

## Solve multiband Eliashberg equations

—— Outline —
This software provides three programs:
<ol> <li>ebmb itself solves the multiband Eliashberg equations (Eqs. 1 or 4) on a cut-off imaginary axis and optionally continues the results to the real axis via Padé approximants. The normal-state equations (Eq. 7) can also be solved on the real axis.</li> </ol>
A material is defined by nothing but an ELIASHBERG spectral function or, as fallback, an EINSTEIN phonon frequency and intra- and interband electron-phonon couplings, Coulomb pseudo-potentials and, if desired, the band densities of Bloch states, otherwise assumed to be constant.
<ol> <li>critical finds the critical point via the bisection method varying a parameter of choice. Superconductivity is defined by the kernel of the linearized gap equation (Eq. 5 or 6) having an eigenvalue greater than or equal to unity.</li> </ol>
<ol> <li>tc finds the critical temperature for each band separately via the bisection method. Su- perconductivity is defined by the order parameter exceeding a certain threshold. Usually, it is preferable to use critical.</li> </ol>
—— Installation ————
The makefile is designed for the <i>GNU</i> or <i>Intel</i> Fortran compiler:
<pre>\$ make FC=gfortran FFLAGS='-03 -fopenmp' \$ python3 -m pip install -e .</pre>
I/O
Parameters are defined on the command line:
$\ \langle program \rangle \langle key 1 \rangle = \langle value 1 \rangle \langle key 2 \rangle = \langle value 2 \rangle \dots$

- The available keys and default values are listed in Table 1.

   Unless tell=false, the results are printed to standard output.
- Unless file=none, a binary output file is created. For critical and to it simply contains one or more double precision floating point numbers, for ebmb the format defined in Tables 2 and 3 is used.
- The provided *Python* wrapper functions load the results into *NumPy* arrays:

```
import ebmb
results = ebmb.get(\( \)program \>, \( \)file \>, \( \)replace \>,
  \( \)key 1 \>= \( \)value 1 \>, \( \)key 2 \>= \( \)value 2 \>, \( \)...)
```

 $\langle replace \rangle$  decides whether an existing  $\langle file \rangle$  is used or overwritten.

Let  $\hbar=k_{\rm B}=1$ . Fermionic and bosonic Matsubara frequencies are defined as  $\omega_n=(2n+1)\pi T$  and  $\nu_n=2n\pi T$ , respectively. The quantity of interest is the Nambu self-energy matrix<sup>1</sup>

$$\boldsymbol{\Sigma}_{i}(n) = i\omega_{n}[1 - Z_{i}(n)]\mathbf{1} + \underbrace{Z_{i}(n)\,\Delta_{i}(n)}_{\boldsymbol{\phi}_{i}(n)}\boldsymbol{\sigma}_{1} + \chi_{i}(n)\boldsymbol{\sigma}_{3},$$

where the Pauli matrices are defined as usual and i is a band index. Renormalization  $Z_i(n)$ , order parameter  $\phi_i(n)$  and energy shift  $\chi_i(n)$  are determined by the Eliashberg equations<sup>2</sup>

$$Z_{i}(n) = 1 + \frac{T}{\omega_{n}} \sum_{j} \sum_{m=0}^{N-1} \int_{-\infty}^{\infty} d\varepsilon \frac{n_{j}(\varepsilon)}{n_{j}(\mu_{0})} \frac{\omega_{m} Z_{j}(m)}{\Theta_{j}(\varepsilon, m)} \Lambda_{ij}^{-}(n, m),$$

$$\phi_{i}(n) = T \sum_{j} \sum_{m=0}^{N-1} \int_{-\infty}^{\infty} d\varepsilon \frac{n_{j}(\varepsilon)}{n_{j}(\mu_{0})} \frac{\phi_{j}(m)}{\Theta_{j}(\varepsilon, m)} [\Lambda_{ij}^{+}(n, m) - U_{ij}^{*}(m)],$$

$$\chi_{i}(n) = \chi_{C_{i}} - T \sum_{j} \sum_{m=0}^{N-1} \int_{-\infty}^{\infty} d\varepsilon \frac{n_{j}(\varepsilon)}{n_{j}(\mu_{0})} \frac{\varepsilon - \mu + \chi_{j}(m)}{\Theta_{j}(\varepsilon, m)} \Lambda_{ij}^{+}(n, m),$$

$$\Theta_{i}(\varepsilon, n) = [\omega_{n} Z_{i}(n)]^{2} + \phi_{i}^{2}(n) + [\varepsilon - \mu + \chi_{i}(n)]^{2},$$

$$(1)$$

and may then be analytically continued to the real-axis by means of PADÉ approximants.<sup>3</sup> The electron-phonon coupling matrices and the rescaled COULOMB pseudo-potential are connected to the corresponding input parameters via

$$\Lambda_{ij}^{\pm}(n,m) = \lambda_{ij}(n-m) \pm \lambda_{ij}(n+m+1), \qquad \lambda_{ij}(n) = \int_{0}^{\infty} d\omega \frac{2\omega \alpha^{2} F_{ij}(\omega)}{\omega^{2} + \nu_{n}^{2}} \stackrel{=}{\underset{\text{Einstein}}} \frac{\lambda_{ij}}{1 + \left[\frac{\nu_{n}}{\omega_{E}}\right]^{2}}, \\
U_{ij}^{*}(m) = \begin{cases} 2\mu_{ij}^{*}(\omega_{N_{C}}) & \text{for } m < N_{C}, \\ 0 & \text{otherwise,} \end{cases} \frac{1}{\mu_{ij}^{*}(\omega_{N_{C}})} = \frac{1}{\mu_{ij}^{*}} + \ln \frac{\omega_{E}}{\omega_{N_{C}}} \tag{2}$$

with the Eliashberg spectral function  $\alpha^2 F_{ij}(\omega)$  and  $\mu_{ij}^* = \mu_{ij}^*(\omega_{\rm E})$  per definition. Alternatively, if the density of states  $n_i(\varepsilon)$  per spin as a function of energy  $\varepsilon$  is given,

$$\frac{1}{\mu_{ij}^*(\omega_{N_{\mathbb{C}}})} = \frac{1}{\mu_{ij}} + \frac{1}{\pi} \int_{-\infty}^{\infty} d\varepsilon \frac{n_j(\varepsilon)}{n_j(\mu_0)} \begin{cases} \frac{1}{\varepsilon - \mu_0} & \text{arctan } \frac{\varepsilon - \mu_0}{\omega_{N_{\mathbb{C}}}} & \text{for } \varepsilon \neq \mu_0, \\ \frac{1}{\omega_{N_{\mathbb{C}}}} & \text{otherwise,} \end{cases}$$
(3)

where D is the electronic bandwidth.  $\mu_0$  and  $\mu$  are the chemical potentials for free and interacting particles, whose number  $n_0$ , n (including a factor of 2 for the spin) is usually conserved:

$$\sum_{i} \int_{-\infty}^{\infty} d\varepsilon \frac{2n_{i}(\varepsilon)}{e^{(\varepsilon-\mu_{0})/T}+1} = n_{0} \stackrel{!}{=} n = \sum_{i} \int_{-\infty}^{\infty} d\varepsilon \, n_{i}(\varepsilon) \left[ 1 - 4T \sum_{n=0}^{N-1} \frac{\varepsilon - \mu + \chi_{i}(n)}{\Theta_{i}(\varepsilon, n)} - \frac{2}{\pi} \arctan \frac{\varepsilon - \mu + \chi_{C_{i}}}{\omega_{N}} \right].$$

It is unusual but possible to also consider the COULOMB contribution to the energy shift:

$$\chi_{C_i} = \sum_{j} \int_{-\infty}^{\infty} d\varepsilon \frac{n_j(\varepsilon)}{n_j(\mu_0)} \left[ 2T \sum_{m=0}^{N-1} \frac{\varepsilon - \mu + \chi_j(m)}{\Theta_j(\varepsilon, m)} + \frac{1}{\pi} \arctan \frac{\varepsilon - \mu + \chi_{C_j}}{\omega_N} \right] \mu_{ij}.$$

<sup>&</sup>lt;sup>1</sup>Y. Nамви, Phys. Rev. **117**, 648 (1960)

<sup>&</sup>lt;sup>2</sup>G. M. Eliashberg, Soviet Phys. JETP **11**, 696 (1960).

A comprehensive review is given by P. B. ALLEN and B. MITROVIĆ in Solid state physics 37 (1982)

<sup>&</sup>lt;sup>3</sup>H. J. VIDBERG and J. W. SERENE, J. Low Temp. Phys. **29**, 179 (1977)

For a given scalar  $\alpha^2 F(\omega)$ , an effective phonon frequency can be calculated in different ways. We follow ALLEN and DYNES,<sup>4</sup> who define the logarithmic and the second-moment average frequency and use the latter as  $\omega_{\rm E}$  in Eqs. 2 and 3 for rescaling  $\mu^*$ :

$$\omega_{log} = exp \left[ \frac{2}{\lambda} \int_0^\infty \frac{d\omega}{\omega} \alpha^2 F(\omega) \ln(\omega) \right], \qquad \overline{\omega}_2 = \sqrt{\frac{2}{\lambda} \int_0^\infty d\omega \, \alpha^2 F(\omega) \, \omega}.$$

Approximating  $n_i(\varepsilon) \approx n_i(\mu_0)$  yields  $\chi_i(n)=0$  and the constant-DOS ELIASHBERG equations

$$Z_{i}(n) = 1 + \frac{\pi T}{\omega_{n}} \sum_{j} \sum_{m=0}^{N-1} \frac{\omega_{m}}{\sqrt{\omega_{m}^{2} + \Delta_{j}^{2}(m)}} \Lambda_{ij}^{-}(n, m),$$

$$\Delta_{i}(n) = \frac{\pi T}{Z(n)} \sum_{j} \sum_{m=0}^{N-1} \frac{\Delta_{j}(m)}{\sqrt{\omega_{m}^{2} + \Delta_{j}^{2}(m)}} [\Lambda_{ij}^{+}(n, m) - U_{ij}^{*}(m)].$$
(4)

At the critical temperature,  $\phi_i(m)$  is infinitesimal and negligible relative to  $\omega_m$ . This yields

$$\phi_{i}(n) = \sum_{j} \sum_{m=0}^{N-1} K_{ij}(n, m) \,\phi_{j}(m),$$

$$K_{ij}(n, m) = T \int_{-\infty}^{\infty} d\varepsilon \frac{n_{j}(\varepsilon)}{n_{j}(\mu_{0})} \frac{\Lambda_{ij}^{+}(n, m) - U_{ij}^{*}(m)}{\Theta_{j}(\varepsilon, m)},$$
(5)

where  $\Theta_i(\varepsilon, m)$  is obtained from Eqs. 1 for  $\phi_i(m) = 0$ . Similarly, in the constant-DOS case,

$$\Delta_{i}(n) = \sum_{j} \sum_{m=0}^{N-1} K_{ij}(n, m) \Delta_{j}(m),$$

$$K_{ij}(n, m) = \frac{1}{2m+1} [\Lambda_{ij}^{+}(n, m) - \delta_{ij} \delta_{nm} D_{i}^{N}(n) - U_{ij}^{*}(m)],$$

$$D_{i}^{N}(n) = \sum_{j} \sum_{m=0}^{N-1} \Lambda_{ij}^{-}(n, m) \stackrel{N=\infty}{=} \sum_{j} [\lambda_{ij} + 2 \sum_{m=1}^{n} \lambda_{ij}(m)].$$
(6)

 $Z_i(n)$  is not biased by the cutoff if  $D_i^{\infty}(n)$  is used in place of  $D_i^N(n)$  in the kernel  $K_{ij}(n,m)$ . The Eliashberg equations can also be solved on the real axis, which allows for exact analytic continuation without Padé approximants. They are implemented for the normal state:

$$\Sigma_{11i}(\omega) = \underbrace{\sum_{j} \int_{-\infty}^{\infty} d\varepsilon \frac{A_{j}(\varepsilon)}{n_{j}(\mu_{0})} \left[ \mu_{ij} \left( \frac{1}{2} - f(\varepsilon) \right) + \int_{0}^{\infty} d\omega' \alpha^{2} F_{ij}(\omega') \sum_{\pm} \pm \frac{f(\varepsilon) + n(\pm \omega')}{\omega - \varepsilon \pm \omega'} \right]}_{\chi_{C_{i}}}$$
(7)

with the Fermi function  $f(\varepsilon)=1/(\mathrm{e}^{\varepsilon/T}+1)$  and the Bose function  $n(\omega)=1/(\mathrm{e}^{\omega/T}-1)$ . The quasiparticle density of states  $A_i(\omega)=-\frac{1}{\pi}\operatorname{Im} G_i(\omega+\mathrm{i}\eta)$  follows from the Green function

$$G_i(\omega) = -\int_{-\infty}^{\infty} d\varepsilon \, n_i(\varepsilon) \frac{\omega Z_i(\omega) + \varepsilon - \mu + \chi_i(\omega)}{\Theta_i(\varepsilon, \omega)} \stackrel{\phi=0}{=} \int_{-\infty}^{\infty} d\varepsilon \frac{n_i(\varepsilon)}{\omega - \varepsilon + \mu - \Sigma_{11i}(\omega)}.$$

<sup>&</sup>lt;sup>4</sup>P. B. ALLEN and R. C. DYNES, Phys. Rev. B 12, 905 (1975)

 $<sup>^5</sup>$ D. J. Scalapino, J. R. Schrieffer and J. W. Wilkins, Phys. Rev. 148, 263 (1966). See also L. X. Benedict, C. D. Spataru and S. G. Louie, Phys. Rev. B 66, 085116 (2002)

Acknowledgment
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Parts of the program are inspired by the EPW code <sup>6</sup> and work of Malte Rösner.
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Any feedback may be directed to jan.berges@uni-bremen.de.

<sup>&</sup>lt;sup>6</sup>See F. Giustino, M. L. Cohen and S. G. Louie, Phys. Rev. B **76**, 165108 (2007) for a methodology review. Results related to Eliashberg theory are given by E. R. Margine and F. Giustino, Phys. Rev. B **87**, 024505 (2013)

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power	true	_	_	power method for single band?	_	_	+	_

**Table 1:** Input parameters.

- The columns ebmb, tc and critical show which keys are used by these programs.
- The rightmost column indicates which parameters may be chosen as variable for critical. The variable is marked with a negative sign; its absolute value is used as initial guess. If no parameter is negative, the critical temperature is searched for.
- lambda, muStar, and muC expect flattened square matrices of equal size the elements of which are separated by commas. It is impossible to vary more than one element at once.
- dos has lines  $\varepsilon/\text{eV} \ n_1/\text{eV}^{-1} \ n_2/\text{eV}^{-1} \dots$  with  $\varepsilon$  ascending but not necessarily equidistant.
- ullet a2F has lines  $\omega/\mathrm{eV}$   $\alpha^2F_{1,1}$   $\alpha^2F_{2,1}$  ... with  $\omega$  ascending but not necessarily equidistant.
- The relative change in the sample spacing of the real-axis frequencies between  $\omega = 0$  and  $\omega = x$  is logscale  $\omega = x$ . Thus, logscale = 0 corresponds to equidistant sampling.

 $\langle \text{CHARACTERS key} \rangle : \langle n_1 \times \ldots \times n_r \text{ NUMBERS value} \rangle$  associate key with value

DIM:  $\langle \text{INTEGER } r \rangle \langle r \text{ INTEGERS } n_1 \dots n_r \rangle$  define shape (column-major)

INT: take NUMBERS as INTEGERS
REAL: take NUMBERS as DOUBLES

**Table 2:** Statements allowed in binary output. The data types CHARACTER, INTEGER and DOUBLE take 1, 4 and 8 bytes of storage, respectively.

imaginaru-a	imaginary-axis results					
iomega	Matsubara frequency (without i)	$\omega_n$				
Delta	gap $\Delta_i(n)$					
Z	renormalization					
chi	energy shift (*)	$Z_i(n)$ $\chi_i(n)$				
chiC	COULOMB part of energy shift (*) $\chi_{C_i}$					
phiC	COULOMB part of order parameter $\phi_{C_i}$					
status	status (steps till convergence or $-1$ )					
occupancy i	(*) DOS given					
states	integral of density of states	$\frac{\sum_{i} \int d\varepsilon  n_{i}(\varepsilon)}{\sum_{i} \int d\varepsilon  n_{i}(\varepsilon)}$				
inspect	integral of spectral function (**) $\sum_{i=1}^{\infty} \int_{0}^{\infty} d\omega A_{i}(\omega)$					
n0	initial 1	$n_0$				
n	final } occupancy number	n				
mu0	initial )					
mu	final } chemical potential					
effective par	a2F given					
lambda	electron-phonon coupling	$\lambda_{ij}$				
omegaE	EINSTEIN frequency	$\omega_{E}$				
omegaLog	logarithmic average frequency	$\omega_{ m log}$				
omega2nd	second-moment average frequency	$\overline{\omega}_2$				
real-axis re	(**) points $> 0$					
omega	frequency	ω				
Re[Delta]	real } gap	$arDelta_i(\omega)$				
<pre>Im[Delta]</pre>	imaginary $\int g^{ap}$	$\Delta_l(\omega)$				
Re[Z]	real } renormalization	$Z_i(\omega)$				
Im[Z]	imaginary J Tenormatization	$\mathcal{L}_{l}(\omega)$				
Re[chi]	real } energy shift (*)	$\chi_i(\omega)$				
<pre>Im[chi]</pre>	illaginary )					
DOS	quasiparticle density of states (*)	$A_i(\omega)$				
measurable		measurable=true				
Delta0	measurable gap	$\Delta_{0i} = \text{Re}[\Delta_i(\Delta_{0i})]$				
status0	status of measurable gap	_				

Table 3: Keys used in binary output.