³He library

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Usage

This library provides constant and functions for various ³He properties. Supported Languages:

- Fortran-77. You should use he3.fh include file and libhe3 library for fortran programs. All functions and arguments are real*8. Use -fno-range-check compiler flag to allow NaN values.
- Fortran-90. You should include he3.f90h instead.
- C. You should use he3.h include file and libhe3 library in C programs. Lowercase names with underscore should be used (like he3_pf_). Type of arguments is always double*, type of returned value is double.
- Matlab, Octave. mex files are located in the matlab folder. Lowercase names should be used. In function arguments you can mix numbers with arrays or matrices of the same size. Example:

```
ttc=0:0.1:1; p=10;
I = he3_nu_b(p, ttc);
```

To bild mex files you can use make octave, make matlab or make matlab64 in the matlab folder. Fixed rpath is used for the library. For building this documentation octave files are needed.

• Command line. You can use all the functions via a command line interface. To get a list of all functions run the program he3 without arguments:

> he3

To get information about specific function run the program with the function name as an argument:

```
> he3 he3_nu_b
```

You can use matlab-style vectors (value1:step:value2) to get a table of values (if step is missing then 20 points will be printed):

```
> he3 he3_nu_b 0.1:0.1:1 0
```

will give you Leggett frequency vs temperature at zero pressure.

Location

```
GIT https://github.com/slazav/he3lib
```

```
ROTA /home/slazav/he3lib/lib
/rota/programs/src/he3lib - old, "stable" version.
Use "addpath /rota/programs/src/he3lib/matlab" in Matlab.
```

General

```
Avogadro constant N_A = 6.02214129 \cdot 10^{23} [1/\text{mol}]
const_na
                Boltsman constant k_B=1.3806488\cdot 10^{-16}~[\mathrm{erg/K}]
const\_kb
                R-gas constant R = 8.314472 \cdot 10^7 \text{ [sgs]}
const_r
                Plank constant h = 6.62606957 \cdot 10^{27} \text{ [sgs]}
const_h
const_hbar \hbar = h/2\pi = 1.054571726 \cdot 10^{27} \; [\mathrm{sgs}]
const_pi
                \pi = 3.1415926535897932
                2\pi
const_2pi
                <sup>3</sup>He atom mass, m_3 = 5.0079 \cdot 10^{-24} [g]
he3_amass
                <sup>3</sup>He molar mass, \mu = N_A m_3 = 3.0158 [g/mol]
he3\_mmass
                <sup>3</sup>He gyromagnetic ratio \gamma = 20378 [(G s)<sup>-1</sup>],
he3_gyro
```

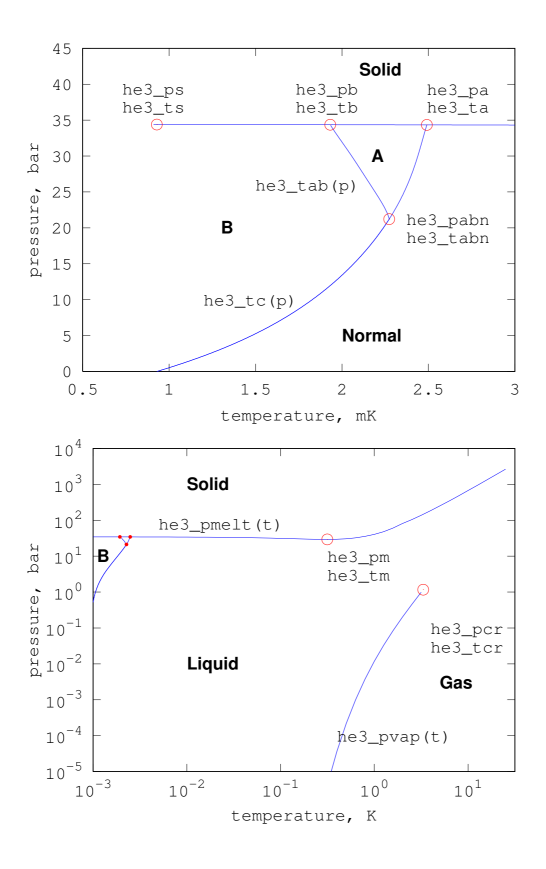
Phase diagram

Vapor pressure and critical point are from the 1962 3 He scale of temperatures. Melting curve $P_m(T)$ at 0.9-250 mK, $T_c(P)$, $T_{AB}(P)$ and all tricritical points are from Greywall-86 temperature scale. Melting curve minimum is from PLTS-2000 temperature scale. Melting curve for higher temperatures is from Osborne, Abraham, Weinstock-1951 and Mills, Grilly-1955 with interpolation in the regions 0.25-0.5K and 1.5-2.0K. PLTS-2000 melting curve and critical points on it are also available. See more information in the source file he3_phase.f.

he3_pvap(T)	Vapor pressure [bar] vs temperature [K], $T=0.2-3.324~\mathrm{K}$
he3_pcr	Gas-liquid critical point pressure, 1.16317 [bar]
he3_tcr	Gas-liquid critical point temperature, $3.324~[\mathrm{K}]$
he3_pmelt(T)	Melting pressure [bar] vs temperature [K], $T = 0.0009 - 31$ K
he3_pm	Melting curve minimum pressure, 29.3113 [bar]
he3_tm	Melting curve minimum temperature, 0.31524 [K]
he3_pa	Superfluid trans. at melting curve, pressure, 34.3380 [bar]
he3_ta	Superfluid trans. at melting curve, temp., $2.491~[\mathrm{mK}]$
he3_pb	A-B trans. at melting curve, pressure, 34.3580 [bar]
he3_tb	A-B trans. at melting curve, temp., 1.932 $[mK]$
he3_ps	Neel transition at melting curve, pressure, 34.3905 [bar]
he3_ts	Neel transition at melting curve, temp., $0.9291~[\mathrm{mK}]$
he3_tc(P)	Superfluid transition temperature [mK] vs pressure [bar], $P=0-34.358~\mathrm{bar}$
he3_tab(P)	A-B transition temperature [mK] vs pressure [bar], $P=0-34.3609~{\rm bar,~below~21.22~bar~is~equal~to~he3_tc}$
he3_tabn	A-B-Normal point temperature, 2.2311 $[mK]$
he3_pabn	A-B-Normal point pressure, 21.22 [bar]
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PLTS-2000 temperature scale:

he3_pmelt_plts(T)	PLTS-2000 melting presure [bar] vs temperature [K].								
he3_pa_plts	Superfluid trans. at melting curve, pressure, 34.3407 [bar]								
he3_ta_plts	Superfluid trans. at melting curve, temp., $2.444~[\mathrm{mK}]$								
he3_pb_plts	A-B trans. at melting curve, pressure, 34.3609 [bar]								
he3_tb_plts	A-B trans. at melting curve, temp., 1.896 [mK]								
he3_ps_plts	Neel transition at melting curve, pressure, $34.3934~[\mathrm{bar}]$								
he3_ts_plts	Neel transition at melting curve, temp., 0.902 [mK]								



Fermi-liquid parameters

Fermi-liquid parameters according to Wheatley-75 table. Values are calculated using experimental data for molar volume, speed of sound, magnetic temperature T^* , collected by Wheatley-75 and heat capacity $(\gamma_f = C/RT)$ measured by Graywall-86. Argument is pressure, [bar] in the range 0 – 34.4. See more information in the source file he3_fermi.f.

Measured values:

he3_vm(P) Molar volume
$$v_m$$
, [cm³/mol]

he3_c1(P) First sound velocity,
$$c_1$$
, [cm/s]

he3_gammaf(P) R-gas constant
$$\gamma_f = C/RT$$
, $[1/(\text{K mol})]$

he3_tmag(P) Magnetic temperature
$$T^*$$
, [K]

Derived values:

he3_rho(P) Density
$$\rho = \mu/v_m$$
, [g/cm³]

he3_2n0(P)
$$2N(0) = \frac{\gamma_f}{v_m} \frac{3N_A}{k_B \pi^2}, [1/(\text{erg cm}^3)]$$

$$\mbox{he3_pf(P)} \qquad p_F = h \left(\frac{3}{8\pi} \; \frac{N_A}{v_m} \right)^{1/3}, \, [\mbox{g cm/s}] \label{eq:pf_P}$$

$$\label{eq:mass_mass_mass_mass_mass} \text{he3.meff(P)} \quad m^\star = \frac{h^3}{8\pi} \; \frac{2N(0)}{p_F}, \, [\text{g}]$$

he3_mm(P)
$$m^*/m_3$$

he3_vf(P)
$$v_F = p_F/m^\star, \, [\mathrm{cm/s}]$$

$$\label{eq:chin} \mbox{he3_chin(P)} \ \chi_N = 2N(0) \frac{(\gamma \hbar)^2}{4(1+F_0^a)}.$$

he3_f0s(P)
$$F_0^s = 3 \ m^{\star} m_3 \ c_1^2/p_F^2 - 1$$

he3_f1s(P)
$$F_1^s = 3(m^*/m_3 - 1)$$

he3_f0a(P)
$$F_0^a = Z_0/4 = 3k_B T^{\star} m^{\star}/p_F^2 - 1$$

he3_f1a(P)
$$F_1^c$$

he3_a(P) Average atomic spacing,
$$a = (v_m/N_A)^{1/3}$$
, Å

he3_gdk(P) Average dipolar coupling enegy,
$$g_d/k_B = \frac{2\pi\gamma^2\hbar^2}{3v_mk_B}$$
, [K]

he3_tfeff(P) Effective Fermi temperature,
$$T_{F_{eff}} = \frac{\pi^2}{2\gamma_f}$$
, [K]

TODO:

Recalculate F_0^a F_1^a from experimental data (Now there are some extra not-consistent data)! (see Ramm-1970, Halperin-92 ...). Add F_2^a (zero or from Halperin-95). Check g_d .

Pressure-dependent values:

P, bar	0	3	6	9	12	15	18	21	24	27	30	33
T_c , mK	0.929	1.290	1.560	1.769	1.934	2.067	2.177	2.267	2.339	2.395	2.438	2.474
T_{AB} , mK	_	_	_	_	_	_	_	_	2.217	2.137	2.056	1.969
$v_m, \mathrm{cm}^3/\mathrm{mol}$	36.84	33.95	32.03	30.71	29.71	28.89	28.18	27.55	27.01	26.56	26.17	25.75
c_1 , m/s	182.9	227.4	259.9	285.8	307.9	327.3	344.8	360.6	375.2	389.2	402.9	416.1
γ_f , 1/(K mol)	2.784	2.981	3.158	3.321	3.475	3.625	3.772	3.918	4.065	4.211	4.355	4.496
T^{\star} , mK	358.8	305.8	276.2	255.7	238.7	224.0	212.4	204.0	197.8	191.9	184.9	178.4
ρ , g/cm ³	0.082	0.089	0.094	0.098	0.102	0.104	0.107	0.109	0.112	0.114	0.115	0.117
$2N(0), 10^{38}$	1.002	1.164	1.307	1.434	1.551	1.663	1.774	1.885	1.995	2.102	2.207	2.315
$p_F, 10^{-20} \text{ g cm/s}$	8.280	8.509	8.675	8.798	8.896	8.979	9.054	9.122	9.182	9.234	9.280	9.330
$m^{\star}, 10^{-23} \text{ g}$	1.401	1.584	1.744	1.887	2.018	2.144	2.269	2.392	2.515	2.635	2.753	2.872
m^{\star}/m_3	2.797	3.162	3.482	3.767	4.030	4.282	4.530	4.777	5.022	5.261	5.497	5.735
v_F , m/s	59.11	53.73	49.74	46.64	44.08	41.87	39.91	38.13	36.51	35.05	33.71	32.49
$\chi_N, 10^{-9}$	38.1	48.5	56.9	64.1	71.0	77.8	84.1	89.6	94.2	98.8	104.1	109.7
F_0^s	9.27	15.99	22.52	28.91	35.32	41.81	48.43	55.17	62.10	69.32	76.95	84.82
F_1^s	5.39	6.49	7.45	8.30	9.09	9.85	10.59	11.33	12.07	12.78	13.49	14.21
F_0^a	-0.70	-0.72	-0.73	-0.74	-0.75	-0.75	-0.76	-0.76	-0.76	-0.75	-0.76	-0.76
F_1^a	-0.55	-0.73	-0.80	-0.85	-0.90	-0.95	-0.99	-1.00	-0.99	-0.99	-0.99	-1.00
a, Å	3.940	3.834	3.761	3.708	3.668	3.634	3.604	3.577	3.553	3.533	3.516	3.497
$g_d/k_B,\mu\mathrm{K}$	114.5	124.3	131.7	137.4	142.0	146.0	149.7	153.1	156.2	158.8	161.2	163.9
$T_{F_{eff}}$, K	1.773	1.656	1.563	1.486	1.420	1.361	1.308	1.260	1.214	1.172	1.133	1.098
$<\!\!W\!\!>$	83.4	104.1	116.0	123.7	130.7	137.2	141.6	142.4	140.6	139.0	140.3	141.8
$<\!W_I\!>$	10.92	14.68	16.37	17.32	18.23	19.08	19.48	19.19	18.41	17.73	17.63	17.60
$\langle W_D \rangle$	24.89	29.75	32.62	34.42	35.98	37.41	38.38	38.56	38.18	37.88	38.23	38.60
$<\!W_L\!>$	-11.42	-8.14	-6.53	-4.70	-2.28	0.29	2.38	3.62	4.10	4.25	4.53	5.19
γ_0	0.131	0.141	0.141	0.140	0.139	0.139	0.138	0.135	0.131	0.127	0.126	0.124
δ_0	0.298	0.286	0.281	0.278	0.275	0.273	0.271	0.271	0.272	0.272	0.272	0.272
w_0	1.211	1.192	1.187	1.185	1.182	1.180	1.179	1.181	1.184	1.187	1.189	1.189
λ_1^a	-0.137	-0.078	-0.056	-0.038	-0.017	0.002	0.017	0.025	0.029	0.031	0.032	0.037
$\tau_N(0,T_c), \mu s$	0.609	0.237	0.137	0.095	0.072	0.057	0.048	0.043	0.039	0.036	0.034	0.031
$\Delta_{WC+}(T=0)/T_c$	1.771	1.795	1.813	1.825	1.834	1.843	1.850	1.856	1.861	1.866	1.870	1.875
$\chi_B(T=0)/\chi_N$	0.378	0.357	0.346	0.340	0.335	0.331	0.328	0.327	0.328	0.329	0.329	0.328
$\nu_B(T=0), kHz$	100.2	140.7	173.6	200.6	223.1	242.6	260.1	276.0	290.4	302.9	313.4	323.5
P, bar	0	3	6	9	12	15	18	21	24	27	30	33

Energy gap

BCS energy gap + trivial strong coupling correction + some values derived from energy gap. See more information in the source file he3_gap.f.

he3_bcsgap(ttc) BCS gap for 3He-B in T_c units, Δ_{BCS}/k_BT_c

 $\verb|he3_bcsgap_fast(ttc)| Einzel approximation for BCS gap (0.5\% accuracy, 70 times faster)|$

(Einzel-1991, f.68)

he3_trivgap(ttc, P) Trivial strong-coupling correction to the BCS gap

he3_yosida(ttc, gap, n) Yosida functions $Y_n(T/T_c, \Delta) = \int_{-\infty}^{\infty} \left(\frac{\xi_k}{E_k}\right)^n \phi_k \ d\xi_k$

Low temperature limit: $Y_n(T, \Delta) = 2\Gamma\left(\frac{n+1}{2}\right) \left(\frac{T}{\Delta}\right)^{\frac{n-1}{2}} \exp\left(-\frac{\Delta}{T}\right)$

Note: type of n parameter in fortran should be real*8 (this was done for standard function handling). If n is integer, result is unpredictable!

he3_yosida_par(ttc, gap) $Y_q^{\parallel}(T/T_c,\Delta)$

he3_yosida_perp(ttc, gap) $Y_q^\perp(T/T_c,\Delta)$

he3_z3(ttc, gap)

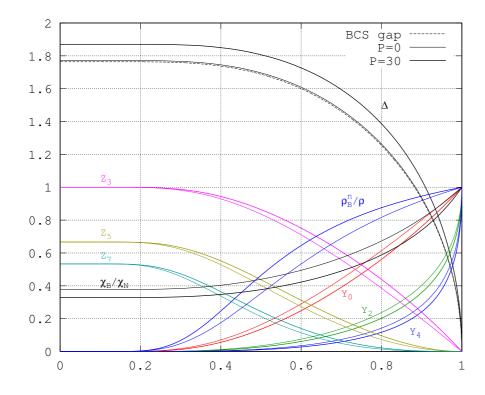
he3_z5(ttc, gap)

he3_z7(ttc, gap)

he3_lambda(ttc, gap)

he3_rho_nb(ttc, p) B-phase normal component density: $\frac{\rho_B^n}{\rho_N} = \frac{(3+F_1^s)Y_0}{3+F_1^sY_0}$

he3_chi_b(ttc, p) B-phase susceptibility: $\frac{\chi_B}{\chi_N} = \frac{(1+F_0^a)(2+Y_0)}{3+F_0^a(2+Y_0)}$



Dipole energy and Legget frequency

Values are given according to Thuneberg's paper JLTP 122, p657 (2001). Temperature independent g_d parameter is restored from experimental data for Ω_B , measured in transverse and longitudinal NMR. See more information in the source file he3_dipole.f.

Dipolar energy:

$$F_D = \lambda_D \int (R_{ii}R_{jj} + R_{ij}R_{ji}) d^3r = 4\lambda_D \int \cos\theta (1 + 2\cos\theta) d^3r$$
$$\lambda_D = \Delta^2 g_d, \qquad \Omega_B = \gamma \sqrt{15\lambda_D/\chi_B}$$

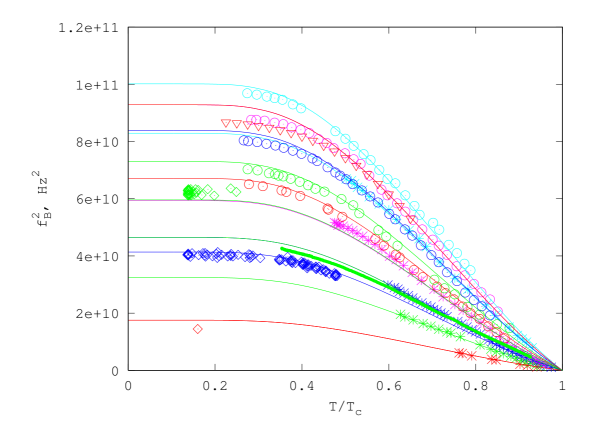
Note: In old papers (Leggett, ...) $g_D/5$ is used instead of λ_G .

he3_gd(p)
$$g_d$$
, $[1/(\text{erg cm}^3)]$

he3_ld(p)
$$\lambda_D = \Delta^2 g_d$$
, [erg/cm³]

he3_nu_b(ttc, p) B-phase Leggett frequency
$$\nu_B = \frac{\gamma}{2\pi} \sqrt{15\Delta^2 g_d/\chi_B}$$
, Hz

he3_nu_b1(ttc, p) Less accurate formula without using
$$g_d$$
, Hz
$$\nu_B = \frac{1}{2\pi} \sqrt{\frac{3\pi}{2\chi}} \; \frac{\gamma^2 \hbar}{2} \; N(0) \; \Delta \log \frac{e_f}{\Delta}$$



Gradient energy and spin wave velocity

Values are given according to Thuneberg's paper JLTP 122, p657 (2001). High order Fermi liquid parameters (F_2^a, F_3^a) are set to 0. See more information in the source file he3_text_gr.f and in doc_tech/egrad.pdf.

$$F_G = \int_V \left[\lambda_{G1} \frac{\partial R_{\alpha i}}{\partial r_i} \frac{\partial R_{\alpha j}}{\partial r_j} + \lambda_{G2} \frac{\partial R_{\alpha j}}{\partial r_i} \frac{\partial R_{\alpha j}}{\partial r_i} \right] \ d^3r + 2\lambda_{G2} \int_S s_k R_{\mu k} \frac{\partial R_{\mu j}}{\partial r_j} \ d^2r$$

 $(s_k \text{ is directed inside the volume})$

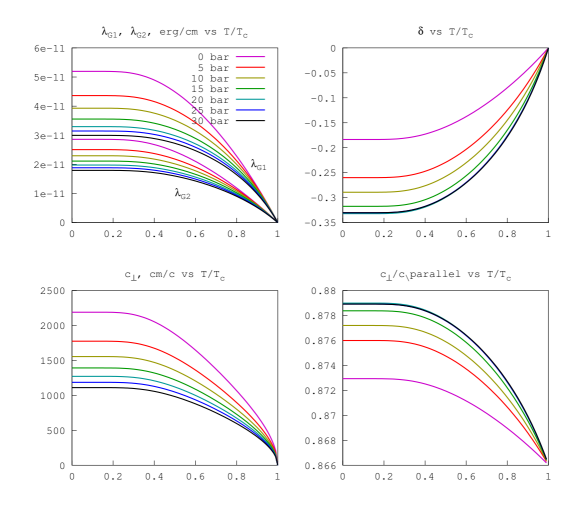
he3_text_lg2(ttc, p)
$$\lambda_{G2} = \frac{\hbar^2 \rho}{40m^* m} \frac{(1 + F_1^a/3)(1 - Y_0)}{1 + F_1^a(2 + 3Y_0)/15}, \, [\mathrm{erg/cm}]$$

he3_text_lg1(ttc, p) $\lambda_{G1}=(2+\delta)\lambda_{G2},\,[{\rm erg/cm}]$

he3_text_delta(ttc, p)
$$\delta = \frac{F_1^a(1-Y_0)}{3+F_1^aY_0}$$

he3_text_cperp(ttc, p) Fomin's perpendicular spin wave velocity. $c_{\perp} = \gamma \sqrt{(6+\delta)\lambda_{G2}/\chi}, \, [\text{cm/s}]$

he3_text_cpar(ttc, p) Fomin's parallel spin wave velocity. $c_{||} = \gamma \sqrt{(8+2\delta)\lambda_{G2}/\chi}, \, [{\rm cm/s}]$



Other textural parameters

Values are given according to Thuneberg's paper JLTP 122, p657 (2001). High order Fermi liquid parameters (F_2^a, F_3^a) are set to 0. See more information in the source file he3_text.f.

Bulk terms:

$$F_{DH} = -a \int_{V} (\mathbf{n} \cdot \mathbf{H})^{2} d^{3}r$$

$$F_{DV} = -\lambda_{DV} \int_{V} [\mathbf{n} \cdot (\mathbf{v_{s}} - \mathbf{v_{n}})]^{2} d^{3}r \ll F_{HV} \text{ at } H \approx 200 \text{ Oe}$$

$$F_{HV} = -\lambda_{HV} \int_{V} [\mathbf{H} \cdot R \cdot (\mathbf{v_{s}} - \mathbf{v_{n}})]^{2} d^{3}r$$

Surface terms:

$$F_{SH} = -d \int_{S} [\mathbf{H} \cdot R \cdot \mathbf{s}]^{2} d^{2}r$$

$$F_{SG} = -\lambda_{SG} \int_{S} s_{j} R_{\alpha j} \frac{\partial R_{\alpha i}}{\partial r_{i}} d^{2}r$$

Vortex term:

$$F_{LH} = \frac{\lambda_{LH}}{2\Omega} \int_{L} |\omega_s| [\mathbf{H} \cdot R \cdot \mathbf{l}]^2 d^3r$$

$$a = \frac{5g_d}{2} \left[\frac{\hbar \gamma/2}{1 + F_0^a (2 + Y_0)/3} \right]^2 \left(5 - 3 \frac{Z_3}{Z_5} \right), \, [\text{erg/cm}^3 \, \text{G}^{-2}]$$

$$\text{he3_text_d(ttc, p)} \qquad d, \, [\text{erg/(cm}^2 \, \text{G}^2)] - \text{GL extrapolation!}$$

$$he3_\text{text_ldv(ttc, p)} \qquad \lambda_{DV} = 5g_d \left(\frac{m^* v_F}{1 + F_1^s Y_0/3} \right)^2 \left(1 - \frac{3Z_5}{2Z_3} \right), \, [\text{erg/cm}^3 \, (\text{cm/s})^{-2}]$$

$$\text{he3_text_lhv(ttc, p)} \qquad \lambda_{HV} = \frac{\rho}{\Delta^2} \frac{1 + F_1^s/3}{(1 + F_1^s Y_0/3)^2} \left(\frac{\hbar \gamma/2}{1 + F_0^a (2 + Y_0)/3)} \right)^2 \left(Z_3 - \frac{9}{10} Z_5 + \frac{9}{10} \, \frac{Z_5^2}{Z_3} - \frac{3}{2} Z_7 \right)$$

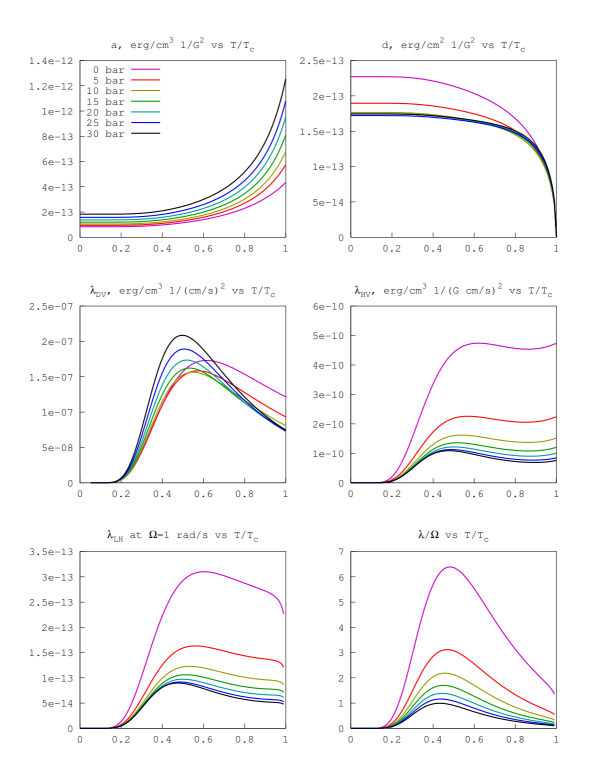
$$\text{he3_text_lhv(ttc, p, omega)} \qquad \lambda_{LH} = \frac{\hbar}{2m} \Omega \lambda_{HV} \left(\ln \frac{R}{r} - \frac{3}{4} \right), \, [\text{erg/(cm}^3 \, \text{G}^2 \, \text{s})] - \text{counterflow part only!}$$

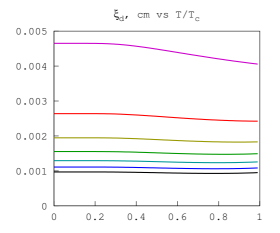
he3_text_lo(ttc, p, omega) lambda/omega used in the texture library: $\frac{\lambda}{\Omega} = \frac{5\lambda_{LH}}{2a\Omega}$, $(rad/s)^{-1}$

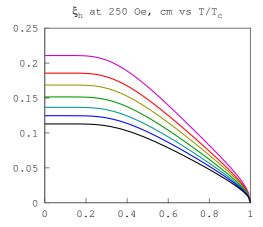
he3_text_xih(ttc, p, h) Magnetic length $\xi_H = \sqrt{65\lambda_{G2}/(8aH^2)}$

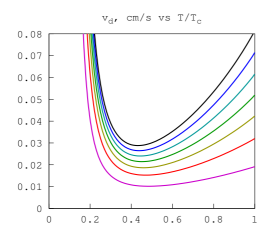
he3_text_xid(ttc, p) Dipolar length (according to Hakonen-1989) $\xi_D = \sqrt{13\lambda_{G2}/12\lambda_D}$. Note: in Thuneberg's paper $\xi_D = \sqrt{\lambda_{G2}/\lambda_D}$ is used.

he3_text_vd(ttc, p) Dipolar velocity (according to Thuneberg-2001) $v_D = \sqrt{2a/5\lambda_{HV}}$









Transport properties in the normal phase

See more information in the source file he3_transp_n.f.

Crossections and scattering factors:

he3_scatt_w(P) Scattering crossection < W > (Einzel-1978, f.82)

 $< W_I >$ (Einzel-1978, f.82) he3_scatt_wi(P)

 $< W_D >$ (Einzel-1978, f.82) he3_scatt_wd(P)

he3_scatt_wl(P) $< W_L > (Einzel-1978, f.71)$

he3_scatt_g0(P) Scattering factor $\gamma_0 = \langle W_I \rangle / \langle W \rangle$ (Einzel-1978, f.66)

Scattering factor $\delta_0 = \langle W_D \rangle / \langle W \rangle$ (Einzel-1978, f.67) he3_scatt_d0(P)

he3_scatt_w0(P) $w_0 = 1 - 3/2\gamma_0 + \delta_0$ (Einzel-1978, f.79)

he3_scatt_l1a(P) Scattering factor λ_1^a used in spin diffusion transport time. Can be neglected.

(Einzel-1978, f.74, p.350).

Normal state quasiparticle lifetime at the Fermi level, s. he3_tau_n0(ttc,p)

 $\tau_N(0,T) = \tau_N(0,T_c) \left(\frac{T_c}{T}\right)^2 = \frac{32E_F\hbar}{\langle W \rangle (\pi k_B T)^2}$ (Fingel 1991, 1992)

(Einzel-1991, p.325)

he3_tau_n_av(ttc,p) Thermal average of normal state quasiparticle lifetime, s.

 $\bar{\tau}_N = \left\langle \frac{\tau_N(0, T)}{1 + (\xi_k / \pi T)^2} \right\rangle = \frac{3}{4} \tau_N(0, T)$ (Einzel-1991, f.4-5)

Spin diffusion transport time for a normal Fermi-liquid, s he3_tau_nd(ttc,p)

 $\tau_{ND} = \tau_1 = \bar{\tau}_N / (1 - \lambda_1^a)$ (Einzel-1991, p.328)

Hydrodynamic spin diffusion in normal liquid, D_0 , cm²/s he3_diffn_hydr(ttc, p)

(Einzel JLTP84 (1991) f.23)

he3_diffn_perp(ttc, p, nu0) Perpendicular component of spin diffusion, reduced because of the

Leggett-Rice effect, D_N^{\perp} , cm²/s (Einzel JLTP84 (1991) f.22)

TODO: Temperature is measured in T_c units which is strange for the normal phase. Maybe it is better to use K of T_F units here.

Transport properties in the B phase

Values for he3_tau0, he3_tau_av, he3_diff_* are extended to $T > T_c$ region using normal phase functions. See more information in the source file he3_transp_b.f and in doc_tech/spin_diff.pdf.

Collision integral for Bogoliubov quasiparticles:

```
he3_coll_int(xi, ttc, gap, g0, d0) Full temperature range approximation
(Einzel, Wolfle, Hirschfeld, JLTP80 (1990), p.66)

he3_coll_int_lt(xi, ttc, gap, g0, d0) Collision integral for low temp (good for < 0.7T<sub>c</sub>)
(Einzel, JLTP84 (1991), f.76)

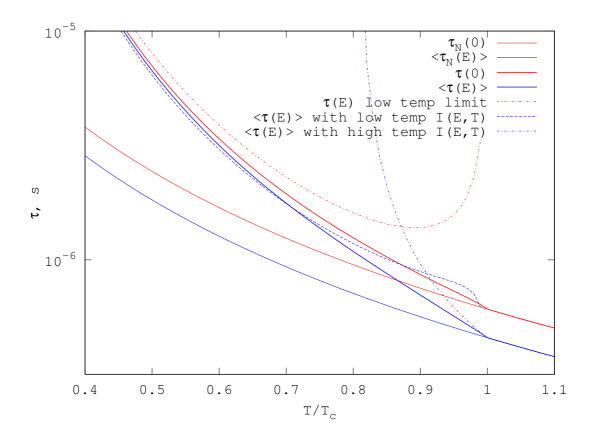
he3_coll_int_ht(xi, ttc, gap, g0, d0) Collision integral for high temp
(Einzel, JLTP84 (1991), f.75)
```

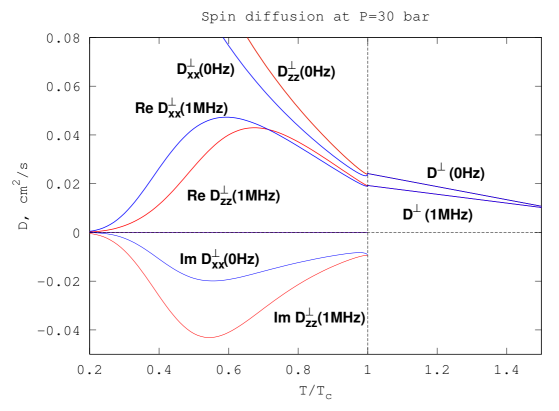
Bogoliubov quasiparticles lifetime and mean free path:

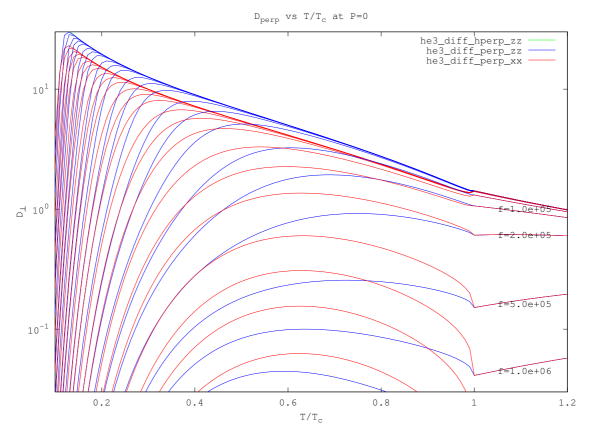
```
Lifetime at the Fermi level, s. 1/\tau(0) = \frac{I(0,T)}{\tau_N(0,T)},
he3_tau0(ttc, p)
                                (Einzel-1991, f.74,76. Einzel-1990 f.A1)
                                1/\tau, T \to 0 limit (does not depend on energy). (Einzel-1978, f.79)
he3_tau0lt(ttc, p)
                               Thermal average lifetime, s. 1/\bar{\tau} = \frac{1}{Y_0} \int_{-\infty}^{\infty} \frac{\phi_k \ d\xi}{\tau}
he3_tau_av(ttc, p)
                                (Einzel-1991, f.77)
he3_fpathv(ttc, p)
                               Mean free path, cm
                                (Einzel JLTP32 (1978) f.84)
                               Spin diff. transport time, s. \tau_D^{\perp,\parallel} = \bar{\tau}/(1-\lambda_1^a Y^{\perp,\parallel})
he3_tau_dperp(ttc, p)
he3_tau_dpar(ttc, p)
                                (Einzel-1991, f.90,96)
   Spin diffusion:
                                              Spin diffusion in hydrodynamic limit (\omega_L \tau \ll 1), D_{zz}^{\perp,\parallel}(\omega_L = 0),
he3_diff_hperp_zz(ttc, p)
he3_diff_hpar_zz(ttc, p)
                                              cm<sup>2</sup>/s. According to Einzel's paper (see f.105) nonhydrodynamic
                                              effects do not affect D^{\parallel}. In Mukharsky paper they do.
                                              (Einzel JLTP84 (1991) f.102)
he3_diff_perp_xx(ttc, p, nu0)
                                              Components of spin diffusion tenzor, cm<sup>2</sup>/s
he3_diff_perp_zz(ttc, p, nu0)
                                              (Bunkov et al, 1990 f.3; Einzel, 1991, f.108;
he3_diff_perp_xx_im(ttc, p, nu0)
                                              Markelov, Mukharsky, 1992, f.7,8)
he3_diff_perp_zz_im(ttc, p, nu0)
he3_diff_par_xx(ttc, p, nu0)
he3_diff_par_zz(ttc, p, nu0)
```

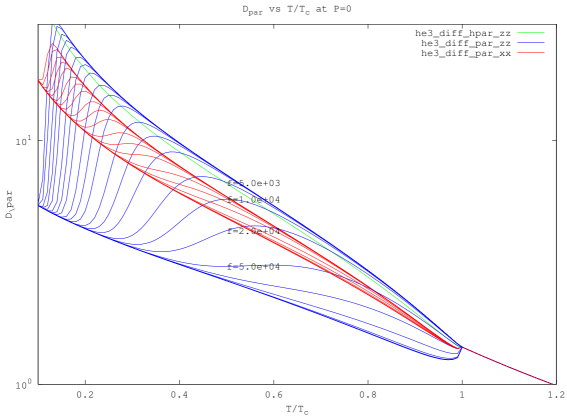
Problem with D^{\parallel} . According to Einzel-1991 (see f.105) nonhydrodynamic effects do not affect D^{\parallel} and you should use he3_diff_hpar_zz for any frequency. In Markelov-Mukharsky paper the result for D^{\parallel} is completely different (it is used in he3_diff_par_xx, _zz). D^{\perp} is the same in all papers. TODO:

Exchange coupling strength is in the $F_1^a=0$ limit. (small difference from normal phase values) How does he3_diff_perp_??_im extends to normal phase?









Other

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he3_xigl(ttc, p) Extrapolated GL coherence length, cm
he3_vneq(ttc, p, omega, r) Equilibrium vortex number
```

Normal liquid beyond zero temperature limit

he3_cv_n(t, p) Normal phase heat capacity C_v , Greywall-1983 TODO: V_m

ROTA specific functions

$$\label{eq:cns} \begin{array}{ll} \text{Nuclear stage heat capacity } [\text{J/K}] \text{ vs } \text{T[K] and I[A]} \\ C_{ns} = 9.66 \cdot 10^{-5} \; [\text{J/T}^2] \; (0.113 \; [\text{T/A}] \; I/T)^2 \\ \\ \text{rota_fork_cal(w, p, n)} \quad \text{Calibration of fork } n, T/T_c, \text{ vs fork width } w \; [\text{Hz}] \text{ and pressure } P \; [\text{bar}] \\ w = a_n \exp \left(-\frac{\Delta_{\text{WC+}}(P, T/T_c)}{T/T_c} \right), \\ a = \alpha_n \; p_F^4(P), \text{ where } \alpha_n \text{ is a geometrical factor} \\ N = 1: \qquad \text{Fork K, calibration } 30.4.2010, 29 \; \text{bar, a} = 11700 \\ N = 2: \qquad \text{Fork E, calibration } 30.4.2010, 29 \; \text{bar, a} = 17543 \\ \end{array}$$