appropriate binary wheel
```

1.2.2 C++

The build is cmake based. There are targets available for an interface library, etc. Please see CMakeLists.txt

CHAPTER

TWO

C++ INTERFACE

2.1 Introduction

The abstract base class defining the public C++ interface of teqp is documented in AbstractModel. This interface was developed because re-compilation of the core of teqp is VERY slow, due to the heavy use of templates, which makes the code very flexible, but difficult to work with when doing development. Especially users that would like to only *use* the library but not be forced to pay the price of recompilation benefit from this approach.

As a user, a new model instance (a std::unique_ptr<teqp::AbstactModel*>) can be created by passing properly formatted JSON data structure to the make model() function.

2.2 Object Model

The object model in teqp is convoluted because of the requirements to have models that use templated types to allow the use of automatic differentiation types. Instances of classes with templated methods cannot be stored directly in generic STL containers like std::vector or std::list (though they can be stored in std::tuple, but tuple cannot be constructed at runtime because they have complete type knowledge and C++ is strongly typed). Thus, some sort of wrapping is required (in C++ the technical term is type erasure) to store objects of a homogenous interface in dynamic containers like std::vector.

A number of type-erasure classes are defined, especially the DerivativeAdapter class which does type erasure on a model that it holds. This DerivativeAdapter class has an interface that takes STL types (and Eigen arrays in some cases) as input arguments, and then calls lower-level methods that can operate with a range of different numerical types, and call the templated methods of a model.

As a developer/implementer of a thermodynamic model, the class implementing the thermodynamic model for a contribution to α must satisfy the following requirements:

- It must have a method called alphar that takes three arguments that are all generic types. The first argument is the temperature, the second argument is the molar density, and the third is the mole fractions. In the case of some equations of state for model potentials, the temperature and density are treated as being in reduced units. The function should be called alphar even for Helmholtz energy contributions that are for ideal gases. You can think of the r in alphar standing for reduced instead of residual if that helps.
- It must have a method called R that takes a single argument that is the mole fractions of the components. It then returns the molar gas constant of the mixture. For most models it suffices to return 8.31446261815324, which is the CODATA value of the molar gas constant, and is available in the teqp::constants namespace. The reason the R method must be implemented is the multiparameter models in which the molar gas constant of differt components is slightly different based upon when the EOS was published. Also, some of the other models used different values of R (or Avogadro's constant) when being developed and if you want to get perfect reproducibility these details matter.

This model instance is then passed to one of two methods in the teqp::cppinterface::adapter namespace: teqp::cppinterface::adapter::make_owned() or teqp::cppinterface::adapter::make_cview(). As the name suggests, if you pass the class instance to the make_owned function, it takes ownership of the model and the argument passed to the function is invalidated. On the contrary, the make_oview method is just a "viewer" of the model without taking ownership, so you need to watch out that the lifetime of the model you pass to this function is longer than the time you are using the wrapper model.

For instance this minimal working model of the van der Waals EOS demonstrates some of the things to be aware of:

```
/// A (very) simple implementation of the van der Waals EOS
class myvdWEOS1 {
public:
    const double a, b;
   myvdWEOS1(double a, double b) : a(a), b(b) {};
    /// \brief Get the universal gas constant
   template<class VecType>
   auto R(const VecType& /*molefrac*/) const { return constants::R_CODATA2017; }
   /// The evaluation of f \alpha^{\rm r}=a/(RT) \f$
   /// \param T The temperature
   /// \param rhotot The molar density
    /// \param molefrac The mole fractions of each component
   template<typename TType, typename RhoType, typename VecType>
   auto alphar(const TType &T, const RhoType& rhotot, const VecType &molefrac) const
        return teqp::forceeval(-log(1.0 - b * rhotot) - (a / (R(molefrac) * T)) *_
→rhotot);
    }
};
```

The name of the class is entirely arbitrary, you could call it just as well GreatVdWModel instead of myvdWEOS1.

A complete example could then read:

```
#include <catch2/catch_test_macros.hpp>
#include "teqp/cpp/teqpcpp.hpp"
#include "tegp/cpp/deriv_adapter.hpp"
#include "teqp/types.hpp"
#include "tegp/constants.hpp"
/// A (very) simple implementation of the van der Waals EOS
class myvdWEOS1 {
public:
    const double a, b;
    myvdWEOS1(double a, double b) : a(a), b(b) {};
    /// \brief Get the universal gas constant
    template<class VecType>
    auto R(const VecType& /*molefrac*/) const { return constants::R_CODATA2017; }
   /// The evaluation of f \alpha^{\rm r}=a/(RT) \f$
    /// \param T The temperature
    /// \param rhotot The molar density
    /// \param molefrac The mole fractions of each component
    template<typename TType, typename RhoType, typename VecType>
    auto alphar(const TType &T, const RhoType& rhotot, const VecType &molefrac) const
                                                                          (continues on next page)
```

In this runnable example (runnable once the include paths are correct and the code is linked against the teqpcpp C++ library), a new factory function is registered with the add_model_pointer_factory_function() function and then this function is used to generate a std::unique_ptr<AbstractModel*>. Once the model has been created, it is possible to cast it back to the original type, but you must know the type of the class that you are holding (at compile time). The teqp::cppinterface::adapter::get_model_cref() is a convenience function to do this casting.

2.3 C++ Details

2.3.1 Don't return expressions

The most important thing to be sure of when developing models in teqp is that you do not return expressions from functions. For instance in the simple function:

```
template<typename T1, typename T2>
auto alphar(const T1 &v1, const T2& v2) {
    return v1 + v2;
}
```

if the types of T1 and T2 are both $\mathtt{autodiff:::real}$ (the same problem occurs for other autodiff types), the value of $\mathtt{v1} + \mathtt{v2}$ is an expression type that is lazily evaluated, and the expression holds references to the actual values of the variables $\mathtt{v1}$ and $\mathtt{v2}$. This lazy evaluation is how autodiff can be so fast. Once the expression is returned from this function, the variables that it was pointing to are no longer valid because they have fallen out of scope and you can silently be pointing to invalid memory locations.

In order to avoid this problem you can use the function teqp::forceeval to force the evaluation of the expression, copying all the variables into the expression, and removing the possibility of dangling references after the function returns.

One way to ensure that you are not running into this problem is to enable the Address Sanitizer option "Detect Use of stack after return" in XCode (its in the Diagnostic panel of the "Edit Scheme..." option). Other address sanitizer tools have similar functionality.

2.3. C++ Details 5

2.3.2 Generic return types

Taking the example shown above, in the function alphar all the arguments have templated type. Sometimes you will need to make use of one or more of the types in intermediate calculations within the function, and you might need to determine the type of an expression to for instance allocate a vector of this type. As an example, let's say that we are going to multiply three different variables together. In the alphar context, let's assume that T is of type double, rhomolar is of type std::complex<double> and molefracs is of type Eigen::ArrayXcd. In the case of the expression T*rhomolar*molefracs[0], the result will be calculated based on the type promotion to a std::complex<double>, so the result type of this product is std::complex<double>. If you want to let the compiler determine this type for you, you can do:

and if you need want to work with the types of the variables, usually because you need to cover all your bases for all the templat permutations, you can do instead

and if you need to remove the const of your variable types, you can do with std::decay_t< >.

CHAPTER

THREE

DERIVATIVES

3.1 Thermodynamic Derivatives

3.1.1 Helmholtz energy derivatives

Thermodynamic derivatives are at the very heart of teqp. All models are defined in the form $\alpha^r(T, \rho, z)$, where ρ is the molar density, and z are mole fractions. There are exceptions for models for which the independent variables are in simulation units (Lennard-Jones and its ilk).

Thereofore, to obtain the residual pressure, it is obtained as a derivative:

$$p^r = \rho RT \left(\rho \left(\frac{\partial \alpha^r}{\partial \rho} \right)_T \right)$$

and other residual thermodynamic properties are defined likewise.

We can define the concise derivative

$$\Lambda_{xy}^r = (1/T)^x (\rho)^y \left(\frac{\partial^{x+y} (\alpha^r)}{\partial (1/T)^x \partial \rho^y} \right)$$

so we can re-write the derivative above as

$$p^r = \rho RT\Lambda_{01}^r$$

Similar definitions apply for all the other thermodynamic properties, with the tot superscript indicating it is the sum of the residual and ideal-gas (not included in teqp) contributions:

$$\frac{p}{\rho RT} = 1 + \Lambda_{01}^r$$

Internal energy (u = a + Ts):

$$\frac{u}{RT} = \Lambda_{10}^{\text{tot}}$$

Enthalpy $(h = u + p/\rho)$:

$$\frac{h}{RT} = 1 + \Lambda_{01}^r + \Lambda_{10}^{\rm tot}$$

Entropy $(s \equiv -(\partial a/\partial T)_v)$:

$$\frac{s}{B} = \Lambda_{10}^{\rm tot} - \Lambda_{00}^{\rm tot}$$

Gibbs energy (g = h - Ts):

$$\frac{g}{RT} = 1 + \Lambda_{01}^r + \Lambda_{00}^{\text{tot}}$$

Derivatives of pressure:

$$\left(\frac{\partial p}{\partial \rho}\right)_T = RT\left(1 + 2\Lambda_{01}^r + \Lambda_{02}^r\right)$$

$$\left(\frac{\partial p}{\partial T}\right)_{\rho} = R\rho \left(1 + \Lambda_{01}^{r} - \Lambda_{11}^{r}\right)$$

Isochoric specific heat $(c_v \equiv (\partial u/\partial T)_v)$:

$$\frac{c_v}{R} = -\Lambda_{20}^{\mathrm{tot}}$$

Isobaric specific heat $(c_p \equiv (\partial h/\partial T)_p)$; see Eq. 3.56 from Span for the derivation):

$$\frac{c_p}{R} = -\Lambda_{20}^{\text{tot}} + \frac{(1 + \Lambda_{01}^r - \Lambda_{11}^r)^2}{1 + 2\Lambda_{01}^r + \Lambda_{02}^r}$$

In teqp, these derivatives are obtained from methods like

- get_Arxy()
- get_Ar06n()

where the A in this context indicates the variable Λ above. This naming is perhaps not ideal, since A is sometimes the total Helmholtz energy, but it was a close visual mnemonic to the character Λ .

```
[1]: import teqp
    teqp.__version__
[1]: '0.19.1'
```

[2]: import numpy as np

```
[3]: Tc_K = [300]
pc_Pa = [4e6]
acentric = [0.01]
model = teqp.canonical_PR(Tc_K, pc_Pa, acentric)
```

```
[4]: z = np.array([1.0])
    model.get_Ar01(300,300,z)

[4]: -0.06836660379313926
```

And there are additional methods to obtain all the derivatives up to a given order:

```
[5]: model.get_Ar06n(300,300,z) # derivatives 00, 01, 02, ... 06
[5]: array([-6.96613834e-02, -6.83666038e-02, 2.53578225e-03, -1.57011622e-04, 1.68186288e-05, -2.23059409e-06, 3.82592585e-07])
```

But more derivatives are slower than fewer:

```
[6]: %timeit model.get_Ar01(300,300,z)
%timeit model.get_Ar04n(300,300,z)
```

```
593 ns ± 1.2 ns per loop (mean ± std. dev. of 7 runs, 1,000,000 loops each)

1.08 µs ± 1.14 ns per loop (mean ± std. dev. of 7 runs, 1,000,000 loops each)
```

Note: calling overhead is usually on the order of 1 microsecond

3.1.2 Virial coefficients

Virial coefficients represent the thermodynamics of the interaction of two-, three-, ... bodies interacting with each other. They can be obtained rigorously if the potential energy surface of interaction is fully known. In general, such a surface can only be constructed for small rigid molecules. Many simple thermodynamic models do a poor job of predicting the thermodynamics captured by the virial coefficients.

The i-th virial coefficient is defined by

$$B_i = \frac{(\alpha^r)^{(i-1)}}{(i-2)!}$$

with the concise derivative term

$$(\alpha^r)^{(i)} = \lim_{\rho \to 0} \left(\frac{\partial^i \alpha^r}{\partial \rho^i} \right)_{T, \vec{x}}$$

teqp supports the virial coefficient directly, there is the get_B2vir method for the second virial coefficient:

```
[7]: model.get_B2vir(300, z)
[7]: -0.00023661263734465424
```

And the get_Bnvir method that allows for the calculation of higher virial coefficients:

```
[8]: model.get_Bnvir(7, 300, z)

[8]: {2: -0.0002366126373446542,
3: 3.001768410777936e-08,
4: -3.2409760373816364e-12,
5: 3.961781646633723e-16,
6: -4.5529239838367004e-20,
7: 5.375927851118494e-24}
```

The get_Bnvir method was implemented because when doing automatic differentiation, all the intermediate derivatives are also retained.

There is also a method to calculate temperature derivatives of a given virial coefficient

3.1.3 Isochoric Thermodynamics Derivatives

In the isochoric thermodynamics formalism, the EOS is expressed in the Helmholtz energy density Ψ as a function of temperature and molar densities $\vec{\rho}$. This formalism is handy because it allows for a concise mathematical structure, well suited to implementation in teqp. For instance the pressure is obtained from (see https://doi.org/10.1002/aic.16074):

$$p = -\Psi + \sum_{i=1}^{N} \rho_i \mu_i$$

with the chemical potential μ_i obtained from

$$\mu_i = \left(\frac{\partial \Psi}{\partial \rho_i}\right)_{T, \rho_{i \neq i}}$$

The molar densities ρ_i are related to the total density and the mole fractions:

$$\rho_i = x_i \rho$$

In teqp, the isochoric derivative functions like <code>get_fugacity_coefficients</code>, <code>get_partial_molar_volumes</code> take as arguments the temperature T and the vector of molar concentrations <code>rhovec= $\vec{\rho}$ </code>, which are obtained by multiplying the mole fractions by the total density.

Example:

```
[10]: model = teqp.build_multifluid_model(["CO2","Argon"], teqp.get_datapath())
   T, rhovec = 300, np.array([0.3,0.4])*300 # K, mol/m^3
   display(model.get_fugacity_coefficients(T, rhovec))
   display(model.get_partial_molar_volumes(T, rhovec))
   array([0.97884567, 0.99866748])
   array([0.00470644, 0.00480351])
```

3.2 Composition derivatives

For other mixture calculations composition derivatives of the form

$$\Lambda_{xyz_i}^r = (1/T)^x (\rho)^y \left(\frac{\partial^{x+y+z_i} (\alpha^r)}{\partial (1/T)^x \partial \rho^y \partial \mathbf{Z}_i^{z_i}} \right)$$

are needed. This function is exposed in teqp (as of version 0.19) as the function get_ATrhoXi. In order to limit the binary size and compilation time, x has a max of 2 and y does as well. z_i can be up to 3, and must be at least 1, otherwise you can use the other derivative functions that do not require any composition derivatives.

The mixed composition derivative of the form

$$\Lambda_{xyz_iz_j}^r = (1/T)^x (\rho)^y \left(\frac{\partial^{x+y+z_i+z_j} (\alpha^r)}{\partial (1/T)^x \partial \rho^y \partial \mathbf{Z}_i^{z_i} \partial \mathbf{Z}_j^{z_j}} \right)$$

supports x and y either 0 or 1, with at most two composition derivatives. The triple composition derivative of the form

$$\Lambda_{xyz_iz_jz_k}^r = (1/T)^x (\rho)^y \left(\frac{\partial^{x+y+z_i+z_j+z_k} (\alpha^r)}{\partial (1/T)^x \partial \rho^y \partial \mathbf{Z}_i^{z_i} \partial \mathbf{Z}_j^{z_j} \partial \mathbf{Z}_k^{z_k}} \right)$$

supports x and y either 0 or 1, with up to first derivatives in each composition variable. If this is not enough derivatives, open a feature request here: https://github.com/usnistgov/teqp/issues

3.2.1 au and δ derivatives

In the multi-fluid modeling approach used in NIST REFPROP and the GERG-2004 GERG-2008 models, the derivatives are in the form

$$\Lambda^{r}_{xyz_{i}} = \tau^{x} \delta^{y} \left(\frac{\partial^{x+y+z_{i}}(\alpha^{r})}{\partial \tau^{x} \partial \delta^{y} \partial \mathbf{Z}_{i}^{z_{i}}} \right)$$

with $\tau = T_{\rm red}(\mathbf{Z})/T$ and $\delta = \rho/\rho_{\rm red}(\mathbf{Z})$. The higher derivatives are similarly equal to

$$\Lambda_{xyz_iz_j}^r = \tau^x \delta^y \left(\frac{\partial^{x+y+z_i+z_j} (\alpha^r)}{\partial \tau^x \partial \delta^y \partial \mathbf{Z}_i^{z_i} \partial \mathbf{Z}_j^{z_j}} \right)$$

$$\Lambda^{r}_{xyz_{i}z_{j}z_{k}} = \tau^{x}\delta^{y} \left(\frac{\partial^{x+y+z_{i}+z_{j}+z_{k}}(\alpha^{r})}{\partial \tau^{x}\partial \delta^{y}\partial \mathbf{Z}_{i}^{z_{i}}\partial \mathbf{Z}_{j}^{z_{j}}\partial \mathbf{Z}_{k}^{z_{k}}} \right)$$

The same limitations on the numbers of derivatives are used for the derivatives with (1/T) and ρ as independent variables.

The Python methods are documented here:

- get_ATrhoXi()
- get_ATrhoXiXj()
- get_ATrhoXiXjXk()
- get_AtaudeltaXi()
- get AtaudeltaXiXi()
- get AtaudeltaXiXiXk()

3.2.2 xN (in)dependent

Let's suppose that some quantity Υ depends on mole fractions. If all the mole fractions are considered to be independent, the total differential is obtained from

$$\mathrm{d}\Upsilon = \sum_{j} \left(\frac{\partial \Upsilon}{\partial x_{j}} \right)_{x_{i \neq j}} \mathrm{d}x_{j}$$

If instead the last mole fraction is defined to be dependent on the others via

$$x_N = 1 - \sum_{i=1}^{N-1} x_i$$

then the total differential is obtained from

$$\mathrm{d}\Upsilon = \sum_{j=1}^{N-1} \left(\frac{\partial \Upsilon}{\partial x_j}\right)_{x_{i\neq j}} \mathrm{d}x_j + \left(\frac{\partial \Upsilon}{\partial x_N}\right)_{x_{i\neq j}} \mathrm{d}x_N$$

where x_N is considered to be an independent variable in the derivative $\left(\frac{\partial \Upsilon}{\partial x_N}\right)_{x_{i\neq j}}$. Thus derivatives with respect to one of the dependent mole fractions $(x_k$ with k < N) would be equal to

$$\left(\frac{\partial \Upsilon}{\partial x_k}\right)_{x_{j \neq k}} = \left(\frac{\partial \Upsilon}{\partial x_k}\right)_{x_{i \neq k}} - \left(\frac{\partial \Upsilon}{\partial x_N}\right)_{x_{i \neq N}}$$

because

$$\left(\frac{\partial x_N}{\partial x_i}\right) = -1$$

So if the library (e.g., CoolProp and TREND) allows for the fractions to be dependent (either option is allowed in CoolProp, TREND uses N-1 independent mole fractions), you can use the molar composition derivatives with the mole fractions treated as being independent to obtain derivatives with one of the mole fractions dependent on the other N-1 fractions.

```
[1]: import teqp, numpy as np
    teqp.__version_
[1]: '0.19.1'
[2]: j = {
         'components': ["Methane", "Nitrogen", "Oxygen"],
        'root': teqp.get_datapath(),
        'BIP': '',
        'departure': ''
    model = teqp.make_model({'kind':'multifluid', 'model': j})
    T = 300 \# K
    rhomolar = 3000 \# mol/m^3
    z = np.array([0.3, 0.5, 0.2]) # mole fractions
    Tr = model.get_Tr(z)
    rhor = model.get_rhor(z)
    tau = Tr/T
    delta = rhomolar/rhor
    Ntau = 0
    Ndelta = 0
    Nxi = 1
    print(model.get_AtaudeltaXi(tau, Ntau, delta, Ndelta, z, 0, Nxi))
    Ntau = 1
    Ndelta = 0
    Nxi = 1
    print(model.get_AtaudeltaXi(tau, Ntau, delta, Ndelta, z, 0, Nxi))
    Ntau = 0
    Ndelta = 1
    Nxi = 1
    print(model.get_AtaudeltaXi(tau, Ntau, delta, Ndelta, z, 0, Nxi))
    Ntau = 2
    Ndelta = 0
    Nxi = 1
    print(model.get_AtaudeltaXi(tau, Ntau, delta, Ndelta, z, 0, Nxi))
    Ntau = 1
    Ndelta = 1
    Nxi = 1
    print(model.get_AtaudeltaXi(tau, Ntau, delta, Ndelta, z, 0, Nxi))
```

(continues on next page)

```
Ntau = 0
Ndelta = 2
Nxi = 1
print(model.get_AtaudeltaXi(tau, Ntau, delta, Ndelta, z, 0, Nxi))
Ntau = 1
Ndelta = 0
Nxi = 1
Nxj = 1
print(model.get_AtaudeltaXiXj(tau, Ntau, delta, Ndelta, z, 0, Nxi, 1, Nxj))
Ntau = 0
Ndelta = 1
Nxi = 1
Nxi = 1
print(model.get_AtaudeltaXiXj(tau, Ntau, delta, Ndelta, z, 0, Nxi, 1, Nxj))
Ntau = 0
Ndelta = 0
Nxi = 1
Nxj = 1
Nxk = 1
print(model.get_AtaudeltaXiXjXk(tau, Ntau, delta, Ndelta, z, 0, Nxi, 1, Nxj, 2, Nxk))
-0.043587384253511226
-0.2118857998812584
-0.03650566667904927
-0.07488856488580686
-0.2069389009652925
0.014468933385218782
-0.005978809921279949
-0.00279185550001082
0.0
```

With CoolProp, version 6.6.0, the following script in C++:

```
#include "AbstractState.h"
#include "Backends/Helmholtz/MixtureDerivatives.h"
int main(){
    std::shared_ptr<CoolProp::AbstractState> AS(
        CoolProp::AbstractState::factory("HEOS","Methane&Nitrogen&Oxygen")
   );
   AS->set_mole_fractions({0.3, 0.5, 0.2});
   AS->specify_phase(CoolProp::iphase_gas);
   AS->update(CoolProp::DmolarT_INPUTS, 3000, 300);
   auto& HEOS = *dynamic_cast<CoolProp::HelmholtzEOSMixtureBackend*>(AS.get());
   auto xN = CoolProp::x_N_dependency_flag::XN_INDEPENDENT;
   using md = CoolProp::MixtureDerivatives;
   std::cout << md::dalphar_dxi(HEOS, 0, xN) << std::endl;</pre>
   \verb|std::cout| << \verb|md::d2alphar_dxi_dTau(HEOS, 0, xN)| *AS-> tau() << \verb|std::endl|; |
    std::cout << md::d2alphar_dxi_dDelta(HEOS, 0, xN)*AS->delta() << std::endl;</pre>
    std::cout << md::d3alphar_dxi_dTau2(HEOS, 0, xN)*pow(AS->tau(), 2) << std::endl;</pre>
    std::cout << md::d3alphar_dxi_dDelta_dTau(HEOS, 0, xN)*AS->tau()*AS->delta() <<_
    std::cout << md::d3alphar_dxi_dDelta2(HEOS, 0, xN)*pow(AS->delta(), 2) << std::
```

(continues on next page)

```
std::cout << md::d3alphar_dxi_dxj_dTau(HEOS, 0, 1, xN)*AS->tau() << std::endl;
std::cout << md::d3alphar_dxi_dxj_dDelta(HEOS, 0, 1, xN)*AS->delta() << std::endl;
std::cout << md::d3alphardxidxjdxk(HEOS, 0, 1, 2, xN) << std::endl;
}</pre>
```

yields the output:

```
-0.0435874

-0.211886

-0.0365057

-0.0748886

-0.206939

0.0144689

-0.00597881

-0.00279186
```

which is the same as the above

CHAPTER

FOUR

MODELS

4.1 Constructing Models

With a few exceptions, most models are constructed by describing the model in JSON format, and passing the JSON-formatted information to the make_model function. There are some convenience functions exposed for backwards compatibility, but as of version 0.14.0, all model construction should go via this route.

At the C++ level, the returned value from the make_model function is a shared_ptr that wraps a pointer to an AbstractModel class. The AbstractModel class is an abstract class which defines the public C++ interface.

In Python, construction is in two parts. First, the model is constructed, which only includes the common methods. Then, the model-specific attributes and methods are attached with the attach_model_specific_methods method.

The JSON structure is of two parts, the kind field is a case-sensitive string defining which model kind is being constructed, and the model field contains all the information needed to build the model. In the case of hard-coded models, nothing is provided in the model field, but it must still be provided.

Also, the argument to make_model must be valid JSON. So if you are working with numpy array datatypes, make sure to convert them to a list (which is convertible to JSON). Example below.

```
[1]: import teqp, numpy as np
    teqp.__version__
[1]: '0.19.1'
[2]: teqp.make_model({'kind': 'vdW1', 'model': {'a': 1, 'b': 2}})
[2]: <teqp.teqp.AbstractModel at 0x7652ac72bfb0>
[3]: # Fields are case-sensitive
    teqp.make_model({'kind': 'vdW1', 'model': {'a': 1, 'B': 2}})
    RuntimeError
                                                Traceback (most recent call last)
    Cell In[3], line 2
          1 # Fields are case-sensitive
     ----> 2 teqp.make_model({'kind': 'vdW1', 'model': {'a': 1, 'B': 2}})
    File /opt/conda/lib/python3.11/site-packages/teqp/__init__.py:47, in make_model(*args,

    **kwargs)

         42 def make_model(*args, **kwargs):
         43
         44
                 This function is in two parts; first the make_model function (renamed to _
     →make_model in the Python interface)
                is used to make the model and then the model-specific methods are.
                                                                                (continues on next page)
```

```
→attached to the instance
         46
     ---> 47
                AS = _make_model(*args, **kwargs)
         48
                attach_model_specific_methods(AS)
          49
                return AS
    RuntimeError: :{"B":2,"a":1}': required property 'b' not found in object
     |/|\|:{"B":2, "a":1}': validation failed for additional property 'B': instance invalid
     →as per false-schema
[4]: # A hard-coded model
    teqp.make_model({
        'kind': 'AmmoniaWaterTillnerRoth',
        'model': {}
    })
[4]: <teqp.teqp.AbstractModel at 0x765290d8f9b0>
[5]: # Show what to do with numpy array
    Tc_K = np.array([100,200])
    pc_Pa = np.array([3e6, 4e6])
    teqp.make_model({
        "kind": "vdW",
         "model": {
            "Tcrit / K": Tc_K.tolist(),
            "pcrit / Pa": pc_Pa.tolist()
     })
[5]: <teqp.teqp.AbstractModel at 0x765290d8fb90>
[6]: # methane with conventional PC-SAFT
    j = {
         'kind': 'PCSAFT',
         'model': {
             'coeffs': [{
                 'name': 'methane',
                  'BibTeXKey': 'Gross-IECR-2001',
                  'm': 1.00,
                  'sigma_Angstrom': 3.7039,
                  'epsilon_over_k': 150.03,
             } ]
         }
    model = teqp.make_model(j)
```

4.2 General cubics

The reduced residual Helmholtz energy for the main cubic EOS (van der Waals, Peng-Robinson, and Soave-Redlich-Kwong) can be written in a common form (see https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7365965/)

$$\alpha^{r} = \psi^{(-)} - \frac{\tau a_m}{RT_r} \psi^{(+)}$$

$$\psi^{(-)} = -\ln(1 - b_m \rho)$$

$$\psi^{(+)} = \frac{\ln\left(\frac{\Delta_1 b_m \rho + 1}{\Delta_2 b_m \rho + 1}\right)}{b_{rr}(\Delta_1 - \Delta_2)}$$

with the constants given by:

- vdW: $\Delta_1 = 0$, $\Delta_2 = 0$
- SRK: $\Delta_1 = 1, \Delta_2 = 0$
- PR: $\Delta_1 = 1 + \sqrt{2}$, $\Delta_2 = 1 \sqrt{2}$

The quantities a_m and b_m are described (with exact solutions for the numerical coefficients) for each of these EOS in https://pubs.acs.org/doi/abs/10.1021/acs.iecr.1c00847.

The models in teqp are instantiated based on knowledge of the critical temperature, pressure, and acentric factor. Thereafter all quantities are obtained from derivatives of α^r .

The Python class is here: GeneralizedCubic

```
[1]: import teqp
    teqp.__version_
[1]: '0.19.1'
[2]: import json
    import CoolProp.CoolProp as CP
    # Values taken from http://dx.doi.org/10.6028/jres.121.011
    Tc_K = [190.564, 154.581, 150.687]
    pc_Pa = [4599200, 5042800, 4863000]
    acentric = [0.011, 0.022, -0.002]
    # Instantiate Peng-Robinson model
    modelPR = teqp.canonical_PR(Tc_K, pc_Pa, acentric)
    # Instantiate Soave-Redlich-Kwong model
    modelSRK = teqp.canonical_SRK(Tc_K, pc_Pa, acentric)
[3]: # And you can get information about the model in JSON format
    # from the get_meta function
    modelPR.get_meta()
[3]: {'Delta1': 2.414213562373095,
     'Delta2': -0.41421356237309515,
     'OmegaA': 0.4572355289213822,
     'OmegaB': 0.07779607390388846,
     'kind': 'Peng-Robinson'}
```

4.2. General cubics 17

4.2.1 Adjusting k ij

Fine-tuned values of k_{ij} can be provided when instantiating the model, for Peng-Robinson and SRK. A complete matrix of all the k_{ij} values must be provided. This allows for asymmetric mixing models in which $k_{ij} \neq k_{ji}$.

```
[4]: k_12 = 0.01
kmat = [[0,k_12,0],[k_12,0,0],[0,0,0]]
teqp.canonical_PR(Tc_K, pc_Pa, acentric, kmat)
teqp.canonical_SRK(Tc_K, pc_Pa, acentric, kmat)
[4]: <teqp.teqp.AbstractModel at 0x7ee9b01157f0>
```

4.2.2 Superancillary

The superancillary equation gives the co-existing liquid and vapor (orthobaric) densities as a function of temperature. The set of Chebyshev expansions was developed in https://pubs.acs.org/doi/abs/10.1021/acs.iecr.1c00847. These superancillary equations are more accurate than iterative calculations in double precision arithmetic and also at least 10 times faster to calculate, and cannot fail in iterative routines, even extremely close to the critical point.

The superancillary equation is only exposed for pure fluids to remove ambiguity when considering mixtures. The returned tuple is the liquid and vapor densities

```
[5]: teqp.canonical_PR([Tc_K[0]], [pc_Pa[0]], [acentric[0]]).superanc_rhoLV(100)
[5]: (30846.392909514052, 42.480231719002326)
```

4.2.3 a and b

For the cubic EOS, it can be useful to obtain the a and b parameters directly. The b parameter is particularly useful because 1/b is the maximum allowed density in the EOS

```
[6]: import numpy as np
z = np.array([0.3, 0.4, 0.3])
modelPR.get_a(140, z), modelPR.get_b(140, z)
[6]: (0.1874177858906821, 2.1984349667726406e-05)
```

4.2.4 alpha functions

It can be advantageous to modify the alpha function to allow for more accurate handling of the attractive interactions. Coefficients are tabulated for many species in https://pubs.acs.org/doi/10.1021/acs.jced.7b00967 for the Peng-Robinson EOS with Twu alpha function and the values from the SI of that paper are in the csv file next to this file.

```
[7]: import pandas

dfTwu = pandas.read_csv('fitted_Twu_coeffs.csv')

def get_model(INCHIKey):
    row = dfTwu.loc[dfTwu['inchikey']==INCHIKey]
    if len(row) == 1:
        row = row.iloc[0]
    Tc_K = row['Tc_K']
    pc_MPa = row['pc_MPa']
    c = [row['c0'], row['c1'], row['c2']]

(continues on next page)
```

```
# The JSON definition of the EOS,
# here a generic cubic EOS to allow for
# specification of the alpha function(s)
j = {
    'kind': 'cubic',
    'model': {
        'type': 'PR',
        'Tcrit / K': [Tc_K],
        'pcrit / Pa': [pc_MPa*1e6],
        'acentric': [0.1],
        'alpha': [{'type': 'Twu', 'c': c}]
}
model = teqp.make_model(j)
return model, j

# Hexane
model, j = get_model(INCHIKey='VLKZOEOYAKHREP-UHFFFAOYSA-N')
```

```
[8]: # And how about we calculate the pressure and s^+ = -sr/R at NBP of water
model, j = get_model(INCHIKey='XLYOFNOQVPJJNP-UHFFFAOYSA-N') # WATER

T = 373.1242958476844 # K, NBP of water
rhoL, rhoV = model.superanc_rhoLV(T)
z = np.array([1.0])
pL = rhoL*model.get_R(z)*T*(1.0 + model.get_Ar01(T, rhoL, z))
splusL = model.get_splus(T, rhoL*z)
print(pL, splusL)

102739.27983424198 6.03697343297877
```

Also implemented in version 0.17 are the alpha functions of Mathias-Copeman.

$$\alpha = (1 + c_0 x + c_1 x^2 + c_2 x^3)^2$$

with

$$x = 1 + \sqrt{\frac{T}{T_c i}}$$

Parameters are tabulated for many fluids in the paper of Horstmann (https://doi.org/10.1016/j.fluid.2004.11.002) for the SRK EOS (only)

4.2. General cubics

(continues on next page)

```
model = teqp.make_model(j)
T = 373.1242958476844 # K
rhoL, rhoV = model.superanc_rhoLV(T)
z = np.array([1.0])
pL = rhoL*model.get_R(z)*T*(1.0 + model.get_Ar01(T, rhoL, z))
print('And with SRK and Mathias-Copeman parameters:', pL, 'Pa')
And with SRK and Mathias-Copeman parameters: 101639.22259842217 Pa
```

4.3 Quantum PR

The quantum-corrected Peng-Robinson model of Aasen *et al.* (https://doi.org/10.1063/1.5111364) can be used to account for quantum effects by empirical fits to the Feynman-Hibbs corrections.

The conventional Peng-Robinson approach is used, with an adjusted covolume b_i given by

$$b_i = b_{i,PR}\beta_i(T)$$

with

$$\beta_i(T) = \left(\frac{1 + A_i/(T + B_i)}{1 + A_i/(T_{ci} + B_i)}\right)^3$$

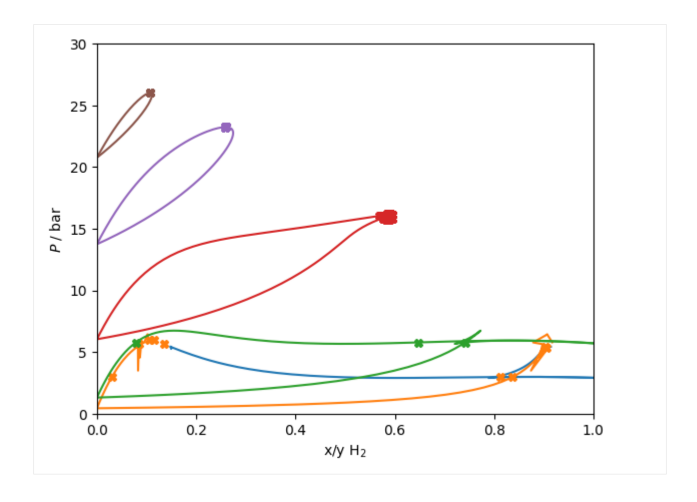
and Twu alpha functions are used to correct the attractive part.

```
[1]: import numpy as np, matplotlib.pyplot as plt, pandas
    import CoolProp.CoolProp as CP
    import teqp
    teqp.__version_
[1]: '0.19.1'
[2]: kij_library = {
         ('H2','Ne'): 0.18,
         ('He', 'H2'): 0.17
    lij_library = {
         ('H2','Ne'): 0.0,
         ('He','H2'): -0.16
    def get_model(names, c_factor=0):
        param_library = {
             'H2': {
                 "Ls": [156.21],
                 "Ms": [-0.0062072],
                 "Ns": [5.047],
                 "As": [3.0696],
                 "Bs": [12.682],
                 "cs / m^3/mol": [c_factor*-3.8139e-6],
                 "Tcrit / K": [33.19],
                 "pcrit / Pa": [12.964e5]
             },
                                                                                  (continues on next page)
```

```
'Ne': {
            "Ls": [0.40453],
            "Ms": [0.95861],
            "Ns": [0.8396],
            "As": [0.4673],
            "Bs": [2.4634],
            "cs / m^3/mol": [c_factor*-2.4665e-6],
            "Tcrit / K": [44.492],
            "pcrit / Pa": [26.79e5]
        },
        'He': {
            "Ls": [0.48558],
            "Ms": [1.7173],
            "Ns": [0.30271],
            "As": [1.4912],
            "Bs": [3.2634],
            "cs / m^3/mol": [c_factor*-3.1791e-6],
            "Tcrit / K": [5.1953],
            "pcrit / Pa": [2.276e5]
    }
    params = [param_library[name] for name in names]
    model = {k: [param[k][0] for param in params] for k in ['Ls','Ms','Ns','As','Bs',
→'cs / m^3/mol','Tcrit / K','pcrit / Pa']}
    if len(names) == 1:
        model['kmat'] = [[0]]
        model['lmat'] = [[0]]
    else:
        kij = kij_library[names]
        model['kmat'] = [[0,kij],[kij,0]]
        lij = lij_library[names]
        model['lmat'] = [[0,lij],[lij,0]]
    j = {
        "kind": "QCPRAasen",
        "model": model
    return teqp.make_model(j), j
model = get_model(('H2','Ne'))[0]
modelH2 = get_model(('H2',))[0]
modelNe = get_model(('Ne',))[0]
def get_traces(T, ipures):
    traces = []
    for ipure in ipures:
        rhovecL0 = np.array([0.0, 0.0])
        rhovecV0 = np.array([0.0, 0.0])
        if ipure == 1:
            rhoL, rhoV = modelNe.superanc_rhoLV(T)
            rhoL, rhoV = modelH2.superanc_rhoLV(T)
        rhovecL0[ipure] = rhoL
        rhovecV0[ipure] = rhoV
        opt = teqp.TVLEOptions();
                                                                           (continues on next page)
```

4.3. Quantum PR 21

```
opt.polish=True;
          opt.integration_order=5; opt.rel_err=1e-10;
          opt.calc_criticality = True;
        opt.crit\_termination=1e-10
        trace = model.trace_VLE_isotherm_binary(T, rhovecL0, rhovecV0, opt)
       traces.append(trace)
    return traces
for T in [24.59, 28.0, 34.66, 39.57, 42.50]:
   if T < 26.0:
       traces = get_traces(T, [0, 1])
    else:
       traces = get_traces(T, [1])
   for trace in traces:
       df = pandas.DataFrame(trace)
        # Plot the VLE solution
       line, = plt.plot(df['xL_0 / mole frac.'], df['pL / Pa']/1e5)
        plt.plot(df['xV_0 / mole frac.'], df['pL / Pa']/1e5, color=line.get_color())
    # Plot the VLLE solution if found
   for soln in model.find_VLLE_T_binary(traces):
        for rhovec in soln['polished']:
           rhovec = np.array(rhovec)
           rhotot = sum(rhovec)
           x = rhovec/rhotot
           p = rhotot*model.get_R(x)*T*(1+model.get_Ar01(T, rhotot, x))
           plt.plot(x[0], p/1e5, 'X', color=line.get_color())
            # print(T, rhovec, x[0], p/1e5, 'bar')
plt.gca().set(xlabel='x/y H$_2$', ylabel='$P$ / bar', xlim=(0,1), ylim=(0,30));
```



4.4 Advanced cubic mixing rules

In the advanced cubic mixing rules, the conventional cubic EOS is taken as the basis for the method (usually Peng-Robinson), but different rules are used for the attractive term a_m . The formulation reads:

$$\frac{a_m}{b_m} = \sum_{i} z_i \frac{a_i}{b_i} + \frac{a_{\text{res}}^{E,\gamma}}{CEoS}$$

where CEoS is a scaling parameter that is in principle linked with the EOS coefficients, but can also be allowed to be an adjustable parameter. The a_i and b_i are the pure fluid values of component i. The z_i are mole fractions. The mixture covolume is given by

$$b_m = \sum_i \sum_j z_i z_j b_{ij}$$

with

$$b_{ij} = \left(\frac{b_i^{1/s} + b_j^{1/s}}{2}\right)^s$$

The heart of the method is the definition of $a_{\rm res}^{E,\gamma}$, the residual contribution (not in the conventional thermodynamic sense) to the excess Helmholtz energy. There are many possible models here, but one that seems to work well is that of Wilson,

for which the expression reads:

$$\frac{a_{\mathrm{res}}^{E,\gamma}}{RT} = -\sum_{i} z_{i} \ln \left(\sum_{j} z_{j} \Omega_{ji}(T) \right) - \sum_{i} z_{i} \ln \left(\frac{\phi_{i}}{z_{i}} \right)$$

with

$$\Omega_{ji} = \frac{v_j}{v_i} \exp(-A_{ij}/T)$$

and

$$\frac{\phi_i}{z_i} = \frac{v_i}{\sum_k z_k v_k}$$

with the $v_i = b_i$. The parameter $A_{ij} \neq A_{ji}$ in general, and is also given temperature dependence, which is also not supposed to be present according to the derivation. Thus, the models for A_{ij} read something like this here:

$$A_{ij} = m_{ij}T + n_{ij}$$

so m is non-dimensional and n has units of temperature.

```
[1]: import numpy, matplotlib.pyplot as plt, numpy as np, pandas
  import teqp
  teqp.__version__
[1]: '0.19.1'
```

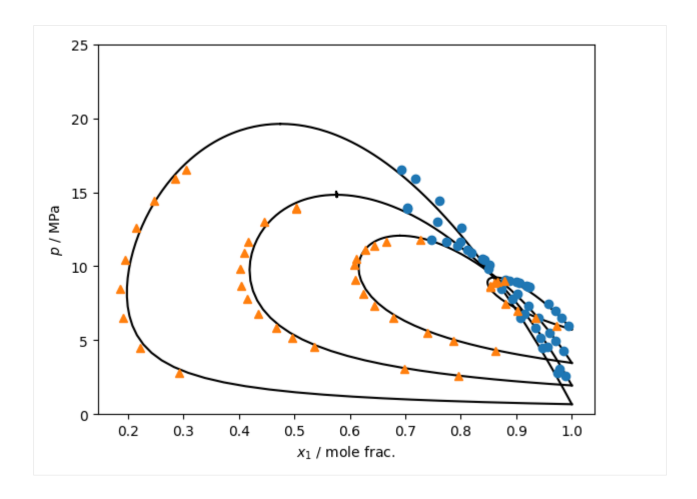
```
[2]: # Four isotherms of experimental data from doi: 10.1016/j.fluid.2016.05.015
    import io, pandas
    dat = pandas.read_csv(io.StringIO("""PointID y1 uy1 x1 ux1 p/bar up T/K
    1 0.0274 0.0007 0.0068 0.0002 59.830 0.053 293.10
    2 0.0664 0.0014 0.0183 0.0004 64.864 0.080 293.10
    3 0.0978 0.0020 0.0298 0.0007 69.772 0.080 293.10
    4 0.1199 0.0024 0.0424 0.0009 74.737 0.080 293.10
    5 0.1219 0.0028 0.1132 0.0023 89.869 0.080 293.10
    6 0.1339 0.0024 0.0995 0.0022 89.198 0.080 293.10
    7 0.1399 0.0026 0.0943 0.0020 88.853 0.080 293.10
    8 0.1461 0.0027 0.0823 0.0019 86.962 0.080 293.10
    9 0.1466 0.0028 0.0778 0.0017 85.942 0.080 293.10
    10 0.1466 0.0028 0.0772 0.0016 85.868 0.080 293.10
    1 0.1378 0.0027 0.0159 0.0004 42.667 0.051 273.08
    2 0.2143 0.0038 0.0297 0.0007 49.547 0.051 273.08
    3 0.2612 0.0043 0.0411 0.0009 55.238 0.051 273.08
    4 0.3209 0.0049 0.0609 0.0013 65.069 0.088 273.08
    5 0.3554 0.0051 0.0786 0.0016 73.395 0.088 273.08
    6 0.3758 0.0052 0.0978 0.0019 81.061 0.088 273.08
    7 0.3903 0.0053 0.1190 0.0023 90.706 0.088 273.08
    8 0.3914 0.0053 0.1477 0.0028 100.966 0.088 273.08
    9 0.3879 0.0053 0.1614 0.0030 104.806 0.088 273.08
    10 0.3724 0.0052 0.1875 0.0033 110.846 0.088 273.08
    11 0.3550 0.0051 0.2068 0.0036 114.105 0.088 273.08
    12 0.2727 0.0044 0.2531 0.0041 118.020 0.088 273.08
    13 0.3343 0.0049 0.2268 0.0038 116.295 0.088 273.08
    1 0.2048 0.0038 0.0106 0.0003 25.754 0.050 253.05
    2 0.3019 0.0049 0.0217 0.0005 30.479 0.050 253.05
    3 0.4638 0.0056 0.0436 0.0010 45.352 0.050 253.05
    4 0.5319 0.0056 0.0647 0.0014 58.188 0.050 253.05
```

24 Chapter 4. Models

(continues on next page)

```
(continued from previous page)
5 0.5854 0.0054 0.1077 0.0021 78.315 0.084 253.05
6 0.5979 0.0054 0.1497 0.0028 98.276 0.084 253.05
7 0.5898 0.0054 0.1801 0.0032 109.241 0.084 253.05
8 0.5042 0.0057 0.0570 0.0012 51.343 0.084 253.05
9 0.5644 0.0055 0.0861 0.0017 67.594 0.084 253.05
10 0.5949 0.0054 0.1267 0.0024 86.883 0.084 253.05
11 0.5826 0.0054 0.2015 0.0035 116.614 0.084 253.05
12 0.5537 0.0055 0.2431 0.0040 129.873 0.084 253.05
13 0.4973 0.0055 0.2971 0.0046 139.161 0.084 253.05
14 0.4971 0.0055 0.2972 0.0046 139.261 0.084 253.05
1 0.7076 0.0050 0.0257 0.0006 27.983 0.056 223.10
2 0.7774 0.0041 0.0522 0.0011 44.918 0.056 223.10
3 0.8077 0.0036 0.0930 0.0019 64.906 0.081 223.10
4 0.8131 0.0035 0.1261 0.0024 84.799 0.081 223.10
5 0.8057 0.0035 0.1584 0.0029 104.410 0.081 223.10
6 0.7843 0.0038 0.1982 0.0035 125.782 0.081 223.10
7 0.7533 0.0041 0.2380 0.0040 144.287 0.081 223.10
8 0.7150 0.0045 0.2813 0.0044 159.015 0.081 223.10
9 0.6942 0.0047 0.3064 0.0047 165.347 0.081 223.10
"""), sep='\s+', engine='python')
j = {
    "kind": "advancedPRaEres",
    "model": {
```

```
[3]: # Model from Lasala, FPE, 2016: https://doi.org/10.1016/j.fluid.2016.05.015
            "Tcrit / K": [304.21, 126.19],
            "pcrit / Pa": [7.383e6, 3395800.0],
            "alphas": [{"type": "PR78", "acentric": 0.22394}, {"type": "PR78", "acentric": __
     \rightarrow 0.037211,
            "aresmodel": {"type": "Wilson", "m": [[0.0, -3.4768], [3.5332, 0.0]], "n": [[0.
     \rightarrow0, 825], [-585, 0.0]]},
            "options": {"s": 2.0, "brule": "Quadratic", "CEoS": -0.52398}
    }
    model = teqp.make_model(j)
    for T in [223.15, 253.05, 273.08, 293.1]:
        ipure = 0
        [rhoL0, rhoV0] = model.superanc_rhoLV(T, ipure)
        rhovecL0 = np.array([0.0, 0.0]); rhovecL0[ipure] = rhoL0
        rhovecV0 = np.array([0.0, 0.0]); rhovecV0[ipure] = rhoV0
        J = model.trace_VLE_isotherm_binary(T, rhovecL0, rhovecV0)
        df = pandas.DataFrame(J)
        plt.plot(df['xL_0 / mole frac.'], df['pL / Pa']/1e6,'k')
        plt.plot(df['xV_0 / mole frac.'], df['pV / Pa']/1e6,'k')
    plt.plot(1-dat['x1'], dat['p/bar']/10, 'o')
    plt.plot(1-dat['y1'], dat['p/bar']/10, '^')
    plt.gca().set(xlabel='$x_1$ / mole frac.', ylabel='$p$ / MPa', ylim=(0, 25))
    plt.show()
```



4.5 RK-PR

The EOS can be given as

$$\alpha^{r} = \psi^{(-)} - \frac{a_m}{RT} \psi^{(+)}$$
$$\psi^{(-)} = -\ln(1 - b_m \rho)$$
$$\psi^{(+)} = \frac{\ln\left(\frac{\Delta_1 b_m \rho + 1}{\Delta_2 b_m \rho + 1}\right)}{b_m(\Delta_1 - \Delta_2)}$$

with the EOS fixed constants of

$$\Delta_1 = \sum_i x_i \delta_{1,i}$$

$$\Delta_2 = \frac{1 - \Delta_1}{1 + \Delta_1}$$

The attractive term goes like

$$a_i = a_{c,i} \left(\frac{2}{3 + T/T_{c,i}} \right)^{k_i}$$

with quadratic mixing rules

$$a_m = \sum_{i} \sum_{j} x_i x_j (1 - k_{ij}) \sqrt{a_i(T)a_j(T)}$$

And the covolume also gets quadratic mixing rules

$$b_m = \sum_{i} \sum_{j} x_i x_j (1 - l_{ij})(b_i + b_j)/2$$

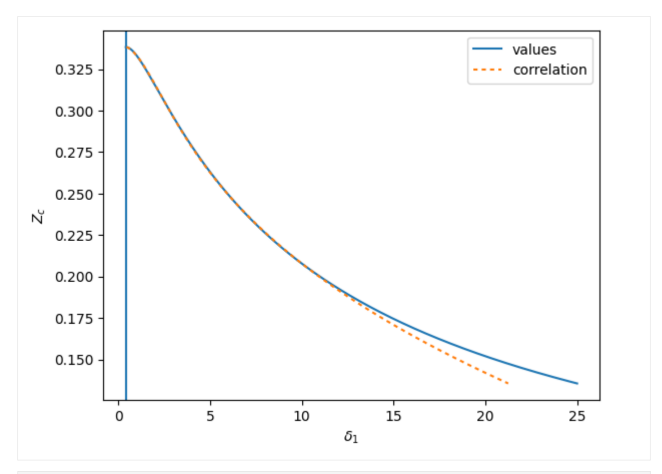
Thus, to implement the RK-PR model in predictive mode, the following steps are required:

- 1. Obtain the critical parameters Tc, pc
- 2. Solve for delta_1 from the experimental critical compressibility factor, begin with the values from the correlation
- 3. Solve for k by fixing the pressure at the T=0.7Tc. In the case (e.g, CO:math:`_2`) that Tt < 0.7Tc, use instead Tr=Tt/Tc

It may be necessary to adjust the values of $\delta_{1,i}$ and k_i for an individual component to better match the behavior of more polar components.

```
[1]: import numpy as np
    import scipy.optimize
    import matplotlib.pyplot as plt
    import teqp, numpy as np
    import CoolProp.CoolProp as CP
    import pandas
    def delta1_correlation(Zc):
         # Eq. B.4 of Cismondi FPE 2005
        d1 = 0.428363
        d2 = 18.496215
        d3 = 0.338426
        d4 = 0.660000
        d5 = 789.723105
        d6 = 2.512392
         return d1 + d2*(d3-Zc)**d4 + d5*(d3-Zc)**d6
    def Zc_delta1(delta1):
         # Eqs. B.1 to B.3 of Cismondi FPE 2005
        d1 = (1+delta1**2)/(1+delta1)
        y = 1 + (2*(1+delta1))**(1/3) + (4/(1+delta1))**(1/3)
         return y/(3*y + d1 - 1)
    DELTA1 = np.linspace(np.sqrt(2)-1, 25, 1000)
    ZZ = Zc_delta1(DELTA1)
    plt.plot(DELTA1, ZZ, label='values')
    DELTA1back = delta1_correlation(ZZ)
    plt.axvline(np.sqrt(2)-1)
    plt.plot(DELTA1back, ZZ, dashes=[2,2], label='correlation')
    plt.gca().set(ylabel='$Z_c$', xlabel='$\delta_1$')
    plt.legend(loc='best')
    plt.show()
     # for Zc in np.linspace(0.2, 0.3383, 1000):
          resid = lambda x: Zc_delta1(x) - Zc
           # print(resid(delta1_correlation(Zc)))
          print(Zc, scipy.optimize.newton(resid, delta1_correlation(Zc)), delta1_
     \hookrightarrow correlation (Zc))
```

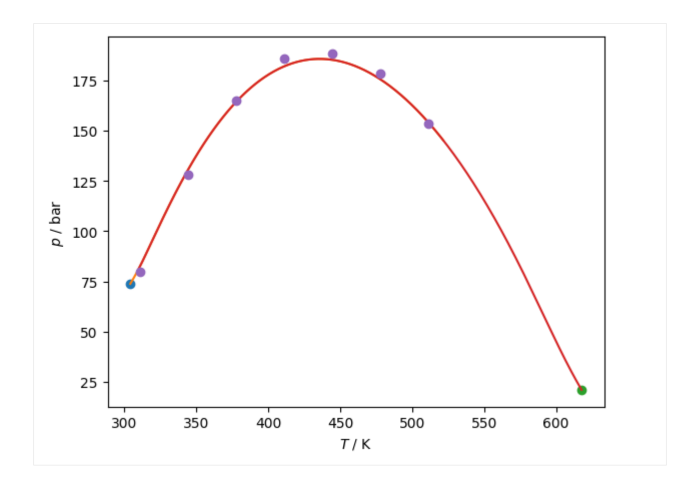
4.5. RK-PR 27



```
[2]: names = ['CO2', 'n-Decane']
    R = 8.31446261815324
    Tc = np.array([CP.PropsSI(k,"Tcrit") for k in names])
    pc = np.array([CP.PropsSI(k,"pcrit") for k in names])
    rhoc = np.array([CP.PropsSI(k,"rhomolar_critical") for k in names])
    Zcexp = pc/(rhoc*R*Tc)
     # Use a rescaled Zc to obtain delta_1
    Zc = 1.168*Zcexp
    delta_1 = [scipy.optimize.newton(lambda x: Zc_delta1(x)-Zc_, delta1_correlation(Zc_))_
     →for Zc_ in Zc]
    def solve_for_k(i, p_target, Tr):
         The value of k for the i-th component is based on getting
        the right vapor pressure, so a rootfinding routing is
        used to obtain these values
         11 11 11
        def objective(k):
             j = {
                 "kind": "RKPRCismondi2005",
                 "model": {
                     "delta_1": [delta_1[i]],
                     "Tcrit / K": [Tc[i]],
                                                                                (continues on next page)
```

```
"pcrit / Pa": [pc[i]],
                "k": [k],
                "kmat": [[0.0]],
                "lmat": [[0.0]],
       model = teqp.make_model(j)
        T = Tr*Tc[i]
        z = np.array([1.0])
        a, b = model.get_ab(T, z)
       anc = teqp.build_ancillaries(model, Tc[i], rhoc[i], 150)
       rhoL, rhoV = model.pure_VLE_T(T, anc.rhoL(T), anc.rhoV(T), 10)
        p = T*R*rhoL*(1+model.get\_Ar01(T, rhoL, z))
       return p-p_target
   return scipy.optimize.newton(objective, 2.1)
Tr = 0.7
i = 1
k_C10 = solve_for_k(i, CP.PropsSI('P', 'T', Tr*Tc[i], 'Q', 0, names[i]), Tr)
model = teqp.make_model({
    "kind": "RKPRCismondi2005",
    "model": {
        "delta_1": delta_1,
       "Tcrit / K": Tc.tolist(),
       "pcrit / Pa": pc.tolist(),
        "k": [2.23854, k_C10],
        "kmat": [[0,0],[0,0]],
        "lmat": [[0,0],[0,0]],
    }
})
# Start at both pures
for ipure in [0, 1]:
   Tc, rhoc = model.solve_pure_critical(300, 5000, {"alternative_pure_index":ipure,
→"alternative_length": 2})
   z = np.array([0.0, 0.0]); z[ipure] = 1.0
   pc = Tc*R*rhoc*(1+model.get_Ar01(Tc, rhoc, z))
   plt.plot(Tc, pc/1e5, 'o')
   opt = teap.TCABOptions(); opt.polish=True; opt.verbosity=100; opt.integration_
→order=5; opt.rel_err=1e-10; opt.abs_err=1e-10
   trace = model.trace_critical_arclength_binary(Tc, z*rhoc, options=opt)
   df = pandas.DataFrame(trace)
   plt.plot(df['T / K'], df['p / Pa']/1e5)
# Overlay the data from Reamer and Sage, Cismondi additional data points not present
→in Reamer and Sage
Tc_K = [310.928, 344.261, 377.594, 410.928, 444.261, 477.594, 510.928]
pc_kPa = np.array([7997.92, 12824.25, 16492.26, 18560.69, 18836.48, 17836.74, 15333.
→ 941)
plt.plot(Tc_K, pc_kPa/1e2, 'o')
plt.gca().set(xlabel='$T$ / K', ylabel='$p$ / bar');
```

4.5. RK-PR 29



4.6 Cubic Plus Association (CPA)

The combination of a cubic EOS with association with the association term. The sum of the terms goes like:

$$\alpha^{\rm r} = \alpha^{\rm r}_{\rm cub} + \alpha^{\rm r}_{\rm assoc}$$

4.6.1 Cubic part

The residual contribution to α is expressed as the sum :

$$\alpha_{\mathrm{cub,rep}}^{\mathrm{r}} + \alpha_{\mathrm{cub,att}}^{\mathrm{r}}$$

where the cubic parts come from

The repulsive part of the cubic EOS contribution:

$$\alpha_{\mathrm{cub},\mathrm{rep}}^{\mathrm{r}} = -\ln(1 - b_{\mathrm{mix}}\rho)$$

The attractive part of the cubic EOS contribution:

$$\alpha_{\mathrm{cub,att}}^{\mathrm{r}} = -\frac{a_{\mathrm{mix}}}{RT} \frac{\ln\left(\frac{\Delta_{1}b_{\mathrm{mix}}\rho + 1}{\Delta_{2}b_{\mathrm{mix}}\rho + 1}\right)}{b_{\mathrm{mix}} \cdot (\Delta_{1} - \Delta_{2})}$$

with the coefficients depending on the cubic type:

SRK:
$$\Delta_1 = 1$$
, $\Delta_2 = 0$

PR:
$$\Delta_1 = 1 + \sqrt{2}$$
, $\Delta_2 = 1 - \sqrt{2}$

The mixture models used for the a_{mix} and b_{mix} are the classical ones:

$$a_{\text{mix}} = \sum_{i} \sum_{j} x_i x_j (1 - k_{ij}) a_{ij}(T)$$

with x the mole fraction, k_{ij} a weighting parameter

$$a_{ij}(T) = \sqrt{a_i a_j}$$

and

$$a_i(T) = a_{0i} \left[1 + c_{1i} (1 - \sqrt{T/T_{\text{crit},i}}) \right]^2$$

and for b:

$$b_{\text{mix}} = \sum_{i} x_i b_i$$

so there are three cubic parameters per fluid that need to be obtained though fitting: b_i , a_{0i} , c_{1i} . The value of a_{ij} depends on temperature while b_{mix} does not.

4.6.2 Association part

For the association, one must have a solid understanding of the association approach that is being applied. To this end, a short discussion of the general approach is required.

1.

4.7 LKP (Lee-Kesler-Plöcker)

The LKP model is a sort of hybrid between corresponding states and multiparameter EOS, simple EOS are developed for a reference fluid, and a simple fluid, and the acentric factor of the mixture is used to weight the two.

The reduced residual Helmholtz energy for the mixture is evaluated from

$$\alpha^{\rm r} = \left(1 - \frac{\omega_{\rm mix}}{\omega_{\rm ref}}\right) \alpha_{\rm simple}^{\rm r} + \frac{\omega_{\rm mix}}{\omega_{\rm ref}} \alpha_{\rm ref}^{\rm r}$$

where the contributions are each of the form

$$\alpha_X^{\rm r}(\tau,\delta) = B\left(\frac{\delta}{Z_c}\right) + \frac{C}{2}\left(\frac{\delta}{Z_c}\right)^2 + \frac{D}{5}\left(\frac{\delta}{Z_c}\right)^5 - \frac{c_4\tau^3}{2\gamma}\left(\gamma\left(\frac{\delta}{Z_c}\right)^2 + \beta + 1\right)\exp\left(-\gamma\left(\frac{\delta}{Z_c}\right)^2\right) + \frac{c_4\tau^3}{2\gamma}(\beta + 1)$$

where X is one of simple or reference (abbreviation: ref) with the matching sets of coefficients taken from this table:

var	simple	reference	
b_1	0.1181193	0.2026579	
b_2	0.265728	0.331511	
b_3	0.154790	0.276550e-1	
b_4	0.303230e-1	0.203488	
c_1	0.236744e-1	0.313385e-1	
c_2	0.186984e-1	0.503618e-1	
c_3	0	0.169010e-1	
c_4	0.427240e-1	0.41577e-1	
d_1	0.155428e-4	0.487360e-4	
d_2	0.623689e-4	0.740336e-5	
β	0.653920	1.226	
γ	0.601670e-1	0.03754	
ω	0.0	0.3978	

The terms in the contributions are given by:

$$B = b_1 - b_2 \tau - b_3 \tau^2 - b_4 \tau^3$$

$$C = c_1 - c_2 \tau + c_3 \tau^3$$

$$D = d_1 + d_2 \tau$$

For density, the reduced density δ is defined by

$$\delta = \frac{\rho}{\rho_{\rm red}} = v_{\rm c,mix} \rho$$

in which the reducing density is the reciprocal of the pseudo-critical volume obtained from

$$v_{\rm c,mix} = \sum_{i=1}^{N-1} \sum_{j=i+1}^{N} x_i x_j v_{ij}$$

$$v_{c,ij} = \frac{1}{8} (v_{c,i}^{1/3} + v_{c,j}^{1/3})^3$$

and the critical volumes are estimated from

$$v_{c,i} = (0.2905 - 0.085\omega_i) \frac{RT_{c,i}}{p_{c,i}}$$

For temperature, the reciprocal reduced density is defined by

$$\tau = \frac{T_{\rm c,mix}}{T}$$

with

$$T_{\text{c,mix}} = \frac{1}{v_{c,mix}^{\eta}} \sum_{i=1}^{N-1} \sum_{j=i+1}^{N} x_i x_j v_{c,ij}^{\eta} T_{c,ij}$$

with $\eta = 0.25$ and

$$T_{c,ij} = k_{ij} \sqrt{T_{c,i} T_{c,j}}$$

Note: the default interaction parameter k_{ij} is therefore 1, rather than 0 in the case of SAFT and cubic models.

Finally the parameter Z_c is defined by

$$Z_c = 0.2905 - 0.085\omega_{\text{mix}}$$

with the mixture acentric factor defined by

$$\omega_{\mathrm{mix}} = \sum_{i} x_i \omega_i$$

```
[1]: import teqp, numpy as np
spec = {
    "Tcrit / K": [190.564, 126.192],
    "pcrit / Pa": [4.5992e6, 3.3958e6],
    "acentric": [0.011, 0.037],
    "R / J/mol/K": 8.3144598,
    "kmat": [[1.0, 0.977], [0.977, 1.0]]
}
model = teqp.make_model({'kind': 'LKP', 'model': spec}, validate=True)
```

```
[2]: # A little sanity check, with the check value from TREND
    expected = -0.18568096994998817
    diff = abs(model.get_Ar00(300, 8000.1, np.array([0.8, 0.2])) - expected)
    assert(diff < 1e-13)</pre>
```

4.8 Model Potentials

These EOS for model potentials are useful for understanding theory, and capture some (but perhaps not all) of the physics of "real" fluids.

```
[1]: import teqp
    teqp.__version__
[1]: '0.19.1'
```

4.8.1 Square-well

The potential is defined by

$$V(r) = \begin{cases} \infty & r < \sigma \\ -\varepsilon & \sigma < r < \lambda \sigma \\ 0 & r > \lambda \sigma \end{cases}$$

from which an EOS can be developed by correlating results from molecular simulation. The EOS is from:

Rodolfo Espíndola-Heredia, Fernando del Río and Anatol Malijevsky Optimized equation of the state of the square-well fluid of variable range based on a fourth-order free-energy expansion J. Chem. Phys. 130, 024509 (2009); https://doi.org/10.1063/1.3054361

```
[2]: model = teqp.make_model({
    "kind": "SW_EspindolaHeredia2009",
    "model": {
        "lambda": 1.3
     }
})
```

4.8. Model Potentials 33

4.8.2 EXP-6

```
[3]: model = teqp.make_model({
    "kind": "EXP6_Kataoka1992",
    "model": {
        "alpha": 12
     }
})
```

4.8.3 Lennard-Jones Fluid

The Lennard-Jones potential is given by

$$V(r) = 4\varepsilon \left((\sigma/r)^{12} - (\sigma/r)^6 \right)$$

and EOS are available from many authors. teqp includes the EOS from Thol, Kolafa-Nezbeda, and Johnson.

4.8.4 Two-Center Lennard-Jones Fluid

```
[5]: model = teqp.make_model({
      'kind': '2CLJF-Dipole',
       'model': {
           "author": "2CLJF_Lisal",
           'L^*': 0.5,
           '(mu^*)^2': 0.1
    })
    print(model.solve_pure_critical(1.3, 0.3))
    model = teqp.make_model({
      'kind': '2CLJF-Quadrupole',
       'model': {
          "author": "2CLJF_Lisal",
          'L^*': 0.5,
           '(0^*)^2': 0.1
    })
    print (model.solve_pure_critical(1.3, 0.3))
    (2.8282972062188056, 0.2005046666634018)
    (2.832574303561834, 0.2003194655463274)
```

4.9 PC-SAFT

The PC-SAFT implementation in teqp is based on the implementation of Gross and Sadowski (https://doi.org/10.1021/ie0003887), with the typo from their paper fixed. It does NOT include the association contribution, only the dispersive contributions.

The model in teqp requires the user to specify the values of sigma, epsilon/kB, and m for each substance. A very few substances are hardcoded in teqp, for testing purposes.

The Python class is here: PCSAFTEOS

```
[1]: import teqp
    import numpy as np
    teqp.__version__
[1]: '0.19.1'
[2]: TeXkey = 'Gross-IECR-2001'
    ms = [1.0, 1.6069, 2.0020]
    eoverk = [150.03, 191.42, 208.11]
    sigmas = [3.7039, 3.5206, 3.6184]
    coeffs = []
    for i in range(len(ms)):
        c = tegp.SAFTCoeffs()
        c.m = ms[i]
        c.epsilon_over_k = eoverk[i]
        c.sigma_Angstrom = sigmas[i]
        coeffs.append(c)
    model = teqp.PCSAFTEOS(coeffs)
[3]: # Here are some rudimentary timing results
    T = 300.0
    rhovec = np.array([3.0, 4.0, 5.0])
    rho = rhovec.sum()
    x = rhovec/np.sum(rhovec)
    %timeit model.get_fugacity_coefficients(T,rhovec)
    %timeit (-1.0) *model.get_Ar20(T, rho, x)
    %timeit model.get_partial_molar_volumes(T, rhovec)
    4.28 \mu s \pm 7.74 ns per loop (mean \pm std. dev. of 7 runs, 100,000 loops each)
    4.3 \mu s ± 49.7 ns per loop (mean ± std. dev. of 7 runs, 100,000 loops each)
```

The model parameters can be queried:

16.9 $\mu s \pm 267$ ns per loop (mean \pm std. dev. of 7 runs, 100,000 loops each)

4.9. PC-SAFT 35

4.9.1 Adjusting k ij

Fine-tuned values of k_{ij} can be provided when instantiating the model. A complete matrix of all the k_{ij} values must be provided. This allows for asymmetric mixing models in which $k_{ij} \neq k_{ji}$.

4.9.2 Superancillary

The superancillary equation for PC-SAFT has been developed, and is much more involved than that of the cubic EOS. As a consequence, the superancillary equation has been provided as a separate package rather than integrating it into to teqp to minimize the binary size of teqp. It can be installed from PYPI with: pip install PCSAFTsuperanc

The scaling in the superancillaries uses reduced variables:

$$\tilde{T} = T/(\epsilon/k_{\rm B})$$
$$\tilde{\rho} = \rho_{\rm N}\sigma^3$$

where ρ_N is the number density, and the other parameters are from the PC-SAFT model

```
[7]: import PCSAFTsuperanc
    sigma_m = 3e-10 \# [meter]
    e_{over_k} = 150.0 \# [K]
    m = 5
    # The saturation temperature
    T = 300
    [Ttilde_crit, Ttilde_min] = PCSAFTsuperanc.get_Ttilde_crit_min(m=m)
    print('Ttilde crit:', Ttilde_crit)
    # Get the scaled densities for liquid and vapor phases
    [tilderhoL, tilderhoV] = PCSAFTsuperanc.PCSAFTsuperanc_rhoLV(Ttilde=T/e_over_k, m=m)
     # Convert back to molar densities
    N_A = PCSAFTsuperanc.N_A # The value of Avogadro's constant used in superancillaries
    rhoL, rhoV = [tilderho/(N_A*sigma_m**3) for tilderho in [tilderhoL, tilderhoV]]
    # As a sanity check, confirm that we got the same pressure in both phases
    c = teqp.SAFTCoeffs()
    c.sigma_Angstrom = sigma_m*1e10
    c.epsilon_over_k = e_over_k
    c.m = m
    model = teqp.PCSAFTEOS([c])
    z = np.array([1.0])
    pL = rhoL*model.get_R(z)*T*(1+model.get_Ar01(T, rhoL, z))
                                                                                (continues on next page)
```

```
pV = rhoV*model.get_R(z)*T*(1+model.get_Ar01(T, rhoV, z))
print('Pressures are:', pL, pV, 'Pa')

Ttilde crit: 2.648680568587752
Pressures are: 227809.12314460654 227809.12314409122 Pa
```

4.9.3 Maximum density

The maximum number density allowed by the EOS is defined based on the packing fraction. To get a molar density, divide by Avogadro's number. The function is conveniently exposed in Python:

```
[8]: max_rhoN = teqp.PCSAFTEOS(coeffs).max_rhoN(130.0, np.array([0.3, 0.3, 0.4]))
    display(max_rhoN)
    max_rhoN/6.022e23 # the maximum molar density in mol/m^3
    1.9139171771761775e+28
[8]: 31782.085306811314
```

4.9.4 Polar contributions

As of teqp version 0.15, quadrupolar and dipolar contributions have been added to the hard chain plus dispersion model which is referred to conventionally as PC-SAFT. The definitions of the reduced dipolar and quadrupolar parameters are not well documented, so they are given here. The work of Stoll, Vrabec, and Hasse (https://doi.org/10.1063/1.1623475) clearly describes the formulation of the star-scaling.

In SI units, the reduced squared dipole moment is defined by

$$(\mu^*)_{\text{conventional}}^2 = \frac{(\mu[Cm])^2}{4\pi\epsilon_0(\varepsilon[J])(\sigma[m])^3}$$

$$(Q^*)_{\rm conventional}^2 = \frac{(\mu[Cm])^2}{4\pi\epsilon_0(\varepsilon[J])(\sigma[m])^5}$$

In the PC-SAFT formulation, the only difference is the addition of dividing the denominator by the number of segments m

$$(\mu^*)^2 = \frac{(\mu[Cm])^2}{4\pi\epsilon_0 m (\varepsilon/k_{\rm B}[K]) k_B (\sigma[m])^3}$$

$$(Q^*)^2 = \frac{(Q[Cm^2])^2}{4\pi\epsilon_0 m(\varepsilon/k_B[K])k_B(\sigma[m])^5}$$

The unit conversions are obtained from

$$(\sigma[m])=(10^{-10}m/A)(\sigma[A])$$

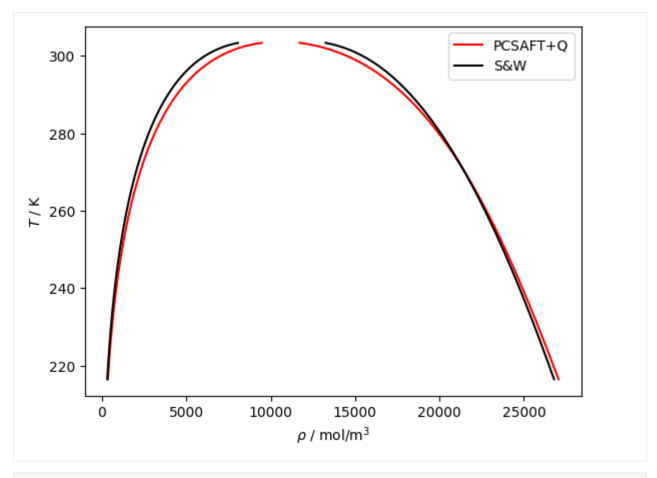
$$(\mu[Cm]) = (3.33564 \times 10^{-30} Cm/D)(\mu[D])$$

and $\epsilon_0 = 8.85419e - 12 \text{ C}^2 \text{ N}^{-1} \text{ m}^{-2}$ is the permittivity of vacuum.

```
[9]: # CO2 with quadrupolar contributions
j = {
    'kind': 'PCSAFT',
    'model': {
    (continues on next page)
```

4.9. PC-SAFT 37

```
'coeffs': [{
            'name': 'CO2',
             'BibTeXKey': 'Gross-AICHEJ',
             'm': 1.5131,
             'sigma_Angstrom': 3.1869,
             'epsilon_over_k': 169.33,
             '(Q^*)^2': 1.26, # modified from the values in Gross and Vrabec since.
→the base model is different
             'nQ': 1
        } ]
    }
}
model = teqp.make_model(j)
Tc, rhoc = model.solve_pure_critical(300, 11000)
T = Tc*0.999
rhoL_, rhoV_ = model.extrapolate_from_critical(Tc, rhoc, T)
rhoL, rhoV = model.pure_VLE_T(T, rhoL_, rhoV_, 10)
import CoolProp.CoolProp as CP
import matplotlib.pyplot as plt
import pandas
0 = []
for T_ in np.linspace(T, 215, 1000):
    rhoL, rhoV = model.pure_VLE_T(T_, rhoL, rhoV, 10)
    try:
        o.append({
          'T': T_, 'rhoL': rhoL, 'rhoV': rhoV,
          'rhoLSW': CP.PropsSI('Dmolar', 'T', T_, 'Q', 0, 'CO2'),
          'rhoVSW': CP.PropsSI('Dmolar','T',T_,'Q',1,'CO2')
        })
    except:
        pass
df = pandas.DataFrame(o)
plt.plot(df['rhoL'], df['T'], 'r', label='PCSAFT+Q')
plt.plot(df['rhoV'], df['T'], 'r')
plt.plot(df['rhoLSW'], df['T'], 'k', label='S&W')
plt.plot(df['rhoVSW'], df['T'], 'k')
plt.legend()
plt.gca().set(xlabel=r'$\rho$ / mol/m$^3$', ylabel='$T$ / K')
plt.show()
```



```
[10]: # Acetone with dipolar contributions
          'kind': 'PCSAFT',
          'model': {
              'coeffs': [{
                  'name': 'acetone',
                   'BibTeXKey': 'Gross-IECR',
                   'm': 2.7447,
                   'sigma_Angstrom': 3.2742,
                   'epsilon_over_k': 232.99,
                   '(mu^*)^2': 1.9, # modified from the values in Gross and Vrabec since.
      →the base model is different
                   'nmu': 1
              } ]
         }
     model = teqp.make_model(j)
     Tc, rhoc = model.solve_pure_critical(300, 11000)
     T = Tc*0.999
     rhoL_, rhoV_ = model.extrapolate_from_critical(Tc, rhoc, T)
     rhoL, rhoV = model.pure_VLE_T(T, rhoL_, rhoV_, 10)
     import CoolProp.CoolProp as CP
     import matplotlib.pyplot as plt
                                                                                 (continues on next page)
```

4.9. PC-SAFT 39

```
import pandas
0 = []
for T_ in np.linspace(T, 215, 1000):
    rhoL, rhoV = model.pure_VLE_T(T_, rhoL, rhoV, 10)
    try:
        o.append({
          'T': T_, 'rhoL': rhoL, 'rhoV': rhoV, 'rhoLSW': CP.PropsSI('Dmolar','T',T_,'Q',0,'acetone'),
          'rhoVSW': CP.PropsSI('Dmolar','T',T_,'Q',1,'acetone')
        })
    except:
        pass
df = pandas.DataFrame(o)
plt.plot(df['rhoL'], df['T'], 'r', label='PCSAFT+D')
plt.plot(df['rhoV'], df['T'], 'r')
plt.plot(df['rhoLSW'], df['T'], 'k', label='S&W')
plt.plot(df['rhoVSW'], df['T'], 'k')
plt.legend()
plt.gca().set(xlabel=r'$\rho$ / mol/m$^3$', ylabel='$T$ / K')
plt.show()
                                                                        PCSAFT+D
    500
                                                                        S&W
    450
     400
    350
     300
    250
             0
                    2000
                             4000
                                      6000
                                               8000
                                                        10000
                                                                 12000
                                                                          14000
                                          \rho / mol/m<sup>3</sup>
```

4.10 SAFT-VR-Mie

The SAFT-VR-Mie EOS of Lafitte et al. (https://doi.org/10.1063/1.4819786) is based on the use of a Mie potential of the form

$$u(r) = C\epsilon \left((\sigma/r)^{\lambda_r} - (\sigma/r)^{\lambda_a} \right)$$

with

$$C = \frac{\lambda_r}{\lambda_r - \lambda_a} \left(\frac{\lambda_r}{\lambda_a}\right)^{\lambda_a/(\lambda_r - \lambda_a)}$$

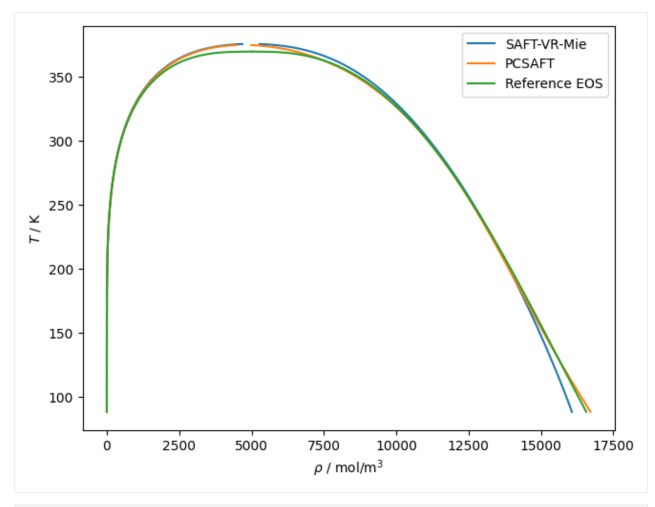
which allows for a better representation of thermodynamic properties in general, but not always.

```
[1]: import teqp
    teqp.__version_
[1]: '0.19.1'
[2]: import numpy as np
    import pandas
    import matplotlib.pyplot as plt
    import CoolProp.CoolProp as CP
    import scipy.integrate
[3]: # Show two ways to instantiate a SAFT-VR-Mie model, the
    # first by providing the coefficients, and the second
    # by providing the name of the species. Only a very small
    # number of molecules are provided for testing, you should
    # plan on providing your own parameters.
    # Show that both give the same result for the residual pressure
    z = np.array([1.0])
    model = teqp.make_model({
        "kind": 'SAFT-VR-Mie',
         "model": {
             "coeffs": [{
                 "name": "Ethane",
                 "BibTeXKey": "Lafitte",
                "m": 1.4373,
                 "epsilon_over_k": 206.12, # [K]
                 "sigma_m": 3.7257e-10,
                 "lambda_r": 12.4,
                 "lambda_a": 6.0
            } ]
        }
    display(model.get_Ar01(300, 300, z))
    model = teqp.make_model({
        "kind": 'SAFT-VR-Mie',
        "model": {
            "names": ["Ethane"]
    })
    display(model.get_Ar01(300, 300, z))
```

4.10. SAFT-VR-Mie 41

```
-0.04926724350863724
-0.04926724350863724
```

```
[4]: # Here is an example of using teqp to trace VLE for propane
    # with the default parameters of PC-SAFT and SAFT-VR-Mie
    # models
    for kind in ['SAFT-VR-Mie', 'PCSAFT']:
        j = {
            "kind": kind,
            "model": {
                 "names": ["Propane"]
        }
        model = teqp.make_model(j)
        z = np.array([1.0])
        Tc, rhoc = model.solve_pure_critical(300, 10000)
        # Extrapolate away from the critical point
        Ti = Tc*0.9997
        rhoL, rhoV = model.extrapolate_from_critical(Tc, rhoc, Ti)
        0 = []
        T = Ti
        while T > 88:
            rhoL, rhoV = model.pure_VLE_T(T, rhoL, rhoV, 10)
            o.append({'rhoL': rhoL, 'rhoV': rhoV, 'T': T})
        df = pandas.DataFrame(o)
        line, = plt.plot(df['rhoL'], df['T'], label=kind)
        plt.plot(df['rhoV'], df['T'], color=line.get_color())
    # From the reference EOS of Lemmon et al. via CoolProp
    name = 'Propane'
    Tc = CP.PropsSI(name, 'Tcrit')
    Ts = np.linspace(88, Tc, 1000)
    rhoL = CP.PropsSI('Dmolar', 'T', Ts, 'Q', 0, name)
    rhoV = CP.PropsSI('Dmolar', 'T', Ts, 'Q', 1, name)
    line, = plt.plot(rhoL, Ts, label='Reference EOS')
    plt.plot(rhoV, Ts, line.get_color())
    plt.gca().set(xlabel=r'$\rho$ / mol/m$^3$', ylabel=r'$T$ / K')
    plt.legend()
    plt.tight_layout(pad=0.2)
    plt.savefig('SAFTVRMIE_PCSAFT.pdf')
    plt.show()
```



```
[6]: # Checking the effective hardness of interaction,
# the neff parameter defined in https://doi.org/10.1063/5.0007583
# SAFT-VR-Mie comes closest to the right behavior
modelVR = teqp.make_model({
        "kind": 'SAFT-VR-Mie',
        "model": { "names": ["Methane"] }
})
(continues on next page)
```

4.10. SAFT-VR-Mie 43

5

10¹

```
(continued from previous page)
modelPCSAFT = teqp.make_model({
        "kind": 'PCSAFT',
        "model": { "names": ["Methane"] }
})
modelMF = teqp.build_multifluid_model(["Methane"], teqp.get_datapath())
for model, label in [(modelVR, 'SAFT-VR-Mie'),
                      (modelPCSAFT, 'PC-SAFT'),
                      (modelMF, 'reference EOS')]:
    z = np.array([1.0])
    rho = 1e-5
    T = np.geomspace(8, 10000, 10000)
    neff = []
    for T_ in T:
        neff.append(model.get_neff(T_, rho, z))
    plt.plot(T, neff, label=label)
plt.xscale('log')
plt.ylim(0, 30)
plt.gca().set(xlabel=r'$T$ / K', ylabel=r'$n_{\rm eff}$')
plt.legend()
plt.show()
    30
                SAFT-VR-Mie
                PC-SAFT
                reference EOS
    25
    20
 7eff
    15
    10
```

T/K

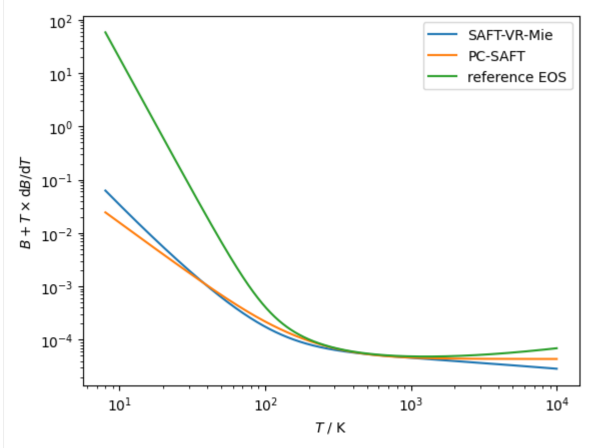
10²

44 Chapter 4. Models

 10^{3}

10⁴

```
"kind": 'SAFT-VR-Mie',
        "model": { "names": [name] }
})
modelPCSAFT = teqp.make_model({
        "kind": 'PCSAFT',
        "model": { "names": [name] }
})
modelMF = teqp.build_multifluid_model([name], teqp.get_datapath())
for model, label in [(modelVR, 'SAFT-VR-Mie'),
                     (modelPCSAFT, 'PC-SAFT'),
                     (modelMF, 'reference EOS')]:
    z = np.array([1.0])
    T = np.geomspace(8, 10000, 10000)
    n = 2
    B, TdBdT, thetan = [],[],[]
    for T_ in T:
        TdBdT.append(model.get_dmBnvirdTm(n, 1, T_, z)*T_)
        B.append(model.get_dmBnvirdTm(n, 0, T_, z))
        thetan.append(B[-1]+TdBdT[-1])
    plt.plot(T, thetan, label=label)
plt.xscale('log')
plt.yscale('log')
plt.gca().set(xlabel=r'$T$ / K', ylabel=r'$B+T\times$$d$B$/d$T$')
plt.legend()
plt.show()
```



4.10. SAFT-VR-Mie 45

4.10.1 Calculation of diameter

The calculation of the diameter is based upon

$$d_{ii} = \int_0^{\sigma_{ii}} (1 - \exp(-\beta u_{ii}^{ ext{Mie}}(r)) ext{d}r$$

but the integrand is basically constant from 0 to some cutoff value of r, which we'll call $r_{\rm cut}$. So first we need to find the value of $r_{\rm cut}$ that makes the integrand take its constant value, which is explained well in the paper from Aasen (https://github.com/ClapeyronThermo/Clapeyron.jl/issues/152#issuecomment-1480324192). Finding the cutoff value is obtained when

$$\exp(-\beta u_{ii}^{\text{Mie}}(r)) = EPS$$

where EPS is the numerical precision of the floating point type. Taking the logs of both sides,

$$-\beta u_{ii}^{\rm Mie} = \ln(EPS)$$

To get a starting value, it is first assumed that only the repulsive contribution contributes to the potential, yielding $u^{\text{rep}} = C\epsilon(\sigma/r)^{\lambda_r}$ which yields

$$-\beta C\epsilon (\sigma/r)^{\lambda_r} = \ln(EPS)$$

and

$$(\sigma/r)_{\rm guess} = (-\ln(EPS)/(\beta C\epsilon))^{1/\lambda_r}$$

Then we solve for the residual R(r)=0, where $R_0=\exp(-u/T)-EPS$. Equivalently we can write the residual in logarithmic terms as $R=-u/T-\ln(EPS)$. This simplifies the rootfinding as you need R, R' and R'' to apply Halley's method, which are themselves quite straightforward to obtain because R'=-u'/T, R''=-u''/T, where the primes are derivatives taken with respect to σ/r .

```
[9]: # Calculation of the residual function (needed for Halley's method) import sympy as sy kappa, j, lambda_r, lambda_a = sy.symbols('kappa, j, lambda_r, lambda_a') u = kappa*(j**lambda_r - j**lambda_a) display(sy.diff(u, j)) display(sy.simplify(sy.diff(u, j, 2))) \kappa \left(-\frac{j^{\lambda_a}\lambda_a}{j} + \frac{j^{\lambda_r}\lambda_r}{j}\right)
```

```
\frac{\kappa \left(-j^{\lambda_a}\lambda_a^2 + j^{\lambda_a}\lambda_a + j^{\lambda_r}\lambda_r^2 - j^{\lambda_r}\lambda_r\right)}{j^2}
```

```
[10]: # Here is a small example of using adaptive quadrature
     # to obtain the quasi-exact value of d for ethane
      # according to the pure-fluid parameters given in
      # Lafitte et al.
     epskB = 206.12 \# [K]
     sigma_m = 3.7257e-10 \# [m]
     lambda_r = 12.4
     lambda_a = 6.0
     C = lambda_r/(lambda_r-lambda_a) * (lambda_r/lambda_a) ** (lambda_a/(lambda_r-lambda_a))
     T = 300.0 \# [K]
      # The classical method based on adaptive quadrature
     def integrand(r m):
         u = C*epskB*((sigma_m/r_m)**(lambda_r) - (sigma_m/r_m)**(lambda_a))
         return 1.0 - np.exp(-u/T)
     print('quasi-exact; (value, error estimate):')
     exact, exact_error = scipy.integrate.quad(integrand, 0.0, sigma_m, epsrel=1e-16,__
      \rightarrowepsabs=1e-16)
     print(exact*1e10, exact_error*1e10)
     j = {"kind": 'SAFT-VR-Mie', "model": {"names": ["Ethane"]}}
     model = teqp.make_model(j)
     d = model.get_core_calcs(T, -1, z)["dmat"][0][0]
     print('tegp; (value, error from quasi-exact in %)')
     print (d, abs (d/(exact*1e10)-1)*100)
     quasi-exact; (value, error estimate):
     3.597838592720949 3.2280056122233332e-12
     teqp; (value, error from quasi-exact in %)
     3.597838640613809 1.331156429529301e-06
```

4.11 SAFT-VR-Mie with polar contributions

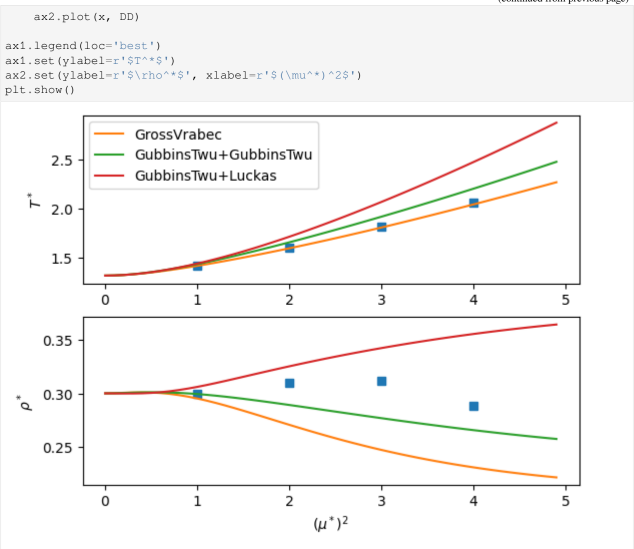
```
[1]: import teqp
teqp.__version__
[1]: '0.19.1'

[2]: import numpy as np
import matplotlib.pyplot as plt
import math

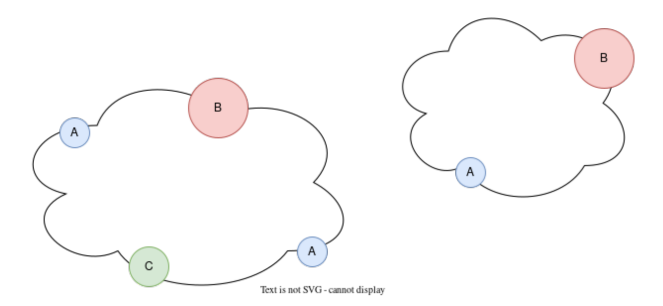
[3]: ek = 100 # [K]
sigma_m = 3e-10

N_A = 6.022e23
fig, (ax1, ax2) = plt.subplots(2, 1)
# # From https://arxiv.org/pdf/mtrl-th/9501001.pdf which pulled from M. van Leeuwen___
(continues on next page)
```

```
→and B. Smit, Phys. Rev. Lett. 71, 3991 (1993)
# These data need to be rescaled according to Hentschke et al. (DOI: https://doi.org/
→10.1103/physreve.75.011506)
\# mustar2 = [2.5, 3.0, 3.5, 4.0]
\# T = [2.63, 3.35, 4.20, 5.07]
# rho = [0.29, 0.25, 0.24, 0.24]
# ax1.plot(mustar2, T, 'd')
# ax2.plot(mustar2, rho, 'd')
# Comparing with Hentschke, DOI: https://doi.org/10.1103/physreve.75.011506
mustar2 = [1, 2, 3, 4]
T = [1.41, 1.60, 1.82, 2.06]
rho = [0.30, 0.31, 0.312, 0.289]
ax1.plot(mustar2, T, 's')
ax2.plot(mustar2, rho, 's')
kB = 1.380649e-23 \# Boltzmann's constant, J/K
epsilon_0 = 8.8541878128e-12 # Vacuum permittivity
for polar_model in ['GrossVrabec','GubbinsTwu+GubbinsTwu','GubbinsTwu+Luckas']:
    x = []; y = []; TT = []; DD = []
   rhostar_guess = 0.27
   Tstar\_guess = 1.5
    for mustar2 in np.arange(0.001, 5, 0.1):
        z = np.array([1.0])
        mu2_C2m2 = 4.0*np.pi*epsilon_0*sigma_m**3*ek*kB*mustar2
        mu_Cm = mu2_C2m2**0.5
        model = teqp.make_model({
            "kind": 'SAFT-VR-Mie',
            "model": {
                "polar_model": polar_model,
                "coeffs": [{
                    "name": "Stockmayer",
                    "BibTeXKey": "me",
                    "m": 1.0,
                    "epsilon_over_k": ek, # [K]
                    "sigma_m": sigma_m,
                    "lambda_r": 12.0,
                    "lambda_a": 6.0,
                    "mu_Cm": mu_Cm,
                    "nmu": 1.0
                } ]
        })
        T, rho = model.solve_pure_critical(Tstar_guess*ek, rhostar_guess/(N_A*sigma_
\rightarrowm**3))
        # Store the values
        x.append(mustar2)
        TT.append(T/ek)
        DD.append(rho*N_A*sigma_m**3)
        # Update the guess for the next calculation
        Tstar\_quess = TT[-1]
        rhostar\_guess = DD[-1]
    ax1.plot(x, TT, label=polar_model)
                                                                           (continues on next page)
```



4.12 Association



4.12.1 Site Interactions

Each unique site type per unique molecule is characterized by a numerical index siteid, which (for consistency with C++) starts with 0. In the example above, the indices would go like:

0: site type A on left molecule (multiplicity of 2)

1: site type B on left molecule (multiplicity of 1)

2: site type C on left molecule (multiplicity of 1)

3: site type A on right molecule (multiplicity of 1)

4: site type B on right molecule (multiplicity of 1)

Within a molecule, the numbering of sites is arbitrary, but the mapping cannot be changed once it is defined.

Association can occur when a site can "dock" with another kind of site. In the most common kind of association used to model hydrogen bonding, there are two classes of sites, positive or negative (e and H in Clapeyron.jl). teqp allows for great flexibility in defining the site types and how they are permitted to interact with each other.

The work of Langenbach and Enders (2012) shows how to construct a counting matrix to make the successive substitution faster because not all sites are included in the summation, rather the sites within a molecule are clustered into groups, since all sites of a similar type will have the same association fractions. Thus a counting matrix \mathbf{D} can be defined, with entries D_{IJ} for the pair of siteid \mathbb{I} and \mathbb{J} with the pseudocode

```
return counts[J]
return 0
```

in which the dictionary interaction_parameters defines which sites are allowed to interact with each other. The typical alcohol+water family would be modeled with:

```
interaction_parameters = {'e': ['H'], 'H': ['e']}
```

and to follow the system considered above, we would have:

```
inv_mapping = {
    0: (0, 'A'),
    1: (0, 'B'),
    2: (0, 'C'),
    3: (1, 'A'),
    4: (1, 'B')
}
counts = [2, 1, 1, 1, 1] # multiplicities for each siteid
```

The definition of the dictionary interaction_parameters would depend on how you want to allow the sites to associate. Sites that are not permitted to interact with each other are removed from the D matrix (are set to zero).

The successive substitution step gives the estimated values with

$$X_{\text{step}} = \frac{1}{1 + \rho_N \sum_J x_J X_J D_{IJ} \Delta_{IJ}}$$

in which ρ_N is the number density (molecules per volume) of the entire mixture, Δ_{IJ} is the interaction strength (volume per site) between site with siteid of I and that with siteid of J and x_J is the mole fraction of the molecule that site J is found in.

Acceleration can be achieved by taking only a partial step of successive substitution, weighted by α :

$$X_{\text{new}} = \alpha X_{\text{old}} + (1 - \alpha) X_{\text{step}}$$

This is the method utilized in Langenbach and Enders.

4.12.2 Interaction strength

The interaction site strength is a matrix with side length of the number of siteid. It is a block matrix because practically speaking the interaction sites are still about molecule-molecule interactions

$$\Delta_{IJ} = gb_{IJ}\beta_{IJ} \left(\exp\left(\frac{\epsilon_{IJ}}{RT}\right) - 1 \right) / N_A$$

Reminder: b, β , and ϵ values are associated with the *molecule*, not the site.

CR1 combining rule

In the CR1 combining rule:

$$b_{IJ} = b_{ij} = \frac{b_i + b_j}{2}$$
$$\beta_{IJ} = \beta_{ij} = \sqrt{\beta_i \beta_j}$$
$$\epsilon_{IJ} = \epsilon_{ij} = \frac{\epsilon_i + \epsilon_j}{2}$$

in which i is the molecule index associated with siteid I and the same for j and J

4.12. Association 51

4.12.3 Radial distribution function

CS:
$$g = \frac{2-\eta}{2(1-\eta)^2}$$

KG: $g = \frac{1}{1-1.9\eta}$

where $\eta = b_{\rm mix} \rho/4$ in which ρ is density with units to match the reciprocal of $b_{\rm mix}$ (so if $b_{\rm mix}$ is mean covolume per atom, then ρ is the number density $\rho_{\rm N}$)

References:

K. Langenbach & S. Enders (2012): Cross-association of multi-component systems, Molecular Physics, 110:11-12, 1249-1260; https://dx.doi.org/10.1080/00268976.2012.668963

```
[1]: import teqp, numpy as np
```

```
[2]: ethanol = {
        "a0i / Pa m^6/mol^2": 0.85164,
        "bi / m^3/mol": 0.0491e-3,
         "c1": 0.7502,
        "Tc / K": 513.92,
        "epsABi / J/mol": 21500.0,
        "betaABi": 0.008,
        "sites": ["e", "H"]
    water = {
        "a0i / Pa m^6/mol^2": 0.12277,
        "bi / m^3/mol": 0.0000145,
        "c1": 0.6736,
        "Tc / K": 647.13,
        "epsABi / J/mol": 16655.0,
        "betaABi": 0.0692,
         "sites": ["e", "e", "H", "H"]
    }
    jCPA = {
        "cubic": "SRK",
        "radial_dist": "CS",
          "combining": "CR1", # No other option is implemented yet
        "pures": [ethanol, water],
         "R_gas / J/mol/K": 8.31446261815324
    }
    model = teqp.make_model({"kind": "CPA", "model": jCPA, "validate": False}, False)
    T = 303.15 \# K
    rhomolar = 1/3.0680691201961814e-5 \# mol/m^3
    molefracs = np.array([0.3, 0.7])
    # Note: passing data back and forth in JSON format is done for convenience and.
     →flexibility, not speed
    res = model.get_assoc_calcs(T, rhomolar, molefracs)
    print('D:', np.array(res['D']))
    print('∆:', np.array(res['Delta']))
    print('X_A:', np.array(res['X_A']))
    print('siteid->(component, name):', res['to_CompSite'])
    print('(component, name)->siteid:', res['to_siteid'])
    print('multiplicities:', np.array(res['counts']))
```

```
D: [[0 1 0 2]
  [1 0 2 0]
  [0 1 0 2]
  [1 0 2 0]]

Δ: [[5.85623687e-27 5.85623687e-27 4.26510827e-27 4.26510827e-27]
  [5.85623687e-27 5.85623687e-27 4.26510827e-27 4.26510827e-27]
  [4.26510827e-27 4.26510827e-27 2.18581242e-27 2.18581242e-27]
  [4.26510827e-27 4.26510827e-27 2.18581242e-27 2.18581242e-27]
  [X_A: [0.062584    0.062584    0.10938445]
  siteid->(component, name): [[0, [0, 'H']], [1, [0, 'e']], [2, [1, 'H']], [3, [1, 'e']]]
  (component, name)->siteid: [[[0, 'H'], 0], [[0, 'e'], 1], [[1, 'H'], 2], [[1, 'e'], ... ... ... ...]]
  multiplicities: [1 1 2 2]
```

```
[3]: # For completeness, here is the worked Python example that was used to develop the.
     →association implementation in teqp:
    import collections
    import numpy as np
    class AssocClass:
        def __init__(self, molecules):
             # Get all the kinds of sites present
            mapping = {}
            counts = {}
            def sort_sites(sites):
                 counts = collections.Counter(sites)
                out = []
                for k in ['B', 'P', 'N']:
                     if k in counts:
                         out += [k]*counts[k]
                 return out
            uid = 0
            for i, molecule in enumerate(molecules):
                 for site in sort_sites(set(molecule)):
                     mapping[(i, site)] = uid
                     counts[uid] = molecule.count(site)
                     uid += 1
            inv_mapping = {v:k for k,v in mapping.items()} # from superindex to (molecule,
     → site pair)
            interaction_partners = {
                     'B': ('N', 'P', 'B'),
                     'N': ('P', 'B'),
                     'P': ('N', 'B')
            def get_DIJ(I, J):
                 """ Return the value of an entry in the D_{IJ} matrix
                 For a given unique site, look at all other sites on all other molecules
                 _, typei = inv_mapping[I]
                                                                               (continues on next page)
```

4.12. Association 53

```
_, typej = inv_mapping[J]
           if typej in interaction_partners[typei]:
               return counts[J]
           return 0
       Ngroups = len(mapping)
       D = np.zeros((Ngroups, Ngroups), dtype=int)
       for I in range(Ngroups):
           for J in range(Ngroups):
               D[I, J] = get_DIJ(I, J)
        # Store variables needed for later use
       self.D = D
       self.counts = counts
       self.inv_mapping = inv_mapping
       self.Ngroups = Ngroups
       # ethanol, water
       self.b_Lmol = np.array([0.0491, 0.0145])
       self.epsilon_barLmol = np.array([215.00, 166.55])
       self.beta = [8e-3, 69.2e-3]
       self.b_m3mol = self.b_Lmol/1e3
       R = 8.31446261815324 \# J/(mo1*K)
       self.epsilon_K = self.epsilon_barLmol*100/R # K, from (bar*L)/mol * (1e5 Pa/
\rightarrowbar) * (Pa / 1000 L), Pa*m^3 = J, then we divide by R to do [J/mol]/[J/mol/K] -> K
   def get_xJ(self, moleculemolefracs):
       Return the fractions of sites within the mixture, not to be confused
       with the mole fractions of molecules within the mixture
       counter = 0
       xJ = np.zeros((self.D.shape[0],))
       for J in range(self.D.shape[0]):
           j, sitej = self.inv_mapping[J] # molecule index and site name
           xJ[J] = self.counts[J]*moleculemolefracs[j]
           counter += xJ[J]
       return xJ/counter
   def get_bmix(self, molefracs):
       return (self.b_m3mol*molefracs).sum()
   def get_bij(self, i, j):
       """ CR1 """
       return (self.b_m3mol[i] + self.b_m3mol[j])/2
   def get_epsilon_k_IJ_CR1(self, *, i, j):
        """ CR1 """
       return (self.epsilon_K[i] + self.epsilon_K[j])/2
   def get_beta_IJ_CR1(self, *, i, j):
       """ CR1 """
       return (self.beta[i]*self.beta[j])**0.5
   def get_DeltaIJ(self, T, rhomolar, molefracs, *, i, j):
       b_ij = self.get_bij(i, j)
                                                                          (continues on next page)
```

```
bmix = self.get_bmix(molefracs)
       eta = bmix*rhomolar/4 # packing fraction
       g_{ij} = (2-eta)/(2*(1-eta)**3)
       beta = self.get_beta_IJ_CR1(i=i,j=j) # dimensionless
       return g_ij*b_ij*beta*(np.exp(eRT)-1.0) # epsilon_k_IJ is in K, beta_IJ is__
→dimensionless
   def get_Delta(self, T, rhomolar, *, molefracs, Ngroups):
       Delta = np.zeros((Ngroups, Ngroups))
       for I in range(Ngroups):
           i, _ = self.inv_mapping[I]
           for J in range(Ngroups):
               j, _ = self.inv_mapping[J]
               Delta[I, J] = self.get_DeltaIJ(T, rhomolar, i=i, j=j,__
→molefracs=molefracs)
       return Delta
   def X_iter_Langenbach(self, T:float, rhomolar:float, molefracs, init):
       """Iterate with successive substitution to obtain the non-bonded fraction of
⊶each site
       Aras:
           T (float): Temperature, K
           rhomolar (float): Molar density, mol/m^3
           molefracs (array): Mole fractions of the components
           init (array): Starting values for X_A
       Returns:
           array: non-bonded fractions for each site as one big array, indexed by-
⇒site family
       TODO: why do we need mole fractions here and site fractions elsewhere?
       # xJ = np.array(self.get_xJ(moleculemolefracs=molefracs), ndmin=2) # row_
→vector
       XXJ = np.array([ molefracs[self.inv_mapping[J][0]] for J in range(self.
→Ngroups)])
       N_A = 6.02214076e23 \# [1/mol]
       Delta = self.get_Delta(T, rhomolar, Ngroups=self.Ngroups,__
→molefracs=molefracs)/N_A
       rhoN = rhomolar*N_A # number density, in 1/m^3
       Y = np.array(init[:], ndmin=2) # copy, row vector
       DD = self.D*Delta # coefficient-wise product
       DDX = XXJ*DD # coefficient-wise product
       for _ in range(30):
           # The naive treatment
           summer = 0.0
           for J in range(self.Ngroups):
               summer += Y[0,J]*XXJ[J]*self.D[:,J]*Delta[:,J]
           # Optimized treatment
           summer2 = (DDX@Y.T).squeeze()
           # print(summer, summer2)
           term = rhoN*summer2
                                                                      (continues on next page)
```

4.12. Association 55

```
Y = 0.5*(Y+1/(1+term))
       return Y
   def X_A_pure_Langenbach(self, i:int, T:float, rhomolar:float):
       """Calculate the association fractions for a pure fluid
       based upon the method of Eq. 20, from
       Langenbach & Enders, Mol. Phys.
       URL: https://www.tandfonline.com/doi/abs/10.1080/00268976.2012.668963
       Args:
           int (int): Index of the pure fluid
           T (float): Temperature, K
           rhomolar (float): Molar density, mol/m^3
           molefracs (_type_): Molar fractions, array
       TODO: why do we need site fractions here and mole fractions elsewhere?
       molefracs = [0]*len(self.b_m3mol)
       molefracs[i] = 1
       xJ = self.get_xJ(moleculemolefracs=molefracs)
       N_A = 6.02214076e23 \# [1/mol]
       Delta = self.get_Delta(T, rhomolar, Ngroups=self.Ngroups,_
→molefracs=molefracs)/N_A
       common = np.array(2*rhomolar*N_A*(xJ@self.D@Delta), ndmin=2).sum(axis=0)
       return (np.sqrt(1+2*common)-1)/common
   def X_A_pure_HuangRadosz(self, *, i:int, T:float, rhomolar:float, klass:str):
       """Use the explicit solutions from Huang and Radosz to obtain the
       association fraction for a pure fluid
       Args:
           i (int): The fluid index for which the method is being applied
           T (float): Temperature, K
           rhomolar (float): Molar density, in mol/m^3
           klass (str): Association class, one in {'2B','3B','4C'}
       Returns:
           float: value of X_A
       b_ij = b_cubic = self.get_bij(i=i,j=i)
       betaABi = self.get_beta_IJ_CR1(i=i,j=i)
       R = 8.31446261815324
       RT = R*T
       epsABi = self.get_epsilon_k_IJ_CR1(i=i,j=i)*R # To get J/mol
       eta = b_ij*rhomolar/4 # packing fraction
       g_vm_ref = (2-eta)/(2*(1-eta)**3)
       DeltaAiBj = g_vm_ref*(np.exp(epsABi/RT) - 1.0)*b_cubic* betaABi
       if klass == '2B':
           X_A = (-1.0 + (1.0 + 4.0 * rhomolar * DeltaAiBj) ** 0.5) / (2.0 * rhomolar_
→* DeltaAiBj)
       elif klass == '3B':
           X_A = ((-(1.0 - rhomolar * DeltaAiBj) + np.sqrt((1.0 + rhomolar *_
→DeltaAiBj) **2 + 4.0 * rhomolar * DeltaAiBj)) / (4.0 * rhomolar * DeltaAiBj))
                                                                         (continues on next page)
```

```
elif klass == '4C':
            X_A = (-1.0 + np.sqrt(1.0 + 8.0 * rhomolar * DeltaAiBj)) / (4.0 *_
→rhomolar * DeltaAiBj)
        return X_A
if __name__ == '__main__':
    a = AssocClass([('B'), ('P', 'N', 'N'), ('P')])
    assert(a.D.tolist() == [[1, 1, 2, 1], [1, 0, 2, 0], [1, 1, 0, 1], [1, 0, 2, 0]])
    #### 4C water
   a = AssocClass([(), ('P', 'P', 'N', 'N')])
   T = 303.15
   rhomolar = 1/1.7915123921401366e-5
   X A Clapeyron = 0.07920738195861185 # version 0.5.9
   X_A_HR = a.X_A_pure_HuangRadosz(i=1, T=T, rhomolar=rhomolar, klass='4C')
   X_A_La = a.X_A_pure_Langenbach(i=1, T=T, rhomolar=rhomolar)[0]
    assert (abs (X_A_HR - X_A_Clapeyron) < 1e-10)</pre>
    assert (abs (X_A_La - X_A_Clapeyron) < 1e-10)</pre>
    a.X_iter_Langenbach(T=T, rhomolar=rhomolar, molefracs=[0,1], init=np.array([1.0,-
\hookrightarrow 1.01)
    ### 2B ethanol
    a = AssocClass([('P','N'), ()])
   T = 303.15
   rhomolar = 1/1.7915123921401366e-5
   X_A_Clapeyron = 0.020464699705843845 # version 0.5.9
   X_A_HR = a.X_A_pure_HuangRadosz(i=0, T=T, rhomolar=rhomolar, klass='2B')
   X_A_La = a.X_A_pure_Langenbach(i=0, T=T, rhomolar=rhomolar)[0]
    assert (abs (X_A_HR - X_A_Clapeyron) < 1e-10)</pre>
    assert (abs(X_A_La - X_A_Clapeyron) < 1e-10)</pre>
    a.X_iter_Langenbach(T=T, rhomolar=rhomolar, molefracs=[1,0], init=np.array([1.0,__
\hookrightarrow 1.0])
    a = AssocClass([('P','N'), ('P','P','N','N')])
   T = 303.15
   print(a.D)
   rhomolar = 1/3.0680691201961814e-5
   print(a.get_Delta(T, rhomolar, molefracs=[0.3, 0.7], Ngroups=4)/6.02214076e23)
   print(a.X_iter_Langenbach(T=T, rhomolar=rhomolar, molefracs=[0.3,0.7], init=np.
\rightarrowarray([1.0, 1.0, 1, 1])))
[[0 1 0 2]
[1 0 2 0]
[0 1 0 2]
[1 0 2 0]]
[[5.85623687e-27 5.85623687e-27 4.26510827e-27 4.26510827e-27]
[5.85623687e-27 5.85623687e-27 4.26510827e-27 4.26510827e-27]
[4.26510827e-27 4.26510827e-27 2.18581242e-27 2.18581242e-27]
[4.26510827e-27 4.26510827e-27 2.18581242e-27 2.18581242e-27]]
[[0.062584    0.062584    0.10938445    0.10938445]]
```

4.12. Association 57

4.13 Multi-fluid EOS

Peering into the innards of teqp

```
[1]: import timeit, json
  import pandas
  import numpy as np
  import teqp
  teqp.__version__
[1]: '0.19.1'
```

4.13.1 Ancillary Equations

Ancillary equations are provided along with multiparameter equations of state. The give a good *approximation* to the phase equilibrium densities. There are routines in teqp to use the ancillary equations provided with the EOS. First a class containing the ancillary equations is obtained, then methods on that class are called

```
[2]: model = teqp.build_multifluid_model(["Methane"], teqp.get_datapath())
anc = model.build_ancillaries()
T = 100.0 # [K]
rhoL, rhoV = anc.rhoL(T), anc.rhoV(T)
print('Densities are:', rhoL, rhoV, 'mol/m^3')
Densities are: 27357.335621492966 42.04100696197727 mol/m^3
```

But those densities do not correspond to the *true* phase equilibrium solution, so we need to polish the solution:

```
[3]: Niter = 10
    rhoLtrue, rhoVtrue = model.pure_VLE_T(T, rhoL, rhoV, Niter)
    print('VLE densities are:', rhoLtrue, rhoVtrue, 'mol/m^3')

VLE densities are: 27357.147019094467 42.047982278351704 mol/m^3
```

And looking the densities, they are slightly different after the phase equilibrium calculation

4.13.2 Ammonia-Water

Tillner-Roth and Friend provided a hard-coded model that is in a form not compatible with the other multi-fluid models. It is available via the high-level factory function

```
[4]: AW = teqp.AmmoniaWaterTillnerRoth()
   AW.get_Ar01(300, 300, np.array([0.9, 0.0]))
[4]: -0.09731055757504622
```

4.13.3 Pure fluid loading

So, how to make it faster? Only do it once and cache

```
[7]: # Here is the set of possible aliases to absolute paths of files
    # Building this map takes a little while (somewhat faster in C++) due to all the file.
     → reads
    # If you know your files will not change, good idea to build this alias map yourself.
    %timeit aliasmap = teqp.build_alias_map(teqp.get_datapath())
    aliasmap = teqp.build_alias_map(teqp.get_datapath())
    list(aliasmap.keys())[0:10] # the first 10 aliases in the dict
    31.6 ms \pm 182 \mus per loop (mean \pm std. dev. of 7 runs, 10 loops each)
[7]: ['1,2-DICHLOROETHANE',
     '1,2-dichloroethane',
     '1-BUTENE',
     '1-Butene',
     '100-41-4',
     '10024-97-2',
     '102687-65-0',
     '106-42-3',
     '106-97-8',
     '106-98-9'1
```

```
[8]: # Then load the absolute paths from the alias map,
    # which will guarantee that you hit exactly what you were looking for,
    # resolving aliases as needed
    identifiers = [aliasmap[n] for n in ["n-C1H4"]]
    %timeit model = teqp.build_multifluid_model(identifiers, teqp.get_datapath())

533 µs ± 3.06 µs per loop (mean ± std. dev. of 7 runs, 1,000 loops each)
```

At some point soon teqp will support in-memory loading of JSON data for the pure components, without requiring reads from the operating system

```
[9]: # And you can also load the JSON that teqp is loading for the pure fluids
pureJSON = teqp.collect_component_json(['Neon','Hydrogen'], teqp.get_datapath())
```

4.13. Multi-fluid EOS 59

4.13.4 Mixture model loading

```
[10]: # Load the default JSON for the binary interaction parameters
     BIP = json.load(open(teqp.qet_datapath()+'/dev/mixtures/mixture_binary_pairs.json'))
[11]: # You can obtain interaction parameters either by pairs of names, where name is the
      →name that teqp uses, the ["INFO"]["NAME"] field
     params, swap_needed = teqp.get_BIPdep(BIP, ['Methane','Ethane'])
     params
[11]: {'BibTeX': 'Kunz-JCED-2012',
      'CAS1': '74-82-8',
      'CAS2': '74-84-0',
      'F': 1.0,
      'Name1': 'Methane',
      'Name2': 'Ethane',
      'betaT': 0.996336508,
      'betaV': 0.997547866,
      'function': 'Methane-Ethane',
      'gammaT': 1.049707697,
      'gammaV': 1.006617867}
[12]: # Or also by CAS#
     params, swap_needed = teqp.get_BIPdep(BIP, ['74-82-8','74-84-0'])
     params
[12]: {'BibTeX': 'Kunz-JCED-2012',
      'CAS1': '74-82-8',
      'CAS2': '74-84-0',
      'F': 1.0,
      'Name1': 'Methane',
      'Name2': 'Ethane',
      'betaT': 0.996336508,
      'betaV': 0.997547866,
      'function': 'Methane-Ethane',
      'gammaT': 1.049707697,
      'gammaV': 1.006617867}
[13]: # But mixing is not allowed
     params, swap_needed = teqp.get_BIPdep(BIP, ['74-82-8','Ethane'])
     params
     ValueError
                                                Traceback (most recent call last)
     Cell In[13], line 2
          1 # But mixing is not allowed
      ----> 2 params, swap_needed = teqp.get_BIPdep(BIP, ['74-82-8','Ethane'])
           3 params
     ValueError: Can't match the binary pair for: 74-82-8/Ethane
```

4.13.5 Estimation of interaction parameters

Estimation of interaction parameters can be used when no mixture model is present. The flags keyword argument allows the user to control how estimation is applied. The flags keyword argument should be a dictionary, with keys of "estimate" to provide the desired estimation scheme as-needed. For now, the only allowed estimation scheme is Lorentz-Berthelot.

If it is desired to force the estimation, the "force-estimate" to force the use of the provided estimation scheme for all binaries, even when a proper mixture model is available. The value associated with "force-estimate" is ignored.

```
[14]: params, swap_needed = teqp.get_BIPdep(BIP, ['74-82-8','74-84-0'], flags={'force-
      →estimate':'yes', 'estimate': 'Lorentz-Berthelot'})
[14]: {'F': 0.0, 'betaT': 1.0, 'betaV': 1.0, 'gammaT': 1.0, 'gammaV': 1.0}
[15]: # And without the force, the forcing is ignored
     params, swap_needed = teqp.get_BIPdep(BIP, ['74-82-8','74-84-0'], flags={'estimate':
     →'Lorentz-Berthelot'})
     params
[15]: {'BibTeX': 'Kunz-JCED-2012',
      'CAS1': '74-82-8',
      'CAS2': '74-84-0',
      'F': 1.0,
      'Name1': 'Methane',
      'Name2': 'Ethane',
      'betaT': 0.996336508,
      'betaV': 0.997547866,
      'function': 'Methane-Ethane',
       'gammaT': 1.049707697,
       'gammaV': 1.006617867}
[16]: # And the same flags can be passed to the multifluid model constructor
     model = tegp.build_multifluid_model(
         ['74-82-8','74-84-0'],
         teqp.get_datapath(),
         flags={'force-estimate':'yes', 'estimate': 'Lorentz-Berthelot'})
```

4.14 Multfluid mutant

These adapted multifluid models are used for fitting departure functions. The pure fluids remain fixed while you can adjust the mixture model, both the interaction parameters as well as the departure function terms

4.14. Multfluid mutant 61

```
[3]: %timeit teqp.build_multifluid_mutant(basemodel, s)

26.4 µs ± 2.53 µs per loop (mean ± std. dev. of 7 runs, 10,000 loops each)
```

```
[4]: mutant.get_Ar01(300, 3.0, np.array([0.5, 0.5]))
[4]: -0.00017517184039893556
```

4.15 REFPROP < 10.0 conversion

As of teqp version 0.19.0, it is possible to read in the .FLD and HMX.BNC of NIST REFPROP 10.0 and load them into teqp multifluid models. There are two approaches; either you can pass paths to the files of interest, or you can load them into JSON once, and pass the converted JSON back into teqp's make_model function.

The conversion code is uses that of REFPROP-interop and the fluid file format of CoolProp is used.

The example is based on the interaction parameters provided in the supporting information of the paper Mixture Model for Refrigerant Pairs R-32/1234yf, R-32/1234ze(E), R-1234ze(E)/227ea, R-1234yf/152a, and R-125/1234yf by Ian Bell

```
[1]: import json
import teqp
teqp.__version__
[1]: '0.19.1'

[2]: # The first approach, we just pass paths to the files, they live in the folder
# containing this notebook, and teqp does the conversion on the fly
jsimple = {
    'kind': 'multifluid',
    'model': {
        'HMX.BNC': 'HMX.BNC',
        'components': ['R152A.FLD', 'NEWR1234YF.FLD'],
    }
}
model = teqp.make_model(jsimple)
```

```
[3]: %timeit teqp.make_model(jsimple)

45.6 ms ± 1.46 ms per loop (mean ± std. dev. of 7 runs, 10 loops each)
```

```
[4]: # Convert each of the FLD files to JSON
FLD0 = teqp.convert_FLD('R152A.FLD', name='R152A')
FLD1 = teqp.convert_FLD('NEWR1234YF.FLD', name='R1234YF')
BIP, DEP = teqp.convert_HMXBNC('HMX.BNC')
```

```
[5]: jconverted = {
    "kind": "multifluid",
    "model": {
        "components": [FLD0, FLD1],
        "BIP": BIP,
        "departure": DEP
    }
}
model = teqp.make_model(jconverted)
```

```
[6]: %timeit teqp.make_model(jconverted)

602 µs ± 2.32 µs per loop (mean ± std. dev. of 7 runs, 1,000 loops each)
```

From this example you can note that the first method is a lot slower because the FLD->JSON conversion needs to happen for each call, while in the second method it is much faster because only the JSON parsing needs to be done in tegp.

```
[7]: # It is also possible to prefix the path to indicate that the
    # indicated file (after the FLD::) should be converted from REFPROP format
    jconverted = {
        "kind": "multifluid",
        "model": {
            "components": ["FLDPATH::R152A.FLD", 'FLDPATH::NEWR1234YF.FLD'],
            "BIP": BIP,
            "departure": DEP
        }
    }
    model = teqp.make_model(jconverted)
```

```
[8]: %timeit teqp.make_model(jconverted)

40.8 ms ± 180 µs per loop (mean ± std. dev. of 7 runs, 10 loops each)
```

4.16 **GERG**

In the GERG-2004 and GERG-2008 models, the pure fluids are modeled with high-accuracy multiparameter EOS. The model is covered exhaustively in the GERG-2004 monograph: https://www.gerg.eu/wp-content/uploads/2019/10/TM15.pdf and in the GERG-2008 paper: https://doi.org/10.1021/je300655b

The following components are supported (case-sensitive) in GERG-2004:

- methane
- nitrogen
- · carbondioxide
- · ethane
- propane
- n-butane

4.16. GERG 63

- isobutane
- · n-pentane
- · isopentane
- · n-hexane
- n-heptane
- · n-octane
- hydrogen
- oxygen
- · carbonmonoxide
- · water
- · helium
- argon

and GERG-2008 adds the components:

- · hydrogensulfide
- n-nonane
- n-decane

(as well as modifying the pure component EOS for carbon monoxide and isopentane).

The interaction parameters and departure functions are not editable (by design) and the EOS parameters are hard-coded. No ancillary equations are available along with the GERG-2004 model, but you can use the on-the-fly ancillary generator of teqp.

The residual portions of these models were added in version 0.18.0, and it is planned to add the ideal-gas portions as well at a later date. The residual portion is enough for many applications like phase equilibria and critical locus tracing.

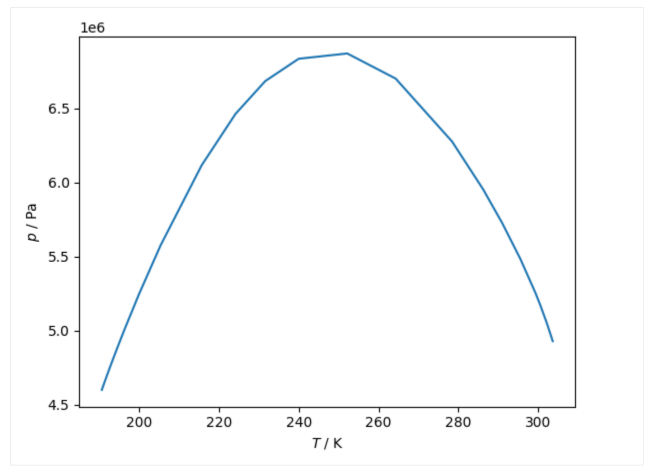
The kind is 'GERG2004resid' for the GERG-2004 residual model and 'GERG2008resid' for the GERG-2008 residual model

```
[1]: import teqp
    import numpy as np
    import pandas
    import matplotlib.pyplot as plt
    teqp.__version__
[1]: '0.19.1'
[2]: model = teqp.make_model({'kind':"GERG2004resid", 'model':{"names": ['methane', 'ethane
     ' ] } } )
[3]: # Note that names are case-sensitive; this doesn't work
    model = teqp.make_model({'kind':"GERG2004resid", 'model':{"names": ['MeThAnE','ethane
     ' ] } } )
    ValueError
                                                 Traceback (most recent call last)
    Cell In[3], line 2
          1 # Note that names are case-sensitive; this doesn't work
     ---> 2 model =<u>-</u>
                                                                                  (continues on next page)
```

```
→teqp.make_model({'kind':"GERG2004resid", 'model':{"names": ['MeThAnE', 'ethane']}})
File /opt/conda/lib/python3.11/site-packages/teqp/__init__.py:47, in make_model(*args,
→ **kwargs)
    42 def make_model(*args, **kwargs):
    43
     44
           This function is in two parts; first the make_model function (renamed to _
→make_model in the Python interface)
           is used to make the model and then the model-specific methods are.
\rightarrowattached to the instance
    46
---> 47
           AS = _make_model(*args, **kwargs)
    48
          attach_model_specific_methods(AS)
    49
           return AS
ValueError: Unable to load pure info for MeThAnE
```

```
[4]: # Here we trace the critical locus for methane+ethane
    rhovec0 = np.array([0.0, 0.0])
    ifluid = 0
    T0 = model.get_Tcvec()[0]
    rhovec0[ifluid] = 1/model.get_vcvec()[0]
    trace = model.trace_critical_arclength_binary(T0=T0, rhovec0=rhovec0)
    df = pandas.DataFrame(trace)
    plt.plot(df['T / K'], df['p / Pa'])
    plt.gca().set(xlabel='$T$ / K', ylabel='$p$ / Pa');
```

4.16. GERG 65



4.17 Extended Corresponding States

This implements the method of Huber and Ely: https://doi.org/10.1016/0140-7007(94)90083-3

It does not include the undocumented temperature and density terms that are included in REFPROP

```
[1]: import teqp
teqp.__version__
```

```
[1]: '0.19.1'
[2]: import numpy as np
    import CoolProp.CoolProp as CP
[3]: # These parameters are from Huber & Ely
    j = {
        "kind": "multifluid-ECS-HuberEly1994",
        "model": {
          "reference_fluid": {
                 "name": teqp.get_datapath() + "/dev/fluids/R134a.json",
                 "acentric": 0.326680,
                 "Z_crit": 4.056e6/(5030.8*8.314471*374.179),
                 "T_crit / K": 374.179,
                 "rhomolar_crit / mol/m^3": 5030.8
           },
           "fluid": {
                 "name": "R143a",
                 "f_T_coeffs": [ -0.22807e-1, -0.64746],
                 "h_T_coeffs": [ 0.36563, -0.26004e-1],
                 "acentric": 0.25540,
                 "T_crit / K": 346.3,
                 "rhomolar_crit / mol/m^3": (1/0.194*1000),
                 "Z_crit": 3.76e6/(346.3*8.314471*(1/0.194*1000))
          }
        }
    }
    model = teqp.make_model(j)
    z = np.array([1.0])
    R = model.get_R(z)
    T, rho = 400, 2600
    p = rho*R*T*(1+model.get_Ar01(T, rho, z))
    display('pressure from ECS:', p)
    display('pressure from EOS:', CP.PropsSI('P','T',T,'Dmolar',rho,'R143a'))
    'pressure from ECS:'
    5556329.442047298
    'pressure from EOS:'
    5478978.746656995
```

4.18 Ideal-gas Models

The collection of ideal-gas contributions are described below. They are summed to yield the ideal-gas contribution from

$$\alpha^{ig} = \sum_{i} x_i \left(\alpha_i^{ig}(T, \rho) + \ln(x_i) \right)$$

Null mole fractions $x_i = 0$ do not contribute to the summation because

$$\lim_{x_i \to 0} x_i \ln(x_i) = 0$$

4.18.1 IdealHelmholtzConstant

JSON arguments: "a"

$$\alpha^{ig} = a$$

4.18.2 IdealHelmholtzLogT

JSON arguments: "a"

$$\alpha^{ig} = a \ln(T)$$

which should be compared with the original form in GERG (and REFPROP and CoolProp)

$$\alpha^{\rm ig} = a^* \ln(\tau)$$

with $\tau = T_r/T$.

4.18.3 IdealHelmholtzLead

JSON arguments: "a_1", "a_2"

$$\alpha^{ig} = \ln(\rho) + a_1 + a_2/T$$

which should be compared with the original form in GERG (and REFPROP and CoolProp)

$$\alpha^{\mathrm{ig}} = \ln(\delta) + a_1^* + a_2^* \tau$$

Note that a_1 contains an additive factor of $-\ln(\rho_r)$ and a_2 contains a multiplicative factor of T_r relative to the former because $\delta = \rho/\rho_r$ and $\tau = T_r/T$.

4.18.4 IdealHelmholtzPowerT

JSON arguments: "n", "t"

$$\alpha^{\rm ig} = \sum_{k} n_k T^{t_k}$$

4.18.5 IdealHelmholtzPlanckEinstein

JSON arguments: "n", "theta"

$$\alpha^{\rm ig} = \sum_k n_k \ln(1 - \exp(-\theta_k/T))$$

4.18.6 IdealHelmholtzPlanckEinsteinGeneralized

JSON arguments: "n", "c", "d", "theta"

$$\alpha^{ig} = \sum_{k} n_k \ln(c_k + d_k \exp(\theta_k/T))$$

4.18.7 IdealHelmholtzGERG2004Cosh

JSON arguments: "n", "theta"

$$\alpha^{\mathrm{ig}} = \sum_{k} n_k \ln(|\cosh(\theta_k/T)|)$$

See Table 7.6 in GERG-2004 monograph

4.18.8 IdealHelmholtzGERG2004Sinh

JSON arguments: "n", "theta"

$$\alpha^{\mathrm{ig}} = \sum_{k} n_k \ln(|\sinh(\theta_k/T)|)$$

4.18.9 IdealHelmholtzCp0Constant

JSON arguments: "c", "T_0"

$$\alpha^{\mathrm{ig}} = c \left(\frac{T - T_0}{T} - \ln \left(\frac{T}{T_0} \right) \right)$$

from a term that is like

$$\frac{c_{p0}}{R} = c$$

4.18.10 IdealHelmholtzCp0PowerT

JSON arguments: "c", "t", "T_0"

$$\alpha^{\text{ig}} = c \left[T^t \left(\frac{1}{t+1} - \frac{1}{t} \right) - \frac{T_0^{t+1}}{T(t+1)} + \frac{T_0^t}{t} \right]$$

from a term that is like

$$\frac{c_{p0}}{R} = cT^t, t \neq 0$$

The C++ classes implementing these functions are at:

- IdealHelmholtzConstant
- IdealHelmholtzLogT
- IdealHelmholtzLead
- IdealHelmholtzPowerT

- IdealHelmholtzPlanckEinstein
- IdealHelmholtzPlanckEinsteinGeneralized
- IdealHelmholtzGERG2004Cosh
- IdealHelmholtzGERG2004Sinh
- IdealHelmholtzCp0Constant
- IdealHelmholtzCp0PowerT

Conversion

Conversion of terms from CoolProp format to teqp format is carried out in the function Cool-Prop2teqp_alphaig_term_reformatter().

For instance the leading term in CoolProp goes like:

$$\alpha = \ln(\delta) + a_1^* + a_2^* \tau$$

with the * indicating the CoolProp formulation. The term reads like

$$\alpha = \ln(\rho) + a_1 + a_2/T$$

in teqp. Refactoring the CoolProp term reads

$$\alpha = \ln(\rho) - \ln(\rho_r) + a_1^* + a_2^* \left(\frac{T_r}{T}\right)$$

so that
$$a_1 = a_1^* - \ln(\rho_r)$$
 and $a_2 = a_2^* T_r$

In some cases reconstitutions of terms are required, as the supported terms in the libraries are somewhat different. The term used in CoolProp to do the offsets to enthalpy and entropy is of the form

$$\alpha = a_1^* + a_2^* \tau = a_1^* + a_2^* \left(\frac{T_r}{T}\right)$$

so that term can be rewritten as an IdealHelmholtzPowerT with coefficients of a_1^* and $a_2^*T_r$ and exponents of 0 and -1.

Most of the remaining terms can be converted in a straightforward fashion, except for some of GERG formulations that are a bit trickier. Mostly, the only conversion required is to multiply or divide by reducing temperatures so that all arguments are in terms of temperature as independent variable.

The mathematics describing how to do the conversion from a term in c_p^0/R follows:

$$\alpha_0 = \frac{a_0}{RT} = -1 + \ln \frac{\rho T}{\rho_0 T_0} + \frac{h_0^0}{RT} - \frac{s_0^0}{R} + \frac{1}{RT} \int_{T_0}^T c_p^0(T) dT - \frac{1}{R} \int_{T_0}^T \frac{c_p^0(T)}{T} dT$$

$$\alpha_0 = \frac{a_0}{RT} = \ln(\rho) + \ln(T) - \ln(\rho_0 T_0) - 1 + \frac{h_0^0}{RT} - \frac{s_0^0}{R} + \frac{1}{RT} \int_{T_0}^T c_p^0(T) dT - \frac{1}{R} \int_{T_0}^T \frac{c_p^0(T)}{T} dT$$

You can set the values of h_0^0 and h_0^0 to any value, including zero. So if you are converting a term from c_p^0/R , then you could do

$$\alpha_0 = \frac{a_0}{RT} = \ln(\rho) + \ln(T) - \ln(\rho_0 T_0) - 1 + \frac{1}{RT} \int_{T_0}^T c_p^0(T) dT - \frac{1}{R} \int_{T_0}^T \frac{c_p^0(T)}{T} dT$$

70 Chapter 4. Models

```
[1]: import teqp, os, numpy as np, json
    display(teqp.__version__)
    '0.19.1'
[2]: path = teqp.get_datapath()+'/dev/fluids/n-Propane.json'
    assert (os.path.exists(path))
    jig = teqp.convert_CoolProp_idealgas(path, 0)
    print('As in the fluid file (matches Lemmon JPCRD 2009 exactly)::::')
    print(json.dumps(json.load(open(path))['EOS'][0]['alpha0'],indent=1))
    print('\n\nAnd after conversion::::')
    print(json.dumps(jig,indent=1))
    As in the fluid file (matches Lemmon JPCRD 2009 exactly)::::
    [
      "a1": -4.970583,
      "a2": 4.29352,
      "type": "IdealGasHelmholtzLead"
     {
      "a": 3,
      "type": "IdealGasHelmholtzLogTau"
     },
     {
      "n": [
       3.043,
       5.874,
       9.337,
       7.922
      ],
      "t": [
       1.062478,
       3.344237,
       5.363757,
       11.762957
      ],
      "type": "IdealGasHelmholtzPlanckEinstein"
    And after conversion::::
     "R": 8.314472,
     "terms": [
       "R": 8.314472,
       "a_1": -13.487776191416238,
       "a_2": 1588.1301128,
       "type": "Lead"
       },
       "R": 8.314472,
       "a": 17.739616992418114,
       "type": "Constant"
      },
       {
                                                                                (continues on next page)
```

```
"R": 8.314472,
  "a": -3.0,
  "type": "LogT"
  "R": 8.314472,
  "n": [
   3.043,
   5.874,
   9.337,
   7.922
  ],
  "theta": [
   392.99998742,
   1236.99982393,
   1984.0000767299998,
   4351.00016473
  ],
  "type": "PlanckEinstein"
]
}
```

Note that the two leading terms of Lemmon generates three terms in teqp because the forms of the terms are slightly different

```
[3]: # As an worked example, the conversions can be carried out like so, with the values_
    →from Lemmon given name of b instead of a
    b_1 = -4.970583
    b_2 = 4.29352
    Tr = 369.89 \# K
    rhor = 5000 \# mo1/m^3
    print('a_1:', b_1-np.log(rhor))
    print('a_2:', b_2*Tr)
    # The 3*ln(tau) term turns into 3*ln(Tr) - 3*ln(T)
    print(np.log(Tr)*3)
    # and the theta values are obtained
    t = np.array([1.062478, 3.344237, 5.363757, 11.762957])
    print((t*Tr).tolist())
    a_1: -13.487776191416238
    a_2: 1588.1301128
    17.739616992418114
    [392.99998742, 1236.99982393, 1984.0000767299998, 4351.00016473]
```

```
[4]: aig = teqp.IdealHelmholtz([jig])
   -aig.get_Ar20(300, 3, np.array([1.0]))
[4]: 7.863830967842212
```

72 Chapter 4. Models

CHAPTER

FIVE

ALGORITHMS

5.1 Phase equilibria

Two basic approaches are implemented in teqp:

- Iterative calculations given guess values
- · Tracing along iso-curves (constant temperature, etc.) powered by the isochoric thermodynamics formalism

```
[1]: import teqp
  import numpy as np
  import pandas
  import matplotlib.pyplot as plt
  teqp.__version__
[1]: '0.19.1'
```

5.1.1 Iterative Phase Equilibria

Pure fluid

For a pure fluid, phase equilibrium between two phases is defined by equating the pressures and Gibbs energies in the two phases. This represents a 2D non-linear rootfinding problem. Newton's method can be used for the rootfinding, and in teqp, automatic differentiation is used to obtain the necessary Jacobian matrix so the implementation is quite efficient.

The method requires guess values, which are the densities of the liquid and vapor densities. In some cases, ancillary or superancillary equations have been developed which provide curves of guess densities as a function of temperature.

For a pure fluid, you can use the pure_VLE_T method to carry out the iteration.

The Python method is here: pure VLE T

```
[2]: # Instantiate the model
  model = teqp.canonical_PR([300], [4e6], [0.1])

T = 250 # [K], Temperature to be used

# Here we use the superancillary to get guess values (actually these are more
# accurate than the results we will obtain from iteration!)
  rhoLO, rhoVO = model.superanc_rhoLV(T)
  display('guess:', [rhoLO, rhoVO])

# Carry out the iteration, return the liquid and vapor densities
```

```
# The guess values are perturbed to make sure the iteration is actually
# changing the values
model.pure_VLE_T(T, rhoL0*0.98, rhoV0*1.02, 10)

'guess:'
[12735.311173407898, 752.4082303122791]
[2]: array([12735.31117341, 752.40823031])
```

Binary Mixture

For a binary mixture, the approach is roughly similar to that of a pure fluid. The pressure is equated between phases, and the chemical potentials of each component in each phase are forced to be the same.

Again, the user is required to provide guess values, in this case molar concentrations in each phase, and a Newton method is implemented to solve for the phase equilibrium. The analytical Jacobian is obtained from automatic differentiation.

The mix_VLE_Tx function is the binary mixture analog to pure_VLE_T for pure fluids.

The Python method is here: mix_VLE_Tx

```
[3]: zA = np.array([0.01, 0.99])
    model = teqp.canonical_PR([300,310], [4e6,4.5e6], [0.1, 0.2])
    model1 = teqp.canonical_PR([300], [4e6], [0.1])
    T = 273.0 \# [K]
     # start off at pure of the first component
    rhoL0, rhoV0 = model1.superanc_rhoLV(T)
     # then we shift to the given composition in the first phase
    # to get guess values
    rhovecA0 = rhoL0*zA
    rhovecB0 = rhoV0*zA
    # carry out the iteration
    code, rhovecA, rhovecB = model.mix_VLE_Tx(T, rhovecA0, rhovecB0, zA,
         1e-10, 1e-10, 1e-10, 1e-10, # stopping conditions
         10 # maximum number of iterations
        )
    code, rhovecA, rhovecB
[3]: (<VLE_return_code.xtol_satisfied: 1>,
     array([ 128.66049209, 12737.38871682]),
     array([ 12.91868229, 1133.77242677]))
```

You can (and should) check the value of the return code to make sure the iteration succeeded. Do not rely on the numerical value of the enumerated return codes!

5.2 Tracing (isobars and isotherms)

When it comes to mixture thermodynamics, as soon as you add another component to a pure component to form a binary mixture, the complexity of the thermodynamics entirely changes. For that reason, mixture iterative calculations for mixtures are orders of magnitude more difficult to carry out. Asymmetric mixtures can do all sorts of interesting things that are entirely unlike those of pure fluids, and the algorithms are therefore much, much more complicated. Formulating phase equilibrium problems is not much more complicated than for pure fluids, but the most challenging aspect is to obtain good guess values from which to start an iterative routine, and the difficulty of this problem increases with the complexity of the mixture thermodynamics.

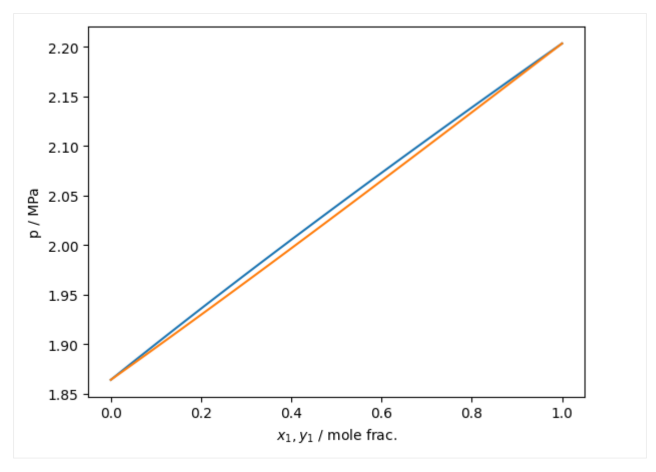
Ulrich Deiters and Ian Bell have developed a number of algorithms for tracing phase equilibrium solutions as the solution of ordinary differential equations rather than carrying out iterative routines for a given state point. The advantage of the tracing calculations is that they can often be initiated at a state point that is entirely known, for instance the pure fluid endpoint for a subcritical isotherm or isobar.

The Python method is here: trace_VLE_isotherm_binary

The C++ implementation returns a string in JSON format, which can be conveniently operated upon, for instance after converting the returned data structure to a pandas. DataFrame. A simple example of plotting a subcritical isotherm for a "boring" mixture is presented here:

```
[4]: model = tegp.canonical_PR([300,310], [4e6,4.5e6], [0.1, 0.2])
    model1 = teqp.canonical_PR([300], [4e6], [0.1])
    T = 273.0 \# [K]
    rhoL0, rhoV0 = model1.superanc_rhoLV(T) # start off at pure of the first component
    j = model.trace_VLE_isotherm_binary(T, np.array([rhoL0, 0]), np.array([rhoV0, 0]))
    display(str(j)[0:100]+'...') # The first few bits of the data
    df = pandas.DataFrame(j) # Now as a data frame
    df.head(3)
    "[{'T / K': 273.0, 'c': -1.0, 'drho/dt': [-0.618312383229212, 0.7690760182230469, -0.
     →1277526773161415..."
       T / K
[4]:
                                                                               dt \
                C
                                                                drho/dt
    0 \quad 273.0 \quad -1.0 \quad [-0.618312383229212, \quad 0.7690760182230469, \quad -0.12... \quad \quad 0.000010
    1 273.0 -1.0 [-0.6183123817120353, 0.7690760162922189, -0.1... 0.000045
       273.0 -1.0 [-0.6183123827116788, 0.7690760173388914, -0.1... 0.000203
                                                                   rhoL / mol/m^3
            pL / Pa
                           pV / Pa
       2.203397e+06 2.203397e+06
                                                       [10697.985891540735, 0.0]
       2.203397e+06 2.203397e+06 [10697.985885357639, 7.690760309421386e-06]
       2.203397e+06 2.203397e+06
                                     [10697.98585753358, 4.229918121248511e-05]
                                      rhoV / mol/m^3
                                                              t xL_0 / mole frac.
    0
                          [1504.6120879290752, 0.0]
                                                      0.000000
                                                                               1.0
       [1504.6120866515366, 9.945415375682985e-07] 0.000010
                                                                               1.0
    1
       [1504.6120809026731, 5.469978386095445e-06] 0.000055
                                                                               1.0
        xV_0 / mole frac.
    0
                      1.0
    1
                      1.0
    2
                      1.0
[5]: plt.plot(df['xL_0 / mole frac.'], df['pL / Pa']/1e6)
```

```
[5]: plt.plot(df['xL_0 / mole frac.'], df['pL / Pa']/1e6)
   plt.plot(df['xV_0 / mole frac.'], df['pL / Pa']/1e6)
   plt.gca().set(xlabel='$x_1,y_1$ / mole frac.', ylabel='p / MPa')
   plt.show()
```



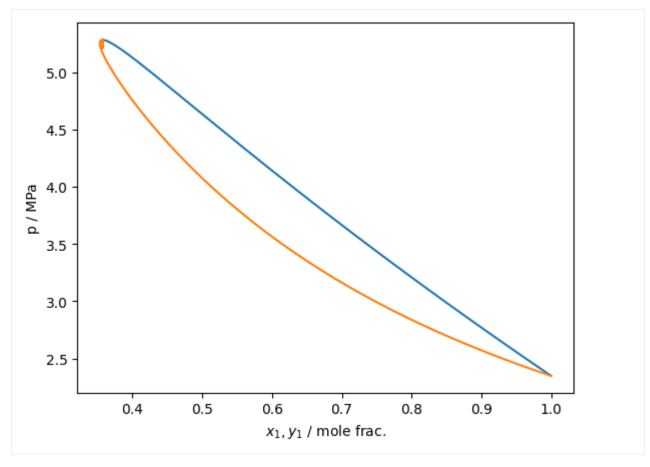
Isn't that exciting!

You can also provide an optional set of flags to the function to control other behaviors of the function, and switch between simple Euler and adaptive RK45 integration (the default)

The options class is here: TVLEOptions

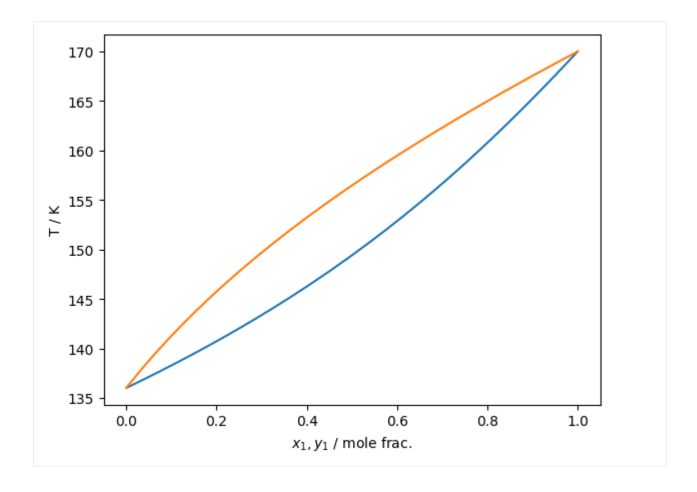
Supercritical isotherms work approximately in the same manner

```
[6]: Tc_K = [190.564, 154.581]
pc_Pa = [4599200, 5042800]
acentric = [0.011, 0.022]
model = teqp.canonical_PR(Tc_K, pc_Pa, acentric)
model1 = teqp.canonical_PR([Tc_K[0]], [pc_Pa[0]], [acentric[0]])
T = 170.0 # [K] # Note: above Tc of the second component
rhoL0, rhoV0 = model1.superanc_rhoLV(T) # start off at pure of the first component
j = model.trace_VLE_isotherm_binary(T, np.array([rhoL0, 0]), np.array([rhoV0, 0]))
df = pandas.DataFrame(j) # Now as a data frame
plt.plot(df['xL_0 / mole frac.'], df['pL / Pa']/1e6)
plt.plot(df['xV_0 / mole frac.'], df['pL / Pa']/1e6)
plt.gca().set(xlabel='$x_1,y_1$ / mole frac.', ylabel='p / MPa')
plt.show()
```



As of version 0.10.0, isobar tracing has been added to teqp. It operates in fundamentally the same fashion as the isotherm tracing and the same recommendations about starting at a pure fluid apply

The tracer function class is here: trace_VLE_isobar_binary



5.3 VLLE

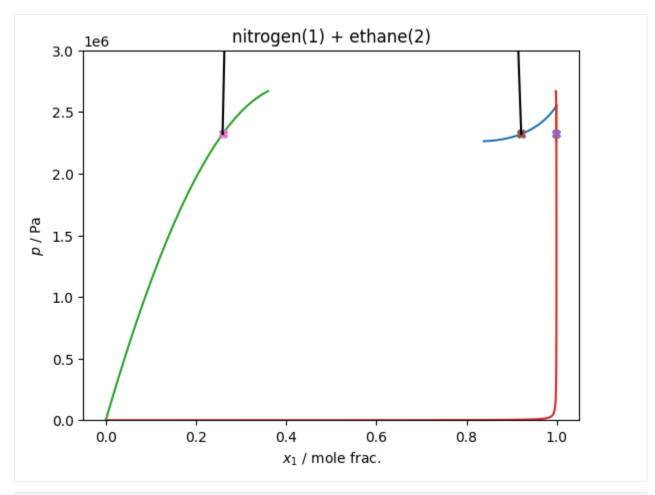
Following the approach described in Bell et al.: https://doi.org/10.1021/acs.iecr.1c04703

for the mixture of nitrogen + ethane, with the default thermodynamic model in teqp, which is the GERG-2008 mixing parameters (no departure function).

Two traces are made, and the intersection is obtained, this gives you the VLLE solution.

```
rhovecL[ipure] = rhoLpure
        rhovecV[ipure] = rhoVpure
        opt = teqp.TVLEOptions()
        opt.p_termination = 1e8
        opt.crit_termination=1e-4
        opt.calc_criticality=True
        j = model.trace_VLE_isotherm_binary(T, rhovecL, rhovecV, opt)
        traces.append(j)
   return model, traces
T = 120.3420
model, traces = get_traces(T=T, ipures=[0,1])
for trace in traces:
   df = pandas.DataFrame(trace)
   plt.plot(df['xL_0 / mole frac.'], df['pL / Pa'])
   plt.plot(df['xV_0 / mole frac.'], df['pV / Pa'])
# Do the VLLE solving
for soln in model.find_VLLE_T_binary(traces):
    print('rhovec / mol/m^3 | p / Pa')
    for rhovec in soln['polished']:
       rhovec = np.array(rhovec)
       rhotot = sum(rhovec)
       x = rhovec/rhotot
       p = rhotot*model.get_R(x)*T*(1+model.get_Ar01(T, rhotot, x))
       plt.plot(x[0], p, 'X')
       print(rhovec, p)
    # And also carry out the LLE trace for the two liquid phases
    j = model.trace_VLE_isotherm_binary(T, np.array(soln['polished'][1]), np.
→array(soln['polished'][2]))
   df = pandas.DataFrame(j)
    plt.plot(df['xL_0 / mole frac.'], df['pL / Pa'], 'k')
   plt.plot(df['xV_0 / mole frac.'], df['pV / Pa'], 'k')
# Plotting niceties
plt.ylim(top=3e6, bottom=0)
plt.gca().set(xlabel='$x_1$ / mole frac.', ylabel='$p$ / Pa', title='nitrogen(1) +_
→ethane(2)')
plt.show()
rhovec / mol/m^3 | p / Pa
[3.66984834e+03 3.25893958e+00] 2321103.087319132
[19890.16767481 1698.86505766] 2321103.087318946
[ 5641.24690517 16140.85769908] 2321103.0873195715
```

5.3. VLLE 79



```
[2]: # Trace from both pure fluid endpoints
    T = 113
    model, traces = get_traces(T=T, ipures = [0,1])
    # Find the VLLE solution for the starting temperature
    solns = model.find_VLLE_T_binary(traces)
    rhovecV, rhovecL1, rhovecL2 = solns[0]['polished']
    # Obtain the VLLE trace towards higher temperatures
    opt = teqp.VLLETracerOptions()
    a = lambda x: np.array(x)
    VLLE = model.trace_VLLE_binary(T, a(rhovecV), a(rhovecL1), a(rhovecL2), opt)
    df = pandas.DataFrame(VLLE)
    # Add the pressure to the DataFrame
    def add_ps(row, key):
        T = row['T / K']
        rhovec = np.array(row[key])
        rhotot = sum(rhovec)
        x = rhovec/rhotot
        p = rhotot*model.get_R(x)*T*(1+model.get_Ar01(T, rhotot, x))
    df['p / Pa'] = df.apply(add_ps, axis=1, key='rhoV / mol/m^3')
    # Plot the p-T curve
                                                                               (continues on next page)
```

```
plt.plot(df['T / K'], df['p / Pa'])
    plt.gca().set(xlabel='$T$ / K', ylabel='$p$ / Pa');
    plt.title('Nitrogen + ethane VLLE curve')
[2]: Text(0.5, 1.0, 'Nitrogen + ethane VLLE curve')
                              Nitrogen + ethane VLLE curve
             1e6
         4.5
         4.0
         3.5
         3.0
         2.5
         2.0
         1.5
                    115
                                  120
                                                125
                                                             130
                                                                            135
                                              T/K
```

5.4 VLLE @ constant pressure

Following the approach described in Bell et al.: https://doi.org/10.1021/acs.iecr.1c04703, but slightly different because the pressure is fixed rather than the temperature, but the same basic principles hold

for the mixture of nitrogen + ethane, with the default thermodynamic model in teqp, which is the GERG-2008 mixing parameters (no departure function).

Two traces are made, and the intersection is obtained, this gives you the VLLE solution.

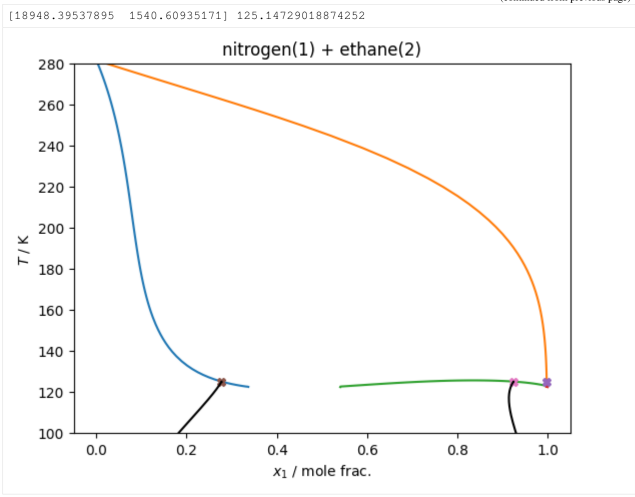
```
[1]: import teqp, numpy as np, matplotlib.pyplot as plt, pandas
import CoolProp.CoolProp as CP

names = ['Nitrogen', 'Ethane']
model = teqp.build_multifluid_model(names, teqp.get_datapath())
pures = [teqp.build_multifluid_model([name], teqp.get_datapath()) for name in names]
p = 29e5 # Pa

(continues on next page)
```

```
# Trace from both pure fluid endpoints
traces = []
for ipure in [1,0]:
    # Init at the pure fluid endpoint
    anc = pures[ipure].build_ancillaries()
   rhoLpure, rhoVpure = [CP.PropsSI('Dmolar','P',p,'Q',Q,names[ipure]) for Q in [0,
→1]]
   T = CP.PropsSI('T', 'P', p, 'Q', 0, names[ipure])
   rhovecL = np.array([0.0, 0.0])
   rhovecV = np.array([0.0, 0.0])
   rhovecL[ipure] = rhoLpure
   rhovecV[ipure] = rhoVpure
    j = model.trace_VLE_isobar_binary(p, T, rhovecL, rhovecV)
   df = pandas.DataFrame(j)
   plt.plot(df['xL_0 / mole frac.'], df['T / K'])
   plt.plot(df['xV_0 / mole frac.'], df['T / K'])
   traces.append(j)
# Do the VLLE solving
for soln in model.find_VLLE_p_binary(traces):
    T = soln['polished'][-1]
   print('rhovec / mol/m^3 | T / K')
    for rhovec in soln['polished'][0:3]:
       rhovec = np.array(rhovec)
        rhotot = sum(rhovec)
        x = rhovec/rhotot
        p = rhotot*model.get_R(x)*T*(1+model.get_Ar01(T, rhotot, x))
        plt.plot(x[0], T, 'X')
        print(rhovec, T)
    # And also carry out the LLE trace for the two liquid phases
   opt = tegp.PVLEOptions()
    opt.integration_order = 5
   opt.init_dt = 1e-10
    # Or could be 1 depending on the initial integration direction, do not know the.
→direction
    # a priori because not starting at a pure fluid endpoint
    for init_dt in [-1]:
        opt.init c = init dt
        rhovecV, rhovecL1, rhovecL2, T = soln['polished']
        j = model.trace_VLE_isobar_binary(p, T, np.array(rhovecL1), np.
→array(rhovecL2), opt)
        df = pandas.DataFrame(j)
        plt.plot(df['xL_0 / mole frac.'], df['T / K'], 'k')
        plt.plot(df['xV_0 / mole frac.'], df['T / K'], 'k')
# Plotting niceties
plt.ylim(top=280, bottom=100)
plt.gca().set(xlabel='$x_1$ / mole frac.', ylabel='$T$ / K', title='nitrogen(1) +_
\rightarrowethane (2)')
plt.show()
rhovec / mol/m^3 | T / K
                9.6755684 ] 125.14729018874252
[4921.97976373
[ 6008.68040253 15630.22353351] 125.14729018874252
```





```
[1]: import scipy.interpolate
  import teqp
  import numpy as np
  import pandas
  import matplotlib.pyplot as plt
  teqp.__version__
[1]: '0.19.1'
```

5.5 Critical curves & points

5.5.1 Pure Fluids

Solving for the critical point involves finding the temperature and density that make

$$\left(\frac{\partial p}{\partial \rho}\right)_T = \left(\frac{\partial^2 p}{\partial \rho^2}\right)_T = 0$$

by 2D non-linear rootfinding. Newton steps are taken, and the analytic Jacobian is used (thanks to the ability to do derivatives with automatic differentiation). This is all handily wrapped up in the <code>solve_pure_critical</code> method which requires the user to provide guess values for temperature and density

```
[2]: # Values taken from http://dx.doi.org/10.6028/jres.121.011
modelPR = teqp.canonical_PR([190.564], [4599200], [0.011])

# Solve for the critical point from a point close to the critical point
T0 = 192.0
# Critical compressibility factor of P-R is 0.307401308698.. (see https://doi.org/10.
--1021/acs.iecr.1c00847)
rhoc = (4599200/(8.31446261815324*190.564))/0.3074
rho0 = rhoc*1.2345 # Perturb to make sure we are doing something in the solver
modelPR.solve_pure_critical(T0, rho0)

[2]: (190.564, 9442.816240022832)
```

If you have a mixture, but want to obtain the critical point of a pure fluid of this mixture, you can specify the index of the component in the mixture, as well as the number of components in the mixture with something like:

```
model.solve_pure_critical(T0, rho0, {"alternative_pure_index": 1,
"alternative_length": 2}) so here, for the second fluid, with 0-based index of 1, in a two-component
mixture
```

5.5.2 Mixtures

A pure fluid has a single vapor-liquid critical point, but mixtures are different:

- They may have multiple (or zero!) critical points for a given mixture composition
- The critical curves may not emanate from the pure fluid endpoints

When it comes to critical points, intuition from pure fluids is not helpful, or sometimes even counter-productive.

teqp has methods for working with the critical loci of binary mixtures (only binary mixtures, for now) and especially, methods for tracing the critical curves emanating from the pure fluid endpoints.

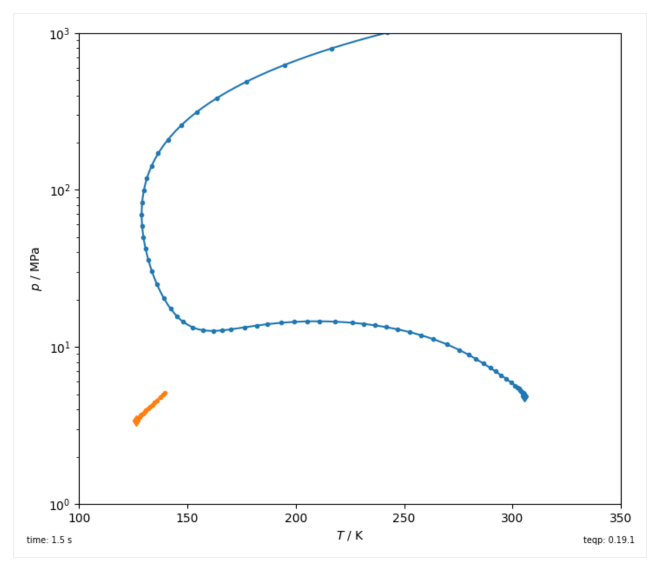
The tracing method in teqp is based explicitly on the isochoric thermodynamics formalism introduced by Ulrich Deiters and Sergio Quinones-Cisneros. It uses the Helmholtz energy density as the fundamental potential and all other properties are derived from it. For critical curves it is based upon the integration of sets of ordinary differential equations; the differential equations are in the form of derivatives of the molar concentrations of each component in the mixture with respect to an integration variable. The set of ODE is then integrated.

Here is an example of the construction of the critical curves emanating from the pure fluid endpoints for the mixture nitrogen + ethane.

```
import timeit
import numpy as np
import matplotlib.pyplot as plt
import pandas
import teqp

def get_critical_curve(ipure):
    """ Return curve as pandas DataFrame """
    names = ['Nitrogen', 'Ethane']
    model = teqp.build_multifluid_model(names, teqp.get_datapath())
    T0 = model.get_Tcvec()[ipure]
    rho0 = np.array([1.0/model.get_vcvec()[ipure]]*2)
    rho0[1-ipure] = 0
    o = teqp.TCABOptions()
    o.init_dt = 1.0 # step in the arclength tracing parameter
    o.rel_err = 1e-8
```

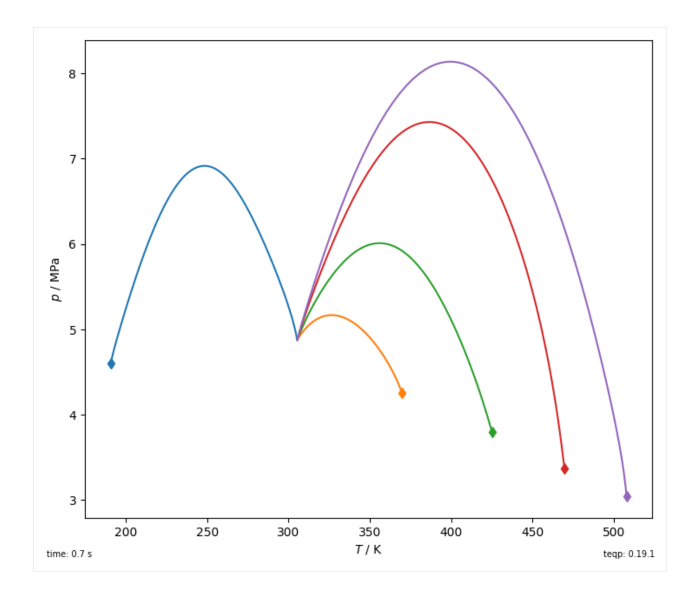
```
o.abs\_err = 1e-5
   o.integration_order = 5
   o.calc_stability = True
   o.polish = True
    curveJSON = model.trace_critical_arclength_binary(T0, rho0, '', o)
   df = pandas.DataFrame(curveJSON)
    rhotot = df['rho0 / mol/m^3']+df['rho1 / mol/m^3']
   df['z0 / mole frac.'] = df['rho0 / mol/m^3']/rhotot
   return df
fig, ax = plt.subplots(1, 1, figsize=(7, 6))
tic = timeit.default_timer()
for ipure in [1,0]:
   df = get_critical_curve(ipure)
   first_unstable = np.argmax(~df['locally stable'])
   df = df.iloc[0:(first_unstable if first_unstable else len(df))]
   line, = plt.plot(df['T / K'], df['p / Pa']/1e6, '.')
    # And interpolate to smooth out the curve using the arclength
    # parameter (which must be monotonically increasing) as
    # the interpolation variable
   tinterp = np.linspace(df['t'].min(), df['t'].max(), 10000)
   Tinterp = scipy.interpolate.interp1d(df['t'], df['T / K'], kind='cubic')(tinterp)
   pinterp = scipy.interpolate.interp1d(df['t'], df['p / Pa'], kind='cubic')(tinterp)
   plt.plot(Tinterp, pinterp/1e6, color=line.get_color())
   plt.plot(df['T / K'].iloc[0], df['p / Pa'].iloc[0]/1e6, 'd',
       color=line.get_color())
elap = timeit.default_timer()-tic
plt.gca().set(xlabel='$T$ / K', ylabel='$p$ / MPa',
    xlim=(100, 350), ylim=(1, 1e3))
plt.yscale('log')
plt.tight_layout(pad=0.2)
plt.gcf().text(0,0,f'time: {elap:0.1f} s', ha='left', va='bottom', fontsize=7)
plt.gcf().text(1,0,f'teqp: {teqp.__version__}}', ha='right', va='bottom', fontsize=7);
```



And now for something a bit more interesting: ethane + alkane critical curves

```
[4]: import timeit
    import numpy as np
    import matplotlib.pyplot as plt
    import pandas
    import teqp
    def get_critical_curve(names, ipure):
        """ Return curve as pandas DataFrame """
        model = teqp.build_multifluid_model(names, teqp.get_datapath())
        T0 = model.get_Tcvec()[ipure]
        rho0 = np.array([1.0/model.get_vcvec()[ipure]]*2)
        rho0[1-ipure] = 0
        o = teqp.TCABOptions()
          print(dir(o))
        o.init_dt = 1.0 # step in the parameter
        o.rel_err = 1e-6 # relative error on the step
        o.abs_err = 1e-6 # absolute error on the step
        o.max_dt = 100 # cap the size of the allowed step
```

```
o.calc_stability = True
   o.polish = True
   curveJSON = model.trace_critical_arclength_binary(T0, rho0, '', o)
   df = pandas.DataFrame(curveJSON)
    rhotot = df['rho0 / mol/m^3']+df['rho1 / mol/m^3']
    df['z0 / mole frac.'] = df['rho0 / mol/m^3']/rhotot
    return df
fig, ax = plt.subplots(1,1,figsize=(7, 6))
tic = timeit.default_timer()
name0 = 'ETHANE'
for othername in ['METHANE','PROPANE','BUTANE','PENTANE','HEXANE']:
    for ipure in [1]:
        df = get_critical_curve([name0, othername], ipure)
        line, = plt.plot(df['T / K'], df['p / Pa']/1e6, '-')
        plt.plot(df['T / K'].iloc[0], df['p / Pa'].iloc[0]/1e6, 'd',
            color=line.get_color())
elap = timeit.default_timer()-tic
plt.gca().set(xlabel='$T$ / K', ylabel='$p$ / MPa')#,xlim=(100, 350), ylim=(1, 1e3))
plt.tight_layout(pad=0.2)
plt.gcf().text(0,0,f'time: {elap:0.1f} s', ha='left', va='bottom', fontsize=7)
plt.gcf().text(1,0,f'teqp: {teqp.__version__}}', ha='right', va='bottom', fontsize=7);
```



5.5.3 Pure fluid EOS with nonanalytic terms

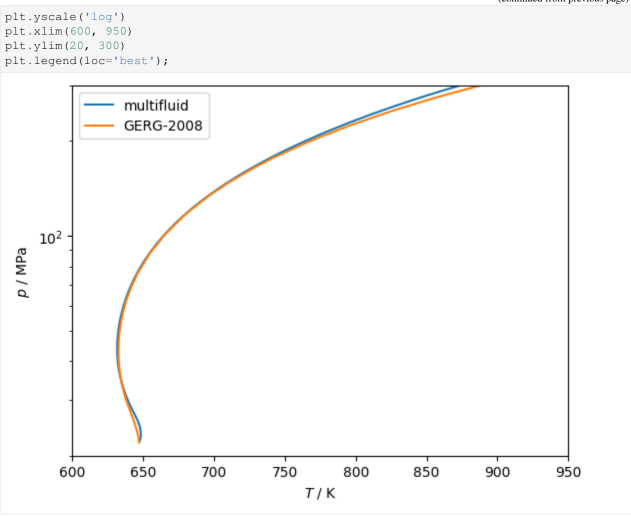
For the highest accuracy EOS for normal water and carbon dioxide, there are non-analytic terms that prevent the initialization of the critical tracing at the pure fluid critical point. Instead, one can start close to, but not AT, the pure fluid endpoint. After deciding on that starting composition, one solves for the critical point and then traces away from it.

You might need to either do tracing in two parts, one with init_c=+1 and then init_c=-1, or one tracing might be good enough.

Here is an example:

```
[5]: def get_critical_curve_composition(names, T0, rhovec0, init_c=-1):
         """ Trace the critical curve from a fixed point along it """
        o = teqp.TCABOptions()
          print(dir(o))
        o.init_dt = 1.0 # step in the parameter
        o.rel_err = 1e-6 # relative error on the step
         o.abs_err = 1e-6 # absolute error on the step
                                                                                (continues on next page)
```

```
o.max_dt = 100 # cap the size of the allowed step
   o.calc_stability = True
   o.polish = True
    o.init_c = init_c # You might need to swap the initial tracing direction by_
→ making this +1.0
    curveJSON = model.trace_critical_arclength_binary(T0, rhovec0, '', o)
    df = pandas.DataFrame(curveJSON)
    rhotot = df['rho0 / mo1/m^3'] + df['rho1 / mo1/m^3']
    df['z0 / mole frac.'] = df['rho0 / mol/m^3']/rhotot
    return df
# Tracing with multi-fluid from an endpoint with non-analytic terms
model = teqp.build_multifluid_model(["Water", "Methane"], teqp.get_datapath())
x0 = 1-1e-6 # ever so slightly away from the pure fluid
molefrac = np.array([x0, 1-x0])
# Solve for the actual critical point at this mole fraction with scipy
y0 = [model.get_Tcvec()[0], 1/model.get_vcvec()[0]]
\texttt{residual} = \textbf{lambda} \ y \colon \texttt{model.get\_criticality\_conditions} \ (y \ [0], \ y \ [1] \ * molefrac)
res = scipy.optimize.fsolve(residual, y0)
T = res[0]
rho0 = res[1]
rhovec0 = rho0*molefrac
# Now trace from this point
curve = get_critical_curve_composition(model, T0=T, rhovec0=rhovec0)
plt.plot(curve['T / K'], curve['p / Pa']/1e6, label='multifluid')
# With GERG-2008, things are much more straightforward...
model = teqp.make_model({'kind': 'GERG2008resid', 'model': {'names': ['water', 'methane
' ] } } )
def get_critical_curve_simple(model, ipure, T0, rho0):
    """ Trace from a pure fluid... """
   rhovec0 = np.array([0, 0])
   rhovec0[ipure] = rho0
   o = teqp.TCABOptions()
   o.init_dt = 1.0 # step in the arclength tracing parameter
   o.rel_err = 1e-8
   o.abs\_err = 1e-5
   o.integration_order = 5
   o.calc_stability = True
   o.polish = True
    curveJSON = model.trace_critical_arclength_binary(T0, rhovec0, '', o)
    df = pandas.DataFrame(curveJSON)
    rhotot = df['rho0 / mo1/m^3'] + df['rho1 / mo1/m^3']
    df['z0 / mole frac.'] = df['rho0 / mol/m^3']/rhotot
    return df
for ifluid in [0]:
   Tci = model.get_Tcvec()[ifluid]
   vci = model.get_vcvec()[ifluid]
   df = get_critical_curve_simple(model, ipure=ifluid, T0=Tci, rho0 = 1.0/vci)
    plt.plot(df['T / K'], df['p / Pa']/1e6, label='GERG-2008')
plt.gca().set(xlabel='$T$ / K', ylabel='$p$ / MPa')
                                                                            (continues on next page)
```



5.6 Information

The algorithms are written in a very generic way; they take an instance of a thermodynamic model, and the necessary derivatives are calculated from this model with automatic differentiation (or similar). In that way, implementing a model is all that is required to enable its use in the calculation of critical curves or to trace the phase equilibria. Determining the starting values, on the other hand, may require model-specific assistance, for instance with superancillary equations.

CHAPTER

SIX

EXAMPLES

6.1 The teqp paper in I&ECR

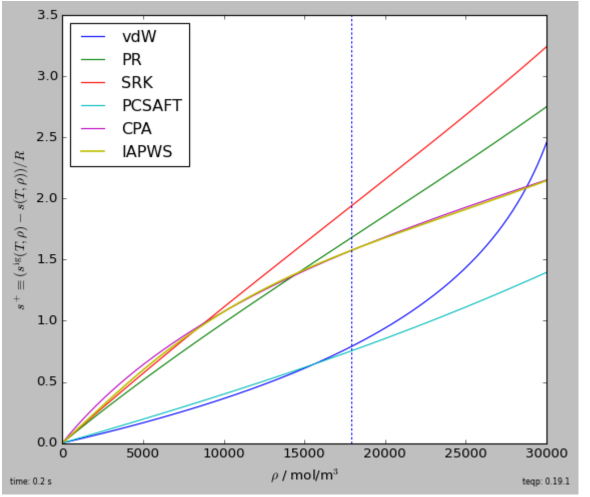
A few minor changes have been made:

- The get_splus method requires the molar concentrations to be a numpy array (to avoid copies) (as of version 0.14.0
- ullet The top-level methods tegp.xxx have been deprecated, and the methods attached to the instance are preferred
- The radial_dist field must always be provided

```
[1]: import timeit, numpy as np
    import matplotlib.pyplot as plt
    plt.style.use('classic')
    import teqp
    def build_models():
        Tc_K, pc_Pa, acentric = 647.096, 22064000.0, 0.3442920843
        water = {
            "a0i / Pa m^6/mol^2": 0.12277 , "bi / m^3/mol": 0.000014515, "c1": 0.67359,
            "Tc / K": 647.096, "epsABi / J/mol": 16655.0, "betaABi": 0.0692, "class": "4C"
        j = {"cubic": "SRK", "pures": [water], "R_gas / J/mol/K": 8.3144598, "radial_dist

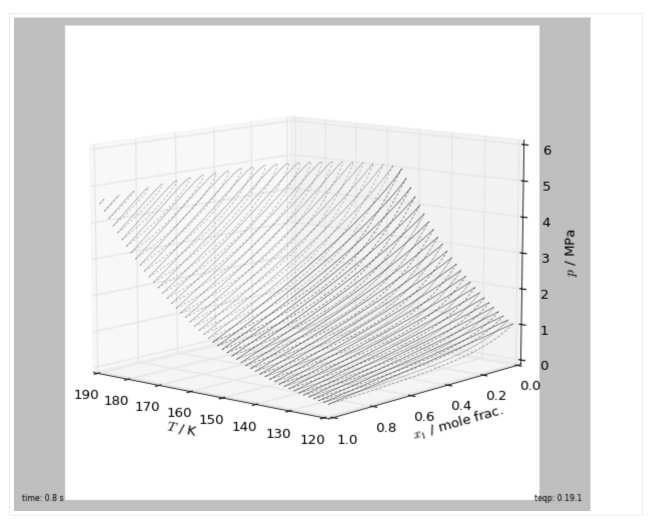
    ": "CS"}
        datapath = teqp.get_datapath()
        def get_PCSAFT():
            c = teqp.SAFTCoeffs()
             # Values from https://doi.org/10.1016/j.fluid.2017.11.015,
             # but association contribution is ignored
            c.name = 'Water'
            c.m = 2.5472
            c.sigma\_Angstrom = 2.1054
            c.epsilon_over_k = 138.63
            return teqp.PCSAFTEOS(coeffs=[c])
        return [
             ('vdW', teqp.vdWEOS([Tc_K], [pc_Pa])),
             ('PR', teqp.canonical_PR([Tc_K], [pc_Pa], [acentric])),
             ('SRK', teqp.canonical_SRK([Tc_K], [pc_Pa], [acentric])),
             ('PCSAFT', get_PCSAFT()),
             ('CPA', teqp.CPAfactory(j)),
             ('IAPWS', teqp.build_multifluid_model(["Water"], datapath))
```

```
fig, ax = plt.subplots(1,1,figsize=(7,6))
T = 700 \# K
rhovec = np.geomspace(0.1, 30e3, 10000) # mol/m^3; critical density is 17873.8... mol/
→m^3
tic = timeit.default_timer()
for abbrv, model in build_models():
    splus = np.array([model.get_splus(T, np.array([rho])) for rho in rhovec])
    plt.plot(rhovec, splus, label=abbrv, lw = 1.5 if abbrv=='IAPWS' else 1)
elap = timeit.default_timer()-tic
plt.axvline(17873.8, dashes=[2,2])
plt.legend(loc='best')
plt.gca().set(xlabel=r'\$\rho\$ / mol/m\$^3\$', ylabel=r'\$s^+\equiv (s^{\rm ig}(T,\rho)-
\hookrightarrows(T,\rho))/R$')
plt.tight_layout(pad=0.2)
plt.gcf().text(0,0,f'time: {elap:0.1f} s', ha='left', va='bottom', fontsize=7)
plt.gcf().text(1,0,f'teqp: {teqp.__version__}', ha='right', va='bottom', fontsize=7)
plt.savefig('splus_water_700K.pdf')
plt.show()
```



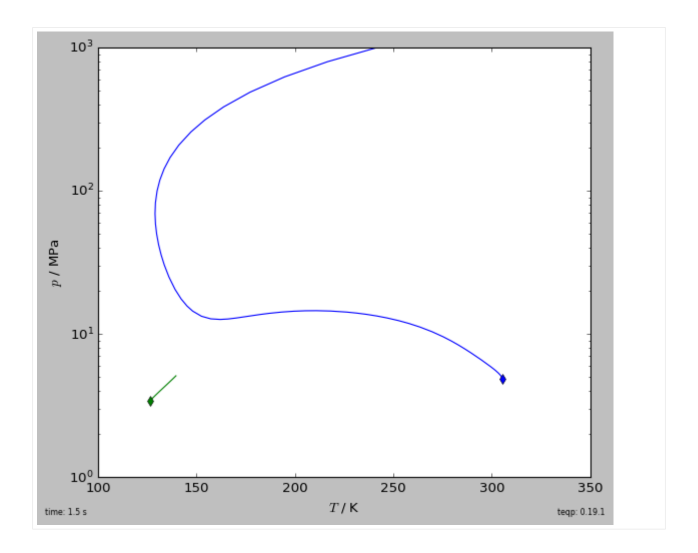
[2]: import json, timeit

```
import pandas, numpy as np, matplotlib.pyplot as plt
plt.style.use('classic')
import teqp
Tc_K = [190.564, 154.581]
pc_Pa = [4599200, 5042800]
acentric = [0.011, 0.022]
model = teqp.canonical_PR(Tc_K, pc_Pa, acentric)
fig, ax = plt.subplots(1,1,figsize=(7, 6), subplot_kw=dict(projection='3d'))
tic = timeit.default_timer()
for ifluid in [0,1]:
   model0 = teqp.canonical_PR([Tc_K[ifluid]], [pc_Pa[ifluid]], [acentric[ifluid]])
    for T in np.linspace(190, 120, 50):
        if T > Tc_K[ifluid]: continue
        [rhoL, rhoV] = model0.superanc_rhoLV(T)
        rhovecL = np.array([0.0, 0.0]); rhovecL[ifluid] = rhoL
        rhovecV = np.array([0.0, 0.0]); rhovecV[ifluid] = rhoV
        opt = teqp.TVLEOptions(); opt.calc_criticality = True
        df = pandas.DataFrame(model.trace_VLE_isotherm_binary(T, rhovecL, rhovecV,__
→opt))
        df['too_critical'] = df.apply(
            lambda row: (abs(row['crit. conditions L'][0]) < 5e-8), axis=1)
        first_too_critical = np.argmax(df['too_critical'])
        df = df.iloc[0:(first_too_critical if first_too_critical else len(df))]
        line, = ax.plot(xs=df['T / K'], ys=df['xL_0 / mole frac.'], zs=df['pL / Pa']/
→1e6,
            lw=0.2, color='k')
        ax.plot(xs=df['T / K'], ys=df['xV_0 / mole frac.'], zs=df['pL / Pa']/1e6,
            dashes=[2,2], color=line.get_color(), lw=0.2)
elap = timeit.default_timer()-tic
ax.view_init(elev=10., azim=130)
ax.set(xlabel='$T$ / K', ylabel='$x_1$ / mole frac.', zlabel='$p$ / MPa')
fig.text(0,0,f'time: {elap:0.1f} s', ha='left', va='bottom', fontsize=7)
fig.text(1,0,f'teqp: {teqp.__version__})', ha='right', va='bottom', fontsize=7)
plt.tight_layout(pad=0.2)
plt.savefig('PR_VLE_trace.pdf')
plt.show()
```



```
[3]: import timeit
    import numpy as np
    import matplotlib.pyplot as plt
    plt.style.use('classic')
    import pandas
    import teqp
    def get_critical_curve(ipure):
        """ Return curve as pandas DataFrame """
        names = ['Nitrogen', 'Ethane']
        model = teqp.build_multifluid_model(names, teqp.get_datapath())
        T0 = model.get_Tcvec()[ipure]
        rho0 = np.array([1.0/model.get_vcvec()[ipure]]*2)
        rho0[1-ipure] = 0
        o = teqp.TCABOptions()
        o.init_dt = 1.0 # step in the parameter
        o.rel\_err = 1e-8
        o.abs\_err = 1e-5
        o.integration_order = 5
        o.calc_stability = True
        o.polish = True
        curveJSON = model.trace_critical_arclength_binary(T0, rho0, '', o)
```

```
df = pandas.DataFrame(curveJSON)
   rhotot = df['rho0 / mo1/m^3'] + df['rho1 / mo1/m^3']
   df['z0 / mole frac.'] = df['rho0 / mol/m^3']/rhotot
    return df
if __name__ == '__main__':
    fig, ax = plt.subplots(1,1,figsize=(7, 6))
    tic = timeit.default_timer()
    for ipure in [1,0]:
        df = get_critical_curve(ipure)
        first_unstable = np.argmax(~df['locally stable'])
        df = df.iloc[0:(first_unstable if first_unstable else len(df))]
        line, = plt.plot(df['T / K'], df['p / Pa']/1e6, '-')
        plt.plot(df['T / K'].iloc[0], df['p / Pa'].iloc[0]/1e6, 'd',
            color=line.get_color())
   elap = timeit.default_timer()-tic
    plt.gca().set(xlabel='$T$ / K', ylabel='$p$ / MPa',
        xlim=(100, 350), ylim=(1, 1e3))
   plt.yscale('log')
   plt.tight_layout(pad=0.2)
    plt.gcf().text(0,0,f'time: {elap:0.1f} s', ha='left', va='bottom', fontsize=7)
   plt.gcf().text(1,0,f'teqp: {teqp.__version__}', ha='right', va='bottom',_
\rightarrowfontsize=7)
    plt.savefig('N2_ethane_critical.pdf')
    plt.show()
```



CHAPTER

SEVEN

FITTING

7.1 Multi-fluid Parameter Fitting

Here is an example of fitting the β_T and γ_T values for the binary pair of propane+n-dodecane with the multi-fluid model. It uses differential evolution to do the global optimization, which is probably overkill in this case as the problem is 2D and other algorithms like Nelder-Mead or even approximate Hessian methods would probably be fine.

In any case, it takes a few seconds to run (when the actual optimization is uncommented), demonstrating how one can fit model parameters with existing tooling from the scientific python stack.

```
[1]: import json
    import teqp, numpy as np, pandas, matplotlib.pyplot as plt
    import scipy.interpolate, scipy.optimize
    import pandas
    data = pandas.read_csv('VLE_data_propane_dodecane.csv')
```

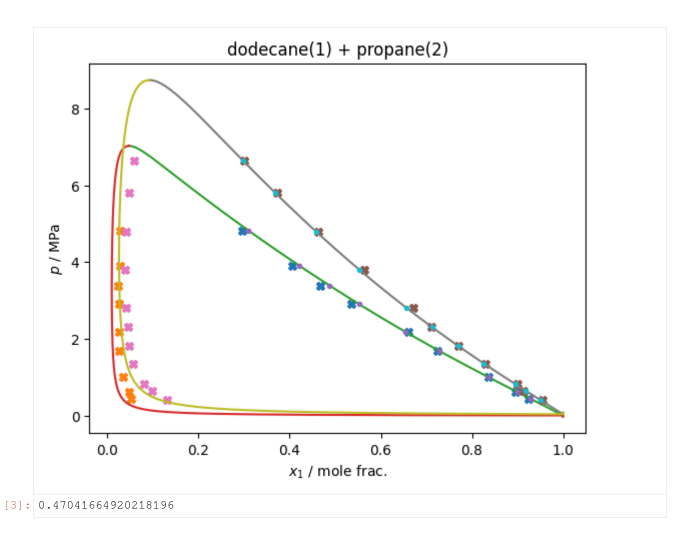
```
[2]: def cost_function(parameters:np.ndarray, plot:bool=False):
        # Fitting some parameters and fixing the others
        betaV, gammaV = 1.0, 1.0
        betaT, gammaT = parameters
         # betaT, gammaT, betaV, gammaV = parameters
        BIP = [{
             'function': '',
             'BibTeX': 'thiswork',
             'CAS1': '112-40-3',
             'CAS2': '74-98-6',
             'F': 0.0,
             'Name1': 'n-Dodecane',
             'Name2': 'n-Propane',
             'betaT': betaT,
             'betaV': betaV,
             'gammaT': gammaT,
             'gammaV': gammaV
        }]
        model = teqp.build_multifluid_model(["n-Dodecane", "n-Propane"], teqp.get_

→datapath(),
            BIPcollectionpath=json.dumps(BIP)
        ancs = [model.build_ancillaries(ipure) for ipure in [0,1]]
```

```
cost = 0.0
   # The O-based index of the fluid to start from. At this temperature, only one.
\hookrightarrowfluid
   # is subcritical, so it has to be that one, but in general you could start
   # from either one.
   ipure = 0
   for T in [419.15, 457.65]:
       # Subset the experimental data to match the isotherm
        # being fitted
       dfT = data[np.abs(data['T / K90'] - T) < 1e-3]
       if plot:
           plt.plot(1-dfT['x[0] / mole frac.'], dfT['p / Pa']/1e6, 'X')
           plt.plot(1-dfT['y[0] / mole frac.'], dfT['p / Pa']/1e6, 'X')
           # Get the molar concentrations of the pure fluid
            # at the starting point
           anc = ancs[ipure]
           rhoL0 = np.array([0, 0.0])
           rhoV0 = np.array([0, 0.0])
           rhoL0[ipure] = anc.rhoL(T)
           rhoV0[ipure] = anc.rhoV(T)
           # Now we do the trace and convert retuned JSON
            # data into a DataFrame
           df = pandas.DataFrame(model.trace_VLE_isotherm_binary(T, rhoL0, rhoV0))
           if plot:
               plt.plot(df['xL_0 / mole frac.'], df['pL / Pa']/1e6)
               plt.plot(df['xV_0 / mole frac.'], df['pL / Pa']/1e6)
            # Interpolate trace at experimental pressures along this
            # isotherm to get composition from the current model
            # The interpolators are set up to put in NaN for out
            # of range values
           x_interpolator = scipy.interpolate.interp1d(
               df['pL / Pa'], df['xL_0 / mole frac.'],
               fill_value=np.nan, bounds_error=False
           y_interpolator = scipy.interpolate.interp1d(
               df['pL / Pa'], df['xV_0 / mole frac.'],
               fill_value=np.nan, bounds_error=False
           # The interpolated values for the compositions
            # along the trace at experimental pressures
           x_model = x_interpolator(dfT['p / Pa'])
           y_model = y_interpolator(dfT['p / Pa'])
           if plot:
               plt.plot(x_model, dfT['p / Pa']/1e6, '.')
            # print(x_model, (1-dfT['x[0] (-)']))
           errTx = np.sum(np.abs(x_model-(1-dfT['x[0] / mole frac.']))))
                                                                          (continues on next page)
```

98 Chapter 7. Fitting

```
errTy = np.sum(np.abs(y_model-(1-dfT['y[0] / mole frac.'])))
           # If any point *cannot* be interpolated, throw out the model,
           # returning a large cost function value.
           # Note: you might need to be more careful here,
           # if the points are close to the critical point, a good model might
           # (but not usually), undershoot the critical point of the
           # real mixture
           # Also watch out for values of compositons in the data that are...
→placeholders
           # with a value of nan, which will pollute the error calculation
           if not np.isfinite(errTx):
               return 1e6
           if not np.isfinite(errTy):
               return 1e6
           cost += errTx + errTy
       except BaseException as BE:
           print (BE)
           pass
   if plot:
       plt.title(f'dodecane(1) + propane(2)')
       plt.xlabel('$x_1$ / mole frac.'); plt.ylabel('$p$ / MPa')
       plt.savefig('n-Dodecane+propane.pdf')
       plt.show()
   return cost
```



100 Chapter 7. Fitting

CHAPTER

EIGHT

TEQP

8.1 teqp package

8.1.1 Submodules

8.1.2 teqp.teqp module

```
TEQP: Templated Equation of State Package
class teap.teap.AbstractModel
      Bases: pybind11_object
      build_Psi_Hessian_autodiff(self: teqp.teqp.AbstractModel, T: float, rhovec:
                                             numpy.ndarray[numpy.float64[m, 1]]) \rightarrow
                                             numpy.ndarray[numpy.float64[m, n]]
      build Psir Hessian autodiff (self: teqp.teqp.AbstractModel, T: float, rhovec:
                                              numpy.ndarray[numpy.float64[m, 1]]) \rightarrow
                                              numpy.ndarray[numpy.float64[m, n]]
      build_Psir_gradient_autodiff(self: teqp.teqp.AbstractModel, T: float, rhovec:
                                                numpy.ndarray[numpy.float64[m, 1]]) \rightarrow
                                                numpy.ndarray[numpy.float64[m, 1]]
      build_d2PsirdTdrhoi_autodiff(self: teqp.teqp.AbstractModel, T: float, rhovec:
                                                numpy.ndarray[numpy.float64[m, 1]]) \rightarrow
                                                numpy.ndarray[numpy.float64[m, 1]]
      dpsatdT pure (self: teqp.teqp.AbstractModel, T: float, rhoL: float, rhoV: float) \rightarrow float
      eigen_problem (self: teqp.teqp.AbstractModel, T: float, rhovec: numpy.ndarray[numpy.float64[m, 1]],
                          alignment\_v0: numpy.ndarray[numpy.float64[m, 1]] | None = None) \rightarrow teqp::EigenData
      extrapolate from critical (self: teqp.teqp.AbstractModel, Tc: float, rhoc: float, T: float, molefrac:
                                           numpy.ndarray[numpy.float64[m, 1]] \mid None = None) \rightarrow
                                           numpy.ndarray[numpy.float64[2, 1]]
      find VLLE T binary (self: teqp.teqp.AbstractModel, traces: List[ison], options:
                                 teqp.teqp.VLLEFinderOptions | None = None) \rightarrow List[json]
      find VLLE p binary (self: teqp.teqp.AbstractModel, traces: List[json], options:
                                 teqp.teqp.VLLEFinderOptions | None = None) \rightarrow List[json]
```

- **get_ATrhoXi** (self: teqp.teqp.AbstractModel, T: float, NT: int, rhomolar: float, Nrho: int, molefrac: numpy.ndarray[numpy.float64[m, 1]], i: int, NXi: int) \rightarrow float
- **get_ATrhoXiXj** (self: teqp.teqp.AbstractModel, T: float, NT: int, rhomolar: float, Nrho: int, molefrac: numpy.ndarray[numpy.float64[m, 1]], i: int, NXi: int, j: int, NXj: int) \rightarrow float
- $\texttt{get_ATrhoXiXjXk}$ (self: teqp.teqp.AbstractModel, T: float, NT: int, rhomolar: float, Nrho: int, molefrac: numpy.ndarray[numpy.float64[m, 1]], i: int, NXi: int, j: int, NXj: int, k: int, NXk: int) \rightarrow float
- get_Ar00 (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
- get_Ar00n (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) → numpy.ndarray[numpy.float64[m, 1]]
- get_Ar01 (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
- get_Ar01n (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow numpy.ndarray[numpy.float64[m, 1]]
- get_Ar02 (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
- get_Ar02n (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow numpy.ndarray[numpy.float64[m, 1]]
- $\texttt{get_Ar03}$ (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
- get_Ar03n (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow numpy.ndarray[numpy.float64[m, 1]]
- $\texttt{get_Ar04}$ (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
- get_Ar04n (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow numpy.ndarray[numpy.float64[m, 1]]
- get_Ar05n (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow numpy.ndarray[numpy.float64[m, 1]]
- $\label{eq:get_Ar06n} \textit{get_Ar06n} (\textit{self:} \ \text{teqp.teqp.AbstractModel}, \ \textit{T:float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]}) \rightarrow \text{numpy.ndarray[numpy.float64[m, 1]]}$
- $\label{eq:get_Ar10} \begin{tabular}{l} \textbf{get_Ar10} (self: teqp.teqp.AbstractModel, T: float, r ho: float, m olefrac: $numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float \\ \end{tabular}$
- $\label{eq:get_Ar11} \textit{(self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]])} \rightarrow \textit{float}$
- $\label{eq:get_Ar12} \begin{tabular}{l} \textbf{get_Ar12} (self: teqp.teqp.AbstractModel, T: float, r ho: float, m olefrac: $numpy.ndarray[numpy.float64[m, 1]]) $$ \to float $$$

102 Chapter 8. teqp

```
get_Ar13 (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, I]]) → float

Tell (self: teap teap AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, I]])
```

- $\texttt{get_Ar14}$ (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
- $\texttt{get_Ar20}$ (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
- $\texttt{get_Ar21}$ (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, I]I) \rightarrow float
- $\texttt{get_Ar22}$ (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
- get_Ar23 (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
- get_Ar24 (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
- **get_Arxy** (*self:* teqp.teqp.AbstractModel, *NT: int*, *ND: int*, *T: float*, *rho: float*, *molefrac:* $numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float$
- $\texttt{get_AtaudeltaXi}$ (self: teqp.teqp.AbstractModel, tau: float, Ntau: int, delta: float, Ndelta: int, molefrac: numpy.ndarray[numpy.float64[m, 1]], i: int, NXi: int) \rightarrow float
- **get_AtaudeltaXiXj** (self: teqp.teqp.AbstractModel, tau: float, Ntau: int, delta: float, Ndelta: int, molefrac: numpy.ndarray[numpy.float64[m, 1]], i: int, NXi: int, j: int, NXj: int) \rightarrow float
- $\label{eq:continuity} \textbf{get_AtaudeltaXiXjXk} \ (self: teqp.teqp.AbstractModel, tau: float, Ntau: int, delta: float, Ndelta: int, molefrac: numpy.ndarray[numpy.float64[m, 1]], i: int, NXi: int, j: int, NXj: int, k: int, NXk: int) \ \rightarrow \ \text{float}$
- $\texttt{get_B12vir} (\textit{self:} \texttt{teqp.teqp.AbstractModel}, \textit{T:} \textit{float}, \textit{molefrac:} \textit{numpy.ndarray}[\textit{numpy.float64[m, 1]]}) \rightarrow \texttt{float}$
- **get_B2vir** (self: teqp.teqp.AbstractModel, T: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
- $\label{eq:get_Bnvir} \begin{subarray}{l} \textbf{get_Bnvir} (self: teqp.teqp.AbstractModel, Nderiv: int, T: float, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow Dict[int, float] \\ \end{subarray}$
- \texttt{get}_{R} (self: teqp.teqp.AbstractModel, molefrac: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
- $\begin{tabular}{ll} \begin{tabular}{ll} \beg$
- $\begin{tabular}{ll} \begin{tabular}{ll} \beg$
- $\begin{tabular}{ll} \begin{tabular}{ll} $\tt get_dchempotdT_autodiff} (self: teqp.teqp.AbstractModel, $T:$ float, $rhovec: numpy.ndarray[numpy.float64[m, 1]]) $\to $$ numpy.ndarray[numpy.float64[m, 1]] $$ \end{tabular}$

8.1. tegp package 103

```
get deriv mat2 (self: tegp.tegp.AbstractModel, T: float, rho: float, molefrac:
                      numpy.ndarray[numpy.float64[m, 1]]) \rightarrow numpy.ndarray[numpy.float64[3, 3]]
get dmBnvirdTm (self: tegp.tegp.AbstractModel, Nderiv: int, NTderiv: int, T: float, molefrac:
                      numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
\texttt{get\_dp\_dT\_crit} (self: teqp.teqp.AbstractModel, T: float, rhovec: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow
get_dpsat_dTsat_isopleth (self: teqp.teqp.AbstractModel, T: float, rhovecL:
                                     numpy.ndarray[numpy.float64[m, 1]], rhovecV:
                                     numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
get_drhovec_dT_crit (self: teqp.teqp.AbstractModel, T: float, rhovec: numpy.ndarray[numpy.float64[m,
                             1]) \rightarrow numpy.ndarray[numpy.float64[m, 1]]
get_drhovecdT_psat (self: teqp.teqp.AbstractModel, T: float, rhovecL: numpy.ndarray[numpy.float64[m,
                            1]], rhovecV: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow
                            Tuple[numpy.ndarray[numpy.float64[m, 1]], numpy.ndarray[numpy.float64[m, 1]]]
get_drhovecdp_Tsat (self: teqp.teqp.AbstractModel, T: float, rhovecL: numpy.ndarray[numpy.float64[m,
                            1]], rhovecV: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow
                            Tuple[numpy.ndarray[numpy.float64[m, 1]], numpy.ndarray[numpy.float64[m, 1]]]
get_fugacity_coefficients (self: teqp.teqp.AbstractModel, T: float, rhovec:
                                      numpy.ndarray[numpy.float64[m, 1]]) \rightarrow
                                      numpy.ndarray[numpy.float64[m, 1]]
get_minimum_eigenvalue_Psi_Hessian (self: teqp.teqp.AbstractModel, T: float, rhovec:
                                                   numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
get_neff (self: teqp.teqp.AbstractModel, T: float, rho: float, molefrac: numpy.ndarray[numpy.float64[m,
             1]]) \rightarrow float
get_partial_molar_volumes (self: teqp.teqp.AbstractModel, T: float, rhovec:
                                      numpy.ndarray[numpy.float64[m, 1]]) \rightarrow
                                      numpy.ndarray[numpy.float64[m, 1]]
get_pr (self: teqp.teqp.AbstractModel, T: float, rhovec: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
get_pure_critical_conditions_Jacobian (self: teqp.teqp.AbstractModel, T: float, rho: float,
                                                        alternative pure index: int | None = None,
                                                        alternative\_length: int \mid None = None) \rightarrow
                                                        Tuple[numpy.ndarray[numpy.float64[m, 1]],
                                                        numpy.ndarray[numpy.float64[m, n]]]
get_splus (self: teqp.teqp.AbstractModel, T: float, rhovec: numpy.ndarray[numpy.float64[m, 1]]) \rightarrow float
mix_VLE_Tp (self: teqp.teqp.AbstractModel, T: float, p_given: float, rhovecL0:
                numpy.ndarray[numpy.float64[m, 1]], rhovecV0: numpy.ndarray[numpy.float64[m, 1]],
                options: teqp.teqp.MixVLETpFlags | None = None) \rightarrow teqp.teqp.MixVLEReturn
mix_VLE_Tx (self: teqp.teqp.AbstractModel, T: float, rhovecL0: numpy.ndarray[numpy.float64[m, 1]],
                rhovecV0: numpy.ndarray[numpy.float64[m, 1]], xspec: numpy.ndarray[numpy.float64[m, 1]],
                atol: float, reltol: float, axtol: float, relxtol: float, maxiter: int) \rightarrow
                Tuple[teap.teap.VLE_return_code, numpy.ndarray[numpy.float64[m, 1]],
                numpy.ndarray[numpy.float64[m, 1]]]
```

104 Chapter 8. teqp

```
mix VLLE_T (self: teqp.teqp.AbstractModel, T: float, rhovecVinit: numpy.ndarray[numpy.float64[m, 1]],
                     rhovecL1init: numpy.ndarray[numpy.float64[m, 1]], rhovecL2init:
                     numpy.ndarray[numpy.float64[m, 1]], atol: float, reltol: float, axtol: float, relxtol: float, maxiter:
                     int) → Tuple[teqp::VLLE::VLLE_return_code, numpy.ndarray[numpy.float64[m, 1]],
                     numpy.ndarray[numpy.float64[m, 1]], numpy.ndarray[numpy.float64[m, 1]]]
     mixture_VLE_px (self: teqp.teqp.AbstractModel, p_spec: float, xmolar_spec:
                          numpy.ndarray[numpy.float64[m, 1]], T0: float, rhovecL0:
                          numpy.ndarray[numpy.float64[m, 1]], rhovecV0: numpy.ndarray[numpy.float64[m, 1]],
                          options: teqp.teqp.MixVLEpxFlags | None = None) \rightarrow Tuple[teqp.teqp.VLE return code,
                          float, numpy.ndarray[numpy.float64[m, 1]], numpy.ndarray[numpy.float64[m, 1]]]
     pure VLE_T (self: teqp.teqp.AbstractModel, T: float, rhoL: float, rhoV: float, max_iter: int, molefrac:
                     numpy.ndarray[numpy.float64[m, 1]] \mid None = None) \rightarrow numpy.ndarray[numpy.float64[2, 1]]
     solve\_pure\_critical (self: teqp.teqp.AbstractModel, T: float, rho: float, flags: json | None = None) \rightarrow
                                  Tuple[float, float]
     trace_VLE_isobar_binary (self: teqp.teqp.AbstractModel, p: float, T0: float, rhovecL0:
                                       numpy.ndarray[numpy.float64[m, 1]], rhovecV0:
                                       numpy.ndarray[numpy.float64[m, 1]], options: teqp.teqp.PVLEOptions |
                                       None = None \rightarrow json
     trace_VLE_isotherm_binary (self: teqp.teqp.AbstractModel, T: float, rhovecL0:
                                          numpy.ndarray[numpy.float64[m, 1]], rhovecV0:
                                          numpy.ndarray[numpy.float64[m, 1]], options: teqp.teqp.TVLEOptions |
                                          None = None \rightarrow json
     trace_VLLE_binary (self: teqp.teqp.AbstractModel, T: float, rhovecV: numpy.ndarray[numpy.float64[m,
                               1]], rhovecL1: numpy.ndarray[numpy.float64[m, 1]], rhovecL2:
                               numpy.ndarray[numpy.float64[m, 1]], options: teqp.teqp.VLLETracerOptions | None
                               = None \rightarrow json
     trace_critical_arclength_binary (self: teqp.teqp.AbstractModel, T0: float, rhovec0:
                                                   numpy.ndarray[numpy.float64[m, 1]], path: str | None = None,
                                                   options: teqp.teqp.TCABOptions | None = None) \rightarrow json
class teqp.teqp.IterationMatrices
     Bases: pybind11_object
     property J
     property v
     property vars
class teap.teap.MixVLEReturn
     Bases: pybind11_object
     property T
     property initial_r
     property message
     property num_fev
```

8.1. tegp package 105

```
property num_iter
    property r
    property return_code
    property rhovecL
    property rhovecV
    property success
class teqp.teqp.MixVLETpFlags
    Bases: pybind11_object
    property atol
    property axtol
    property maxiter
    property reltol
    property relxtol
class teqp.teqp.MixVLEpxFlags
    Bases: pybind11_object
    property atol
    property axtol
    property maxiter
    property reltol
    property relxtol
class teap.teap.MultiFluidVLEAncillaries
    Bases: pybind11_object
    property pL
    property pV
    property rhoL
    property rhoV
class teap.teap.NRIterator
    Bases: pybind11_object
    get_T (self: teqp.teqp.NRIterator) \rightarrow float
    \texttt{get\_molefrac} (self: teqp.teqp.NRIterator) \rightarrow numpy.ndarray[numpy.float64[m, 1]]
    \texttt{get\_rho} (self: teqp.teqp.NRIterator) \rightarrow float
    get_vals (self: teqp.teqp.NRIterator) → numpy.ndarray[numpy.float64[m, 1]]
```

```
get_vars (self: teqp.teqp.NRIterator) → List[str]
    take\_step (self: teqp.teqp.NRIterator) \rightarrow numpy.ndarray[numpy.float64[m, 1]]
    take_steps (self: teqp.teqp.NRIterator, arg0:int) \rightarrow None
class teqp.teqp.PVLEOptions
    Bases: pybind11_object
    property abs_err
    property calc_criticality
    property crit_termination
    property init_c
    property init_dt
    property integration_order
    property max_dt
    property max_steps
    property polish
    property polish_exception_on_fail
    property polish_reltol_rho
    property rel_err
    property terminate_unstable
    property verbosity
class teqp.teqp.SAFTCoeffs
    Bases: pybind11_object
    property BibTeXKey
    property Qstar2
    property epsilon_over_k
    property m
    property mustar2
    property nQ
    property name
    property nmu
    property sigma_Angstrom
class teqp.teqp.TCABOptions
    Bases: pybind11_object
```

8.1. teqp package 107

```
property T_tol
    property abs_err
    property calc_stability
    property init_c
    property init_dt
    property integration_order
    property max_dt
    property max_step_count
    property polish
    property polish_exception_on_fail
    property polish_reltol_T
    property polish_reltol_rho
    property pure_endpoint_polish
    property rel_err
    property skip_dircheck_count
    property small_T_count
    property stability_rel_drho
    property verbosity
class teqp.teqp.TVLEOptions
    Bases: pybind11_object
    property abs_err
    property calc_criticality
    property crit_termination
    property init_c
    property init_dt
    property integration_order
    property max_dt
    property max_steps
    property p_termination
    property polish
    property polish_exception_on_fail
```

```
property polish_reltol_rho
    property rel_err
    property terminate_unstable
    property verbosity
class teqp.teqp.VLEAncillary
    Bases: pybind11_object
    property T_r
    property Tmax
    property Tmin
class teqp.teqp.VLE_return_code
    Bases: pybind11_object
    Members:
    unset
    xtol satisfied
    functol_satisfied
    maxiter_met
    maxfev_met
    notfinite_step
    functol_satisfied = <VLE_return_code.functol_satisfied: 2>
    maxfev_met = <VLE_return_code.maxfev_met: 3>
    maxiter_met = <VLE_return_code.maxiter_met: 4>
    property name
    notfinite_step = <VLE_return_code.notfinite_step: 5>
    unset = <VLE_return_code.unset: 0>
    property value
    xtol_satisfied = <VLE_return_code.xtol_satisfied: 1>
class teap.teap.VLLEFinderOptions
    Bases: pybind11_object
    property max_steps
    property rho_trivial_threshold
class teqp.teqp.VLLETracerOptions
    Bases: pybind11_object
    property T_limit
```

8.1. teqp package 109

```
property abs_err
     property init_dT
     property max_dT
     property max_polish_steps
     property max_step_count
     property max_step_retries
     property polish
     property rel_err
     property terminate_composition
     property terminate_composition_tol
     property verbosity
teqp.teqp.attach_model_specific_methods(arg0:object) \rightarrow None
teqp.teqp.build_alias_map(root: str) \rightarrow Dict[str, str]
teqp.teqp.build_ancillaries (model: teqp.teqp.AbstractModel, Tc: float, rhoc: float, Tmin: float, flags:
                                    json \mid None = None \mid \rightarrow tegp.tegp.MultiFluidVLEAncillaries
teqp.teqp.collect_component_json(identifiers: List[str], root: str) \rightarrow List[json]
teqp.teqp.convert_CoolProp_idealgas (arg0: str, arg1: int) \rightarrow json
teqp.teqp.convert_FLD (component: str, name: str) \rightarrow json
teqp.teqp.convert_HMXBNC(path: str) → Tuple[json, json]
teqp.teqp.get_BIPdep (BIPcollection: json, identifiers: List[str], flags: json = None) \rightarrow Tuple[json, bool]
\texttt{teqp.teqp.get\_departure\_json} \ (\textit{name: str, root: str}) \ \rightarrow \ \texttt{json}
8.1.3 Module contents
teqp.AmmoniaWaterTillnerRoth()
\texttt{teqp.CPAfactory}\,(\textit{spec})
teqp.IdealHelmholtz(model)
teqp.PCSAFTEOS (coeffs, kmat=None)
teqp.build_LJ126_TholJPCRD2016()
teqp.build_Psi_Hessian_autodiff(model, *args, **kwargs)
teqp.build_Psir_Hessian_autodiff(model, *args, **kwargs)
teqp.build_Psir_gradient_autodiff(model, *args, **kwargs)
```

```
teqp.build_d2PsirdTdrhoi_autodiff(model, *args, **kwargs)
teqp.build_multifluid_ecs_mutant(*args, **kwargs)
teqp.build_multifluid_model (components, coolprop_root, BIPcollectionpath=", flags={}, departurepath=")
teqp.build_multifluid_mutant(*args, **kwargs)
teqp.canonical_PR(Tc_K, pc_Pa, acentric, kmat=None)
teqp.canonical_SRK(Tc_K, pc_Pa, acentric, kmat=None)
tegp.deprecated_caller(model, *args, **kwargs)
teqp.eigen_problem(model, *args, **kwargs)
teqp.extrapolate_from_critical(model, *args, **kwargs)
teqp.find_VLLE_T_binary (model, *args, **kwargs)
teqp.get_B2virget_B12vir(model, *args, **kwargs)
teqp.get_chempotVLE_autodiff(model, *args, **kwargs)
teqp.get_criticality_conditions (model, *args, **kwargs)
teqp.get datapath()
     Get the absolute path to the folder containing the root of multi-fluid data
teqp.get_dchempotdT_autodiff(model, *args, **kwargs)
teqp.get_dpsat_dTsat_isopleth(model, *args, **kwargs)
teqp.get_drhovec_dT_crit (model, *args, **kwargs)
teqp.get_drhovecdT_psat (model, *args, **kwargs)
teqp.get_drhovecdp_Tsat (model, *args, **kwargs)
teqp.get_fugacity_coefficients (model, *args, **kwargs)
teqp.get_minimum_eigenvalue_Psi_Hessian(model, *args, **kwargs)
tegp.get partial molar volumes (model, *args, **kwargs)
tegp.get pr(model, *args, **kwargs)
teqp.get_pure_critical_conditions_Jacobian (model, *args, **kwargs)
teqp.get_splus (model, *args, **kwargs)
teqp.make_model(*args, **kwargs)
     This function is in two parts; first the make model function (renamed to make model in the Python interface) is
     used to make the model and then the model-specific methods are attached to the instance
teqp.make_vdW1 (a, b)
teqp.mix_VLE_Tx (model, *args, **kwargs)
teqp.mix_VLLE_T (model, *args, **kwargs)
```

8.1. tegp package 111

```
teqp.mixture_VLE_px (model, *args, **kwargs)

teqp.pure_VLE_T (model, *args, **kwargs)

teqp.solve_pure_critical (model, *args, **kwargs)

teqp.tolist(a)

teqp.trace_VLE_isobar_binary (model, *args, **kwargs)

teqp.trace_VLE_isotherm_binary (model, *args, **kwargs)

teqp.trace_critical_arclength_binary (model, *args, **kwargs)

teqp.trace_critical_arclength_binary (model, *args, **kwargs)

teqp.vdWEOS(Tc_K, pc_Pa)

teqp.vdWEOS1(*args)
```

CHAPTER

NINE

INDICES AND TABLES

- genindex
- modindex
- search

PYTHON MODULE INDEX

```
teqp, 110 teqp. teqp, 101
```

116 Python Module Index

INDEX

A	С
abs_err (teqp.teqp.PVLEOptions property), 107 abs_err (teqp.teqp.TCABOptions property), 108	<pre>calc_criticality(teqp.teqp.PVLEOptions property), 107</pre>
abs_err (teap.teap.TVLEOptions property), 108	calc_criticality (teqp.teqp.TVLEOptions prop-
abs_err (teqp.teqp.VLLETracerOptions property), 109	erty), 108
AbstractModel (class in teap.teap), 101	calc_stability (teqp.teqp.TCABOptions property),
AmmoniaWaterTillnerRoth() (in module teqp),	108
110	canonical_PR() (in module teqp), 111
atol (teqp.teqp.MixVLEpxFlags property), 106	canonical_SRK() (in module teqp), 111
atol (teap.teap.MixVLETpFlags property), 106	<pre>collect_component_json() (in module teqp.teqp),</pre>
attach_model_specific_methods()(in module	110
teqp.teqp), 110	convert_CoolProp_idealgas() (in module
axtol (teqp.teqp.MixVLEpxFlags property), 106 axtol (teqp.teqp.MixVLETpFlags property), 106	teqp.teqp), 110 convert_FLD() (in module teqp.teqp), 110
axcor (teap.teap.tmx v LETpT tags property), 100	convert_HMXBNC() (in module teqp.teqp), 110
В	CPAfactory() (in module teqp), 110
BibTeXKey (teap.teap.SAFTCoeffs property), 107	crit_termination(teqp.teqp.PVLEOptions property),
build_alias_map() (in module teap.teap), 110	107
build_ancillaries() (in module teap.teap), 110	crit_termination (teqp.teqp.TVLEOptions prop-
build_d2PsirdTdrhoi_autodiff() (in module	erty), 108
teqp), 110	D
<pre>build_d2PsirdTdrhoi_autodiff()</pre>	D
(teqp.teqp.AbstractModel method), 101	deprecated_caller() (in module teqp), 111
build_LJ126_TholJPCRD2016() (in module teqp),	<pre>dpsatdT_pure() (teqp.teqp.AbstractModel method),</pre>
110	101
<pre>build_multifluid_ecs_mutant() (in module</pre>	E
build_multifluid_model() (in module teap), 111	eigen_problem() (in module teqp), 111
<pre>build_multifluid_mutant() (in module teqp),</pre>	eigen_problem() (teqp.teqp.AbstractModel method), 101
<pre>build_Psi_Hessian_autodiff() (in module</pre>	epsilon_over_k (teqp.teqp.SAFTCoeffs property), 107
teqp), 110	<pre>extrapolate_from_critical() (in module teqp),</pre>
build_Psi_Hessian_autodiff()	111
(teqp.teqp.AbstractModel method), 101 build_Psir_gradient_autodiff() (in module	<pre>extrapolate_from_critical() (teap.teap.AbstractModel method), 101</pre>
teqp), 110	(teqp.teqp.Abstractimoaet methoa), 101
build_Psir_gradient_autodiff()	F
(teqp.teqp.AbstractModel method), 101	find_VLLE_p_binary() (teqp.teqp.AbstractModel
build_Psir_Hessian_autodiff() (in module	method), 101
teqp), 110	find_VLLE_T_binary() (in module teqp), 111
<pre>build_Psir_Hessian_autodiff()</pre>	find_VLLE_T_binary() (teqp.teqp.AbstractModel
(teqp.teqp.AbstractModel method), 101	method), 101

functol_satisfied (teqp.teqp.VLE_return_code at- tribute), 109	<pre>get_dchempotdT_autodiff() (teap.teap.AbstractModel method), 103</pre>
<i>"</i>	get_departure_json() (in module teqp.teqp), 110
G	<pre>get_deriv_mat2() (teqp.teqp.AbstractModel</pre>
get_Ar00() (teqp.teqp.AbstractModel method), 102	method), 103
get_Ar00n() (teqp.teqp.AbstractModel method), 102	<pre>get_dmBnvirdTm() (teqp.teqp.AbstractModel</pre>
-	method), 104
get_Ar01() (teap.teap.AbstractModel method), 102	get_dp_dT_crit() (teqp.teqp.AbstractModel
get_Ar01n() (teqp.teqp.AbstractModel method), 102	method), 104
get_Ar02() (teqp.teqp.AbstractModel method), 102	get_dpsat_dTsat_isopleth() (in module teqp),
get_Ar02n() (teqp.teqp.AbstractModel method), 102	111
get_Ar03() (teap.teap.AbstractModel method), 102	get_dpsat_dTsat_isopleth()
get_Ar03n() (teqp.teqp.AbstractModel method), 102	(teap.teap.AbstractModel method), 104
get_Ar04() (teqp.teqp.AbstractModel method), 102	
get_Ar04n() (teqp.teqp.AbstractModel method), 102	get_drhovec_dT_crit() (in module teqp), 111
get_Ar05n() (teqp.teqp.AbstractModel method), 102	<pre>get_drhovec_dT_crit() (teqp.teqp.AbstractModel</pre>
get_Ar06n() (teqp.teqp.AbstractModel method), 102	method), 104
get_Ar10() (teqp.teqp.AbstractModel method), 102	get_drhovecdp_Tsat() (in module teqp), 111
get_Ar11() (teqp.teqp.AbstractModel method), 102	<pre>get_drhovecdp_Tsat() (teqp.teqp.AbstractModel</pre>
get_Ar12() (teqp.teqp.AbstractModel method), 102	method), 104
get_Ar13() (teqp.teqp.AbstractModel method), 102	<pre>get_drhovecdT_psat() (in module teqp), 111</pre>
get_Ar14() (teap.teap.AbstractModel method), 103	<pre>get_drhovecdT_psat() (teqp.teqp.AbstractModel</pre>
get_Ar20() (teap.teap.AbstractModel method), 103	method), 104
get_Ar21() (teqp.teqp.AbstractModel method), 103	<pre>get_fugacity_coefficients() (in module teqp),</pre>
get_Ar22() (teqp.teqp.AbstractModel method), 103	111
get_Ar23() (teqp.teqp.AbstractModel method), 103	<pre>get_fugacity_coefficients()</pre>
get_Ar24() (teqp.teqp.AbstractModel method), 103	(teqp.teqp.AbstractModel method), 104
get_Arxy() (teqp.teqp.AbstractModel method), 103	<pre>get_minimum_eigenvalue_Psi_Hessian() (in</pre>
get_AtaudeltaXi() (teqp.teqp.AbstractModel	module teqp), 111
method), 103	<pre>get_minimum_eigenvalue_Psi_Hessian()</pre>
get_AtaudeltaXiXj() (teqp.teqp.AbstractModel	(teqp.teqp.AbstractModel method), 104
method), 103	<pre>get_molefrac() (teap.teap.NRIterator method), 106</pre>
get_AtaudeltaXiXjXk() (teqp.teqp.AbstractModel	get_neff() (teqp.teqp.AbstractModel method), 104
method), 103	<pre>get_partial_molar_volumes() (in module teqp),</pre>
get_ATrhoXi() (teqp.teqp.AbstractModel method), 101	111
	<pre>get_partial_molar_volumes()</pre>
get_ATrhoXiXj() (teqp.teqp.AbstractModel method),	(teap.teap.AbstractModel method), 104
102	get_pr() (in module teqp), 111
get_ATrhoXiXjXk() (teqp.teqp.AbstractModel	get_pr() (teap.teap.AbstractModel method), 104
method), 102	get_pir() (keyp.keyp.nosintcimoter method), 104 get_pure_critical_conditions_Jacobian()
get_B2vir() (teqp.teqp.AbstractModel method), 103	(in module teap), 111
get_B2virget_B12vir() (in module teqp), 111	
get_B12vir() (teqp.teqp.AbstractModel method), 103	get_pure_critical_conditions_Jacobian()
get_BIPdep() (in module teqp.teqp), 110	(teqp.teqp.AbstractModel method), 104
get_Bnvir() (teqp.teqp.AbstractModel method), 103	get_R() (teap.teap.AbstractModel method), 103
<pre>get_chempotVLE_autodiff() (in module teqp),</pre>	get_rho() (teap.teap.NRIterator method), 106
111	<pre>get_splus() (in module teqp), 111</pre>
get_chempotVLE_autodiff()	get_splus() (teqp.teqp.AbstractModel method), 104
(teqp.teqp.AbstractModel method), 103	get_T() (teqp.teqp.NRIterator method), 106
<pre>get_criticality_conditions() (in module</pre>	get_vals() (teqp.teqp.NRIterator method), 106
teqp), 111	get_vars() (teqp.teqp.NRIterator method), 106
get_criticality_conditions()	1
(tegp.tegp.AbstractModel method), 103	1
get_datapath() (in module teqp), 111	IdealHelmholtz() (in module teap), 110
<pre>get_dchempotdT_autodiff() (in module teqp),</pre>	init_c (teap.teap.PVLEOptions property), 107
111	init_c (teap.teap.TCABOptions property), 108
	= \ n n 1 1 2//

118 Index

<pre>init_c (teqp.teqp.TVLEOptions property), 108 init_dt (teqp.teqp.PVLEOptions property), 107 init_dt (teqp.teqp.TCABOptions property), 108</pre>	module teqp, 110 teqp.teqp, 101
<pre>init_dt (teqp.teqp.TVLEOptions property), 108</pre>	MultiFluidVLEAncillaries (class in teqp.teqp),
<pre>init_dT (teap.teap.VLLETracerOptions property), 110</pre>	106
<pre>initial_r (teqp.teqp.MixVLEReturn property), 105</pre>	mustar2 (teqp.teqp.SAFTCoeffs property), 107
integration_order (teap.teap.PVLEOptions prop-	N
erty), 107	
integration_order (teap.teap.TCABOptions prop-	name (teqp.teqp.SAFTCoeffs property), 107
erty), 108	name (teqp.teqp.VLE_return_code property), 109
<pre>integration_order (teap.teap.TVLEOptions prop- erty), 108</pre>	nmu (teqp.teqp.SAFTCoeffs property), 107
IterationMatrices (class in teap.teap), 105	notfinite_step (teqp.teqp.VLE_return_code at- tribute), 109
	nQ (teqp.teqp.SAFTCoeffs property), 107
J	NRIterator (class in teqp.teqp), 106
J (teap.teap.IterationMatrices property), 105	num_fev (teap.teap.MixVLEReturn property), 105
o (teapheaphieranomian tees property), 100	num_iter (teqp.teqp.MixVLEReturn property), 105
M	
m (teap.teap.SAFTCoeffs property), 107	P
make_model() (in module teap), 111	p_termination(teap.teap.TVLEOptions property), 108
make_vdW1() (in module teqp), 111	PCSAFTEOS () (in module teqp), 110
max_dt (teqp.teqp.PVLEOptions property), 107	pL (teap.teap.MultiFluidVLEAncillaries property), 106
max_dt (teqp.teqp.TCABOptions property), 108	polish (teqp.teqp.PVLEOptions property), 107
max_dt (teap.teap.TVLEOptions property), 108	polish (teqp.teqp.TCABOptions property), 108
max_dT (teqp.teqp.VLLETracerOptions property), 110	polish (teap.teap.TVLEOptions property), 108
<pre>max_polish_steps (teqp.teqp.VLLETracerOptions</pre>	polish (teqp.teqp.VLLETracerOptions property), 110
property), 110	polish_exception_on_fail
<pre>max_step_count (teqp.teqp.TCABOptions property),</pre>	(teqp.teqp.PVLEOptions property), 107
108	polish_exception_on_fail
<pre>max_step_count (teqp.teqp.VLLETracerOptions prop-</pre>	(teqp.teqp.TCABOptions property), 108
erty), 110	polish_exception_on_fail
<pre>max_step_retries (teqp.teqp.VLLETracerOptions</pre>	(teqp.teqp.TVLEOptions property), 108
property), 110	polish_reltol_rho (teap.teap.PVLEOptions prop-
max_steps (teap.teap.PVLEOptions property), 107	erty), 107
<pre>max_steps (teap.teap.TVLEOptions property), 108 max_steps (teap.teap.VLLEFinderOptions property),</pre>	polish_reltol_rho (teqp.teqp.TCABOptions property), 108
109	polish_reltol_rho (teap.teap.TVLEOptions prop-
maxfev_met (teap.teap.VLE_return_code attribute), 109	erty), 108
maxiter (teqp.teqp.MixVLEpxFlags property), 106	polish_reltol_T (teap.teap.TCABOptions property),
maxiter (teqp.teqp.MixVLETpFlags property), 106	108
maxiter_met (teap.teap.VLE_return_code attribute),	<pre>pure_endpoint_polish (teqp.teqp.TCABOptions</pre>
109	property), 108
message (teap.teap.MixVLEReturn property), 105	pure_VLE_T() (in module teqp), 112
<pre>mix_VLE_Tp() (teqp.teqp.AbstractModel method), 104</pre>	<pre>pure_VLE_T() (teqp.teqp.AbstractModel method), 105</pre>
mix_VLE_Tx() (in module teqp), 111	pV (teqp.teqp.MultiFluidVLEAncillaries property), 106
<pre>mix_VLE_Tx() (teqp.teqp.AbstractModel method), 104</pre>	PVLEOptions (class in teqp.teqp), 107
mix_VLLE_T() (in module teqp), 111	0
mix_VLLE_T() (teqp.teqp.AbstractModel method), 105	Q
mixture_VLE_px() (in module teqp), 111	Qstar2 (teqp.teqp.SAFTCoeffs property), 107
mixture_VLE_px() (teqp.teqp.AbstractModel	
method), 105	R
MixVLEpxFlags (class in teap.teap), 106	r (teqp.teqp.MixVLEReturn property), 106
MixVLEReturn (class in teap.teap), 105	rel_err (teap.teap.PVLEOptions property), 107
MixVLETpFlags (class in teqp.teqp), 106	

Index 119

rel_err (teqp.teqp.TCABOptions property), 108	<pre>trace_critical_arclength_binary()</pre>
rel_err (teqp.teqp.TVLEOptions property), 109	(teqp.teqp.AbstractModel method), 105
rel_err (teap.teap.VLLETracerOptions property), 110	<pre>trace_VLE_isobar_binary() (in module teqp),</pre>
reltol (teqp.teqp.MixVLEpxFlags property), 106	112
reltol (teap.teap.MixVLETpFlags property), 106	trace_VLE_isobar_binary()
relxtol (teqp.teqp.MixVLEpxFlags property), 106	(teqp.teqp.AbstractModel method), 105
relxtol (teqp.teqp.MixVLETpFlags property), 106	<pre>trace_VLE_isotherm_binary() (in module teqp),</pre>
return_code (teqp.teqp.MixVLEReturn property), 106	112
rho_trivial_threshold	trace_VLE_isotherm_binary()
(teqp.teqp.VLLEFinderOptions property), 109	(teqp.teqp.AbstractModel method), 105
rhoL (teqp.teqp.MultiFluidVLEAncillaries property), 106	trace_VLLE_binary() (teqp.teqp.AbstractModel
rhoV (teqp.teqp.MultiFluidVLEAncillaries property), 106	method), 105
rhovecL (teap.teap.MixVLEReturn property), 106	TVLEOptions (class in teap.teap), 108
rhovecV (teqp.teqp.MixVLEReturn property), 106	U
S	
	unset (teqp.teqp.VLE_return_code attribute), 109
SAFTCoeffs (class in teqp.teqp), 107	V
sigma_Angstrom (teap.teap.SAFTCoeffs property), 107	
skip_dircheck_count (teqp.teqp.TCABOptions	∨ (teqp.teqp.IterationMatrices property), 105
<pre>property), 108 small_T_count (teqp.teqp.TCABOptions property), 108</pre>	value (teqp.teqp.VLE_return_code property), 109
solve_pure_critical() (in module teap), 112	vars (teap.teap.IterationMatrices property), 105
solve_pure_critical() (teqp.teqp.AbstractModel	vdWEOS () (in module teqp), 112
method), 105	vdWEOS1 () (in module teqp), 112
stability_rel_drho (teqp.teqp.TCABOptions prop-	verbosity (teap.teap.PVLEOptions property), 107
erty), 108	verbosity (teap.teap.TCABOptions property), 108
success (teap.teap.MixVLEReturn property), 106	verbosity (teap.teap.TVLEOptions property), 109
Success (reqp.reqp.min vibilitation property), 100	verbosity (teap.teap.VLLETracerOptions property),
Т	
	VLE_return_code (class in teap.teap), 109
T (teqp.teqp.MixVLEReturn property), 105 T_limit (teqp.teqp.VLLETracerOptions property), 109	VLEAncillary (class in teqp.teqp), 109 VLLEFinderOptions (class in teqp.teqp), 109
T_r (teqp.teqp.VLEAncillary property), 109	VLLETracerOptions (class in teap.teap), 109
T_tol (teap.teap.TCABOptions property), 107	VIDETTACETOPCTOMS (class in teap.teap), 10)
take_step() (teap.teap.NRIterator method), 107	X
take_steps() (teqp.teqp.NRIterator method), 107	
TCABOptions (class in teap.teap), 107	xtol_satisfied (teqp.teqp.VLE_return_code at-
teqp	tribute), 109
module, 110	
teqp.teqp	
module, 101	
terminate_composition	
(teqp.teqp.VLLETracerOptions property), 110	
terminate_composition_tol	
(teap.teap.VLLETracerOptions property), 110	
terminate_unstable (teap.teap.PVLEOptions prop-	
erty), 107	
terminate_unstable (teqp.teqp.TVLEOptions prop-	
erty), 109	
Tmax (teqp.teqp.VLEAncillary property), 109	
Tmin (teqp.teqp.VLEAncillary property), 109	
tolist() (in module teqp), 112	
trace_critical_arclength_binary() (in	
module teqp), 112	

120 Index