# Module fermi\_eos

# **Functions**

```
def beta_equilibrium_condition(k_p, k_n)
```

Condition of beta equilibrium in order to compute  $k_p$  as a function of  $k_n$ . If it returns zero, then the Fermi momentum of the proton is the expected one, since it means that chemical potentials of neutrons, protons and electrons match the equilibrium condition for the (beta) weak interactions.

#### **Parameters**

k p : float

Fermi momentum of protons in g cm / s.

k n : float

Fermi momentum of neutrons in g cm / s.

#### Returns

float

Chemical potential in erg.

```
def compute_energy_density_emp(p)
```

Compute energy density from pressure.

# **Parameters**

p : float

Pressure in dyne/cm^2.

## Raises

ValueError

If energy density is negative.

#### Returns

float

Energy density in erg/cm<sup>3</sup>.

```
def compute_energy_density_from_u_emp(u)
```

Compute energy density from  $u = n / n_0$ .

#### **Parameters**

```
\mathbf{u}: float Dimensionless \mathbf{u} = \mathbf{n} / \mathbf{n}_{-}\mathbf{0}.
```

## Returns

float

Energy density in erg/cm<sup>3</sup>.

```
def compute_energy_density_from_u_sky(u)
```

Compute energy density from  $u = n / n_0$ .

#### **Parameters**

```
\mathbf{u}: float Dimensionless \mathbf{u} = \mathbf{n} / \mathbf{n}_{-} \mathbf{0}.
```

## Returns

float

Energy density in erg/cm<sup>3</sup>.

```
def compute_energy_density_from_x_gen(x)
```

Compute energy density from x = k / (m c).

## **Parameters**

```
\mathbf{x}: float
Dimensionless \mathbf{x} = \mathbf{k} / (\mathbf{m} \ \mathbf{c}).
```

# Raises

ValueError

If energy density is negative.

## Returns

float

Energy density in erg/cm<sup>3</sup>.

```
{\tt def} \ \ \textbf{compute\_energy\_density\_from\_x\_npe} \ (x \text{, e\_0})
```

Compute energy density from x = k / (mc) and prefactor  $e_0$ .

#### **Parameters**

x : float

Dimensionless x.

e 0 : float

Prefactor in erg/cm<sup>3</sup>.

## Returns

float

Energy density in erg/cm<sup>3</sup>.

```
def compute_energy_density_gen(p)
```

Compute energy density from pressure.

Parameters: - p : float Pressure in dyne/cm^2.

Returns: - float Energy density in erg/cm^3.

```
def compute_energy_density_npe(k n)
```

In presence of protons, electrons and neutrons, compute energy density from Fermi momentum of neutrons.

## **Parameters**

```
k n : float
```

Fermi momentum of neutrons in g cm / s.

## Raises

ValueError

If energy density is negative.

Returns:

float

Energy density in erg/cm<sup>3</sup>.

```
def compute_energy_density_pe(k_p)
```

In presence of protons and electrons (no neutrons), compute energy density from Fermi momentum of protons.

# **Parameters**

```
k n : float
```

Fermi momentum of protons in g cm / s.

#### Raises

```
ValueError
```

If energy density is negative.

## Returns

float

Energy density in erg/cm<sup>3</sup>.

```
def compute_energy_density_sky(p)
```

Compute energy density from pressure.

## **Parameters**

p : float

Pressure in dyne/cm^2.

#### Raises:

ValueError If energy density is negative.

#### Returns

float

Number density in erg/cm<sup>3</sup>.

```
def compute_fermi_momentum_nonrel(p)
```

Compute Fermi momentum from pressure in the non-relativistic limit.

# **Parameters**

**p** : float

Pressure in dyne/cm^2.

#### Returns

float

Fermi momentum in g cm/s.

```
def compute_k_n(n)
```

Compute Fermi momentum of neutrons from number density.

# **Parameters**

n : float

Number density in 1/cm<sup>3</sup>

## Returns

float

Fermi momentum of neutrons in g cm / s.

```
def compute_k_p(k_n)
```

Compute k\_p from k\_n. he fsolve algorithm is used to find the right value of k\_p that matches the beta equilibrium condition for chemical potential.

#### **Parameters**

k\_n : float

Fermi momentum of neutrons in g cm / s.

#### Raises

ValueError

If Fermi momentum of protons is negative.

#### Returns

float

Fermi momentum of protons in g cm / s.

```
def compute number density emp(p)
```

Compute number density from pressure.

## **Parameters**

**p** : float

Pressure in dyne/cm^2.

## Raises

ValueError

If number density is negative.

# Returns

float

Number density in erg/cm<sup>3</sup>.

```
def compute_number_density_gen(p)
```

Compute number density from pressure

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Parameters: - p : float Pressure in dyne/cm^2.

Returns: - float Number density in erg/cm^3.

```
def compute_number_density_nonrel(p)
```

Compute number density from pressure in the non-relativistic limit.

## **Parameters**

**p** : float

Pressure in dyne/cm^2.

#### Returns

float

Number density in 1/cm<sup>3</sup>.

```
def compute_number_density_sky(p)
```

Compute number density from pressure.

# **Parameters**

**p** : float

Pressure in dyne/cm^2.

## Raises

ValueError

If number density is negative.

## Returns:

float Energy density in erg/cm<sup>3</sup>.

```
def compute_pressure_from_u_emp(u, p)
```

Compute calculated pressure minus expected pressure from the latter and  $u = n / n_0$ . If it returns zero, then the pressure is exactly the desidered one.

#### **Parameters**

 $\mathbf{u}$  : float

Dimensionless  $u = n / n_0$ .

 ${\bf p}$  : float

Pressure in dyne/cm^2.

## Returns

float

Calculated pressure minus expected.

```
def compute_pressure_from_u_sky(u, p)
```

Compute calculated pressure minus expected pressure from expected pressure and  $u = n / n_0$ . If it returns zero, then the pressure is exactly the desidered one.

#### **Parameters**

u : float

Dimensionless  $u = n / n_0$ .

p : float

Pressure in dyne/cm^2.

## Returns

float

Calculated pressure minus expected.

```
def compute_pressure_from_x_npe(x, e_0)
```

Compute pressure from x = k / (mc) and prefactor  $e_0$ .

## **Parameters**

 $\mathbf{x}$  : float

Dimensionless x.

e 0 : float

Prefactor in erg/cm<sup>3</sup>.

#### Returns

float

Pressure in dyne/cm^2.

```
def compute_pressure_gen(x, p)
```

Compute calculated pressure minus expected pressure given the latter and x = k / (m c). If it returns zero, then the pressure is exactly the desidered one.

## **Parameters**

x : float

Dimensionless x = k / (m c).

**p** : float

Pressure in dyne/cm^2.

## Returns

float

Calculated pressure minus expected.

```
def compute_pressure_npe(k_n)
```

In presence of protons, electrons and neutrons, compute pressure from Fermi momentum of neutrons.

## **Parameters**

```
k n : float
```

Fermi momentum of neutrons in g cm / s.

#### Raises

ValueError

If pressure is negative.

## Returns

float

Pressure in dyne/cm^2.

```
def compute_pressure_pe(k_p)
```

In presence of protons and electrons (no neutrons), compute pressure from Fermi momentum of protons.

## **Parameters**

 $\mathbf{k}_{\mathbf{p}}$ : float

Fermi momentum of protons in g cm / s.

## Raises

ValueError

If pressure is negative.

## Returns

float

Pressure in dyne/cm^2.

```
def compute_u_emp(p)
```

Compute  $u = n / n_0$  from pressure. The toms748 algorithm is used to find the right value of u that matches the expected pressure with the calculated one.

#### **Parameters**

```
p : float
```

Pressure in dyne/cm^2.

## Raises

```
ValueError
```

If u is negative.

#### Returns

```
float
```

Dimensionless  $u = n / n_0$ .

```
def compute_u_sky(p)
```

Compute  $u = n / n_0$  from pressure. The toms748 algorithm is used to find the right value of u that matches the expected pressure with the calculated one.

## **Parameters**

```
u : float
```

Dimensionless  $u = n / n_0$ .

# Raises

ValueError

If u is negative.

# Returns

float

Energy density in erg/cm<sup>3</sup>.

```
def compute_x_gen(p)
```

Compute x = k / (m c) from pressure. The toms748 algorithm is used to find the right value of x that matches the expected pressure with the calculated one.

#### **Parameters**

p : float

Pressure in dyne/cm^3.

## Raises

ValueError

If x is negative.

#### Returns

float

Dimensionless x = k / (m c).

```
def equation_of_state_emp(pressures)
```

Compute and save to file the equation of state for empirical interactions. The equation of state is in the form energy density as function of pressure. Also number densities are computed and saved to file.

#### **Parameters**

pressures : NumPy array

Numpy array of pressures in dyne/cm^2.

Raises: - ValueError If pressure is negative.

```
def equation_of_state_gen(pressures)
```

Compute and save to file the equation of state for a generic degenerate ideal Fermi gas of neutrons. The equation of state is in the form energy density as function of pressure. Also number densities are computed and saved to file.

#### **Parameters**

pressures : NumPy array

Numpy array of pressures in dyne/cm^2.

Raises: - ValueError If pressure is negative.

```
def equation_of_state_nonrel(pressures)
```

Compute and save to file the equation of state for a non-relativistic degenerate ideal Fermi gas of neutrons. The equation of state is in the form energy density as function of pressure. Also number densities are computed and saved to file.

## **Parameters**

pressures : NumPy array

Numpy array of pressures in dyne/cm^2.

Raises: - ValueError If pressure is negative.

```
def equation of state npe (number densities)
```

Compute and save to file the equation of state for a generic degenerate ideal Fermi gas of neutrons, protons and electrons in beta equilibrium. The equation of state is in the form energy density as function of pressure.

## **Parameters**

number densities : NumPy array

Numpy array of number densities in 1/cm<sup>3</sup>.

Raises: - ValueError If number density is negative.

```
def equation_of_state_sky(pressures)
```

Compute and save to file the equation of state for Skyrme Hartee-Fock interactions. The equation of state is in the form energy density as function of pressure. Also number densities are computed and saved to file.

#### **Parameters**

pressures : NumPy array

Numpy array of pressures in dyne/cm^2.

Raises: - ValueError If pressure is negative.

```
def polytropic_equation_of_state(p, k,
gamma)
```

Compute energy density from pressure using polytropic equation of state. K is the prefactor and gamma is the exponent of energy density.

## **Parameters**

p: float

Pressure in dyne/cm^2.

k : float

K in (dyne/cm<sup>2</sup>)<sup>(1 - gamma)</sup>.

gamma : float

Dimensionless Gamma.

#### Returns

float

Energy density in erg/cm<sup>3</sup>.

#### **Functions**

```
beta_equilibrium_condition
compute_energy_density_emp
compute_energy_density_from_u_emp
compute_energy_density_from_u_sky
compute_energy_density_from_x_gen
compute_energy_density_from_x_npe
compute_energy_density_gen
compute_energy_density_npe
compute_energy_density_pe
compute_energy_density_sky
compute_fermi_momentum_nonrel
compute_k n
```

- compute k p
- compute number density emp
- compute number density gen
- compute number density nonrel
- compute\_number\_density\_sky
- compute pressure from u emp
- compute pressure from u sky
- compute\_pressure\_from\_x\_npe
- compute pressure gen
- compute pressure npe
- compute pressure pe
- compute u emp
- compute\_u\_sky
- compute\_x\_gen
- equation of state emp
- equation of state gen
- equation of state nonrel
- equation of state npe
- equation of state sky
- $\bullet \ \, \texttt{polytropic\_equation\_of\_state} \\$

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