VARIATIONAL DIAGRAMMATIC MONTE CARLO AND THE UNIFORM ELECTRON GAS

Jouvence, May 28, 2024

Support:



Simons Collaboration on the Many Electron Problem



Blavatnik Awards Young Scientists



Advertisement: DFT+embedded DMFT

https://github.com/ru-ccmt/eDMFT Easy to follow **tutorials**, easy to install (Python/C++/Fortran) http://hauleweb.rutgers.edu/tutorials/

One of the first DFT+DMFT implementations with many advanced & unique features:

- High throughput scripts for eDMFT calculation, including magnetic materials
- Exact double-counting between LDA&DMFT (PRL 115, 196403 (2015))
- Forces on all atoms
- Structural relaxations within eDMFT free energy functional
- Phonons within eDMFT
- LAPW precise basis set for all electrons (with Wien2k)



DFT + embedded DMFT Functional*

Developed by Kristjan Haule at Rutgers University, ©Copyright 2007-2024.





Why bother with uniform electron gas?

- Solution of UEG serves as a proof of principle that tests the capability of a method to address realistic materials with long range Coulomb repulsion (beyond simplified models).
- Such solution offers new insights into the ab-intion methods (DFTs and GWs), and more understanding of screening in solids.

Variational Diagrammatic Monte Carlo (VDMC) [1,2] allows

- liquid parameters, spin & charge susceptibilities.
- It also provides XC-kernel needed in TDDFT community [3].
- It settles the debate on bandwidth in electron gas, as relevant for Na metal.
- It is useful in other fields, i.e., warm dense matter field uses the same model at higher temperature, where VDMC performs even better.
- achieved in DFT community, as well as recently by DFT+eDMFT method [4]).

VDMC:

[1] Kun Chen, K. Haule, Nature Communications 10, 3725 (2019) [2] K. Haule, K. Chen, <u>Scientific Reports</u> **12**, 2294 (2022) [3] J. P. F. LeBlanc, K. Chen, N.V. Prokof'ev, K.H., Igor S. Tupitsyn, PRL 129 (24), 246401 (2022).

[4]Kamal Choudhary et.al., npj Computational Materials 6, 1 (2020).

• very precise determination of certain physical observables in electron gas: effective mass, landau-

• VDMC could be developed into electronic structure method for high-throughput calculation (like

Build database of accurate electronic structure calculations

History : Uniform electron gas

Is at the heart of the DFT success for materials property prediction.

 $V_{xc} = \frac{\delta E_{xc}[n]}{\delta n}$

	_
1928	Dirac's relativistic theory of the electron Bloch's theory of electrons in solids Pauli-Sommerfeld free electron theory of metals
1934	Wigner's proposal of the Wigner crystal
1956	Landau's theory of Fermi liquids
1957	BCS theory of superconductivity
1964	Hohenberg-Kohn-Sham DFT
1980	Ceperley-Alder QMC prediction of Exc
	many properties of UEG remain unknown

 $E_{xc}[n]$

$$f_{xc}[n](\mathbf{r},\mathbf{r}',\omega) = \frac{\delta V_{xc}[n](\mathbf{r},\omega)}{\delta n(\mathbf{r}'\omega)}$$

Remains essentially unknown to this day Needed in TDDFT

Very little is known: spin susceptibility, Landau parameters,...

high temperature at warm dense matter (plasma) conditions

Diffusion MC simulation of UEG (trajectories in imaginary time) J. Chem. Phys. **151**, 014108 (2019)





The Solid to Uniform Electron Gas Problem

$$H = \sum_{s} \int d^{3}\mathbf{r} \psi_{s}^{\dagger}(\mathbf{r}) \left[-\frac{\nabla^{2}}{2m} + V_{e-n}(\mathbf{r}) \right] \psi_{s}(\mathbf{r}) + \frac{1}{2} \sum_{ss} V_{e-n}$$
 electron-nuclei interaction

$$H_{n-n} \quad \text{nuclei-nuclei interaction}$$

$$V_{C}(\mathbf{r} - \mathbf{r}') = \frac{1}{4\pi\varepsilon_{0}|\mathbf{r} - \mathbf{r}'|}$$
neglecting spin-orbit coupling
Born-Oppenheimer : H_{n-n} and V_{e-n} just
Uniform electron gas: $V_{e-n}(\mathbf{r}) = -\int d^{3}\mathbf{r}' V_{C}(\mathbf{r})$
e-n and n-n terms diverge, but they cancel out

$$H = \sum_{\mathbf{k},s} \frac{k^2}{2m} \psi_{\mathbf{k},s}^{\dagger} \psi_{\mathbf{k},s} + \frac{1}{2V} \sum_{\mathbf{q}\neq 0,\mathbf{k}\mathbf{k}',ss'} \psi_{\mathbf{k}+\mathbf{q},s}^{\dagger} \psi_{\mathbf{k}'-\mathbf{q},s'}^{\dagger} V_c(\mathbf{q}) \psi_{\mathbf{k}'s'} \psi_{\mathbf{k},s}$$

notice the absence of q=0 term, which is diverging and cancels out.

 $\sum \int d^3 \mathbf{r} d^3 \mathbf{r}' \psi_s^{\dagger}(\mathbf{r}) \psi_{s'}^{\dagger}(\mathbf{r}') V_c(\mathbf{r} - \mathbf{r}') \psi_{s'}(\mathbf{r}') \psi_s(\mathbf{r}) + H_{n-n}$

a classical potentials

 $(\mathbf{r} - \mathbf{r'})n_0$ where n_0 is constant neutralizing density ut exactly, so that the final Hamiltonian is simplified to

Significance of Uniform electron gas for DFT

$$E = \langle \Phi_0 | H | \Phi_0 \rangle = \langle \Phi_0 | T + H_{e-e} + V_{e-n} | \Phi_0 \rangle = \langle \Phi_0 | T + H_{e-e} | \Phi_0 \rangle + \int d^3 \mathbf{r} \, V_{e-n}(\mathbf{r}) \, n(\mathbf{r})$$

Hohenberg-Kohn theorem: Ground state electron density $n(\mathbf{r})$ is V-representable.

The knowledge of $n(\mathbf{r})$ alone gives knowledge of the external potential and hence the Hamiltonian H. If the Hamiltonian is uniquely determined from density, then the ground state is also a functional of the density only. (The ground state might be degenerate, but the universality of the functional can still be proven.)

$$F[\{n\}] \equiv \langle \Phi_0^{n(\mathbf{r})} | T + H_{e-e} | \Phi_0^{n(\mathbf{r})} \rangle$$

$$F[\{n\}] = \langle \Phi_0^{n(\mathbf{r})} | \sum_s \int d^3 \mathbf{r} \,\psi_s^{\dagger}(\mathbf{r}) [-\frac{\nabla^2}{2m}] \psi_s(\mathbf{r}) + \frac{1}{2} \sum_{ss'} \int d^3 \mathbf{r} d^3 \mathbf{r} \,d^3 \mathbf{r}' \,\psi_s^{\dagger}(\mathbf{r}) \psi_{s'}(\mathbf{r}') V_c(\mathbf{r} - \mathbf{r}') \psi_{s'}(\mathbf{r}') \psi_s(\mathbf{r}) | \Phi_0^n | \Phi$$

Universal functional can be computed from the simplest possible interacting model, i.e., the uniform electron gas model???

Hohenberg-Kohn theorem: $\langle \Phi_0 | T + H_{e-e} | \Phi_0 \rangle$ is universal functional of the density $n(\mathbf{r})$, i.e.,



Significance of Uniform electron gas for DFT

 $F[\{n\}]$ The functional is non-local even in UEG:

We want to find a part of the functional for which a local-type approximation is good.

 $E = \int d^3 \mathbf{r} V_{e-n}(\mathbf{r}) n(\mathbf{r}) + E_H[\{n\}] + T_0[\{n\}] + E_{xc}[\{n\}]$ $E_H[\{n\}] = \frac{1}{2} \int d^3 \mathbf{r} d^3 \mathbf{r'} n(\mathbf{r}) V_C(\mathbf{r} - \mathbf{r'}) n(\mathbf{r'})$

- $T_0[\{n\}]$ is not the exact kinetic energy, but just the kinetic energy of the corresponding non-interacting system. We do not even know how to express the total kinetic energy or the exchange energy as a functional of density. They can be expressed exactly with the density matrix.
- $E_{xc}[\{n\}]$ turns out to be a piece that is amenable to local approximation.

$$E_{xc} \approx \int d^3 \mathbf{r} \, n(\mathbf{r}) \varepsilon_{xc}^{UEG} [n(\mathbf{r})]$$

map solid point by point to UEG



LDA:

It is unlikely that we will ever be able to compute functional F[{n}] exactly even for the uniform electron gas.

$$| \equiv \langle \Phi_0^{n(\mathbf{r})} | T + H_{e-e} | \Phi_0^{n(\mathbf{r})} \rangle$$



to compute XC energy and XC potential



DFT is pretty good for ground state properties (exact DFT is exact) But DFT has well known "gap problem" when trying to interpret KS spectra as physical excitations



$$\chi^{-1}(\mathbf{r},\mathbf{r}';\omega) = \chi_{KS}^{-1}(\mathbf{r},\mathbf{r}';\omega)$$

density time response: $\chi(\mathbf{r},\mathbf{r}',\tau) = -\langle \psi^{\dagger}(\mathbf{r},\tau)\psi(\mathbf{r},\tau)\psi^{\dagger}(\mathbf{r}',\tau')\psi(\mathbf{r}',\tau')\rangle$

Kohn-Sham non-interacting response (RPA bubble):

from Richard Martin et.al., Interacting electrons

The same idea was extended by Gross&Kohn in 1985 to compute the excited state properties (PRL 55, 2850):









Hohenberg-Kohn for GS DFT: One can not find two different $V_{\rho-n}$ potentials that give rise to the same electron density $n(\mathbf{r})$ in the ground state.

$$H(t) = T + H_{e-e} + V_{e-e}$$

Runge–Gross theorem (PRL 52, 997, (1984)): One can not find two different $V_{e-n}(t)$ $V'_{e-n}(t)$ potentials that give rise to the same electron density $n(\mathbf{r}, t)$, if $n(\mathbf{r}, t)$ is time evolved by H(t) from the ground state.

Gross&Kohn (PRL 55, 2850, (1985)):

using time-dependent Schroedinger Eq. the response of the interacting electrons is

$$\chi^{-1}(\mathbf{r},\mathbf{r}';\omega) = \chi_{KS}^{-1}(\mathbf{r},\mathbf{r}';\omega) - V_C(\mathbf{r}-\mathbf{r}') - f_{xc}(\mathbf{r},\mathbf{r}',\omega)$$

where
$$f_{xc}[\{n\}](\mathbf{r},\mathbf{r}';\omega) = \frac{\delta V_{xc}[\{n\}](\mathbf{r},\omega)}{\delta n(\mathbf{r}',\omega)}$$
 is

-n(t)add time-dependence to external potential

- Caveat: $V_{e-n}(t)$ has to be expandable in Taylor series (analytic in time) and $V_{e-n}(t)$ and $V'_{e-n}(t)$ differ for more than c(t)

 $f_{xc}[\{n\}]$? But what is universal functional of n...







Original idea was to take the unknown $f_{xc}[\{n\}]$ from the uniform electron gas. But we do not know $f_{xc}[\{n\}]$ in UEG.

If we assume $f_{xc}[\{n\}]$ is local to a point in 3D space and local in time (constant in frequency) than:

$$f_{xc}[\{n\}](\mathbf{r},\mathbf{r}';\omega=0) = \frac{\delta V_{xc}[\{n\}](\mathbf{r},\omega=0)}{\delta n(\mathbf{r}',\omega=0)} = \frac{\delta^2 E_{xc}[\{n\}]}{\delta n^2} \bigg|_{n=n_0} \delta(\mathbf{r}-\mathbf{r}')$$
Adiabatic LDA

Considerably improves (compared to LDA) the excitation energies of molecules

Atom	ω_{exp}	ω_{ALDA}	$\omega_{LDA}^{(0)}$
Не	1.56 Ry	1.552	_
Be	0.388	0.399	0.257
Mg	0.319	0.351	0.249
Ca	0.216	0.263	0.176
Zn	0.426	0.477	0.352
Sr	0.198	0.241	0.163
Cd	0.398	0.427	0.303

 ${}^{1}S \rightarrow {}^{1}P$ excitation energies in two-valence-electron atoms.

Not much better gaps or optical excitations in semiconductors.



Optics is q->0 charge response, which is in TDDFT

If we want a substantial change of optics in semiconductors, than we require the form:

Phenomenological ansatz works really well: $f_{xc}(\mathbf{r},\mathbf{r'}) = -\frac{0.2}{4\pi |\mathbf{r}-\mathbf{r'}|}$

But each semiconductor needs different number

Conclusion: f_{xc} is highly non-local

$$\chi(\mathbf{q},\omega) = \frac{\chi_{KS}(\mathbf{q},\omega)}{1 - \chi_{KS}(\mathbf{q},\omega)[\frac{4\pi e^2}{q^2} + f_{xc}(\mathbf{q},\omega)]}$$

 $\lim_{q\to 0} f_{xc}(\mathbf{q},\omega) = \frac{\alpha(\omega)}{\mathbf{q}^2}$ Should be singular in semiconductors at zero frequency, but not in metals, like UEG.



Nazarov&Vignale&Chang (PRL 102, 113001, (2009)):

Instead of TDDFT for density-density response function, we might use current-current response functions.

Time Dependent Current Density Functional Theory (TDCDFT):

$$\hat{\chi}^{-1}(\mathbf{q},\mathbf{q}',\omega) = \hat{\chi}_{KS}^{-1}(\mathbf{q},\mathbf{q}',\omega) - \hat{f}_{xc}(\mathbf{q},\mathbf{q}',\omega) - \frac{4\pi ec}{\omega^2} \delta_{\mathbf{q},\mathbf{q}'} \frac{\vec{e}_{\mathbf{q}} \otimes \vec{e}_{\mathbf{q}}}{\mathbf{q}^2}$$

where $\hat{\chi}$ is current-current response function. $\hat{f}_{xc} \rightarrow f_{xc}^L \& f_{xc}^T$ has two components: longitunidal and transverse

Local approximation on longitudinal & transverse f_{xc} seems a much better approximation as it leads to desired form for the charge f_{xc} $\lim_{\mathbf{q}\to 0} f_{xc}(\mathbf{q}, 0) = \frac{\alpha(\omega)}{\mathbf{q}^2}$

Namely:
$$\lim_{\mathbf{q}\to 0} f_{xc}(\mathbf{q},\omega) = \frac{1}{n_0^2 q^2} \sum_{\mathbf{G}\neq 0} (\mathbf{G} \cdot \mathbf{e}_{\mathbf{q}})^2 [f_{xc}^L(\mathbf{G},\omega) - f_{xc}^L(\mathbf{G},\omega=0)] |n_0(\mathbf{G})|^2$$

 $f^{L}_{xc}(\omega)$ is not known in uniform electron gas, hence this was not evaluated yet. Only phenomenological kernels are used in practice.

Bandwidth of alkali metals, correspond to r_s~4

Bandwidth of Na metal is controversial for 35 years:

-ARPES bandwidth show reduction for 18-25% [1,2] (newer 2021 data 10%) -some GW calculation reproduce reduction [3], most do not. -DMC shows increased bandwidth, not reduced [5] because of fixed node approximation.

- [1] E. Jensen & E.W. Plummer, PRL 55, 1912–1915, (1985).
- [2] I.-W. Lyo & E.W. Plummer, PRL 60, 1558–1561, (1988).
- [3] J.E. Northrup, M.S. Hybertsen, & S.G. Louie, PRL 59, 819 (1987).
- [4] X. Zhu, & A.W. Overhauser, RPB 33, 925(1986).
- [5] R. Maezono, M.D. Towler, Y Lee, & R.J. Needs, PRB 68, 165103, (2003).

[6] J. McClain, J. Lischner, T. Watson, D.A. Matthews, E. Ronca, S.G. Louie, T.C. Berkelbach, G. K-L Chan, PRB 93, 235139 (2016)



Exp2: D. V. Potorochin, B. Buechner et.al., arXiv:2112.00422

Variational Diagrammatic Monte Carlo

Diagrammatic MC: provided numerically exact solution by summing sufficiently high-order Feynman diagrams*

Variational Diag-MC:

- Yukawa form) to achieve fast convergent series.
- group of Hugenholtz diagrams)

Kun Chen and K. Haule, Nature Communications 10, 3725 (2019).

* N. Prokof'ev, B. Svistunov, PRL 81, 2514 (1998) N. Prokof'ev. B. Svistunov, PRB 77, 020408 (2008)

• variational principle to determine best starting point (such as screening by

• leverage sign blessing: exact summation of diagrams that largely cancel optimizing internal variables (such as the conserving Baym-Kadanoff



Variational Perturbation Theory

Started with *Kleinert & Feynman* Later improved by *Kleinert & Janke*

VOLUME 75

Convergent Strong-Coupling Expansions from Divergent Weak-Coupling Perturbation Theory

Anharmonic oscillator: $V(x) = \frac{1}{2}\omega^2 x^2 + g$ Weak coupling series is diverging at small Rearrange perturbation: $V(x) = \frac{1}{2}$

Perform expansion in powers of ξ : $E^{(1)}$ Principle of minimum sensitivity: $\frac{dE^{(1)}}{d}$ Final expansion: $E^{(1)}$

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$$gx^4$$

all
$$\omega$$
: $E_0 = \frac{\omega}{2} + g \frac{3}{4\omega^2} - g^2 \frac{21}{8\omega^5} + g^3 \frac{333}{16\omega^8} + \cdots$
 $\frac{1}{2}\Omega^2 x^2 + \xi(gx^4 + \frac{1}{2}(\omega^2 - \Omega^2)x^2)$
variational counter-term
 $\xi = 1$ set to unity at the end
 $\xi = 1$ set to unity at the end
 $\xi = 1$ set to unity at the end
 $\xi = 0 \rightarrow \Omega^n_{optimal}$
 $[\Omega^1_{optimal}], E^{(2)}[\Omega^2_{optimal}], \cdots$

Variational Perturbation Theory

Check first order: $H = H_0 + \xi (gx^4 + \frac{1}{2}(\omega^2 - \Omega^2)x^2)$ Expansion: $E^{(1)} = \langle \psi_0 | H | \psi_0 \rangle = \frac{\Omega}{2} + \xi \left(g \frac{3}{4\Omega^2} + \frac{1}{2} \frac{\omega}{2} + \frac{1}{2} \frac{\omega}{2} \right)$ perturba correc $\frac{dE^{(1)}}{d\Omega}$ Principle of minimum sensitivity:

At $\omega =$

Final first order: $E^{(1)}[\Omega_{or}^{(1)}]$

 E^{exact} Exact result:

$$\Omega^3 - \omega^2 \Omega - 6g = 0$$

= 0 $\Omega_{optimal}^{(1)} = (6g)^{1/3}$

$$p_{ptimal}^{(1)}] = g^{1/3} \frac{3}{8} 6^{1/3} \approx g^{1/3} \, 0.68142$$

$$f = g^{1/3} \, 0.66798$$

Turned diverging series into fast converging series

Variational Perturbation Theory

Higher order terms are well behaved and rapidly converging

Even odd term optimization:





Variational Diagrammatic Monte Carlo

Lagrangian + counter-terms:

- achieve fast convergence : screened short-range interaction in solids or DFT+DMFT solution the problem.
- 2) Optimize parameters in ΔL with principal of the minimal sensitivity, or more complicated Lagrangian with counter-terms.
- convergence (use sign blessed groups to avoid sign problem)

 $L = L_0 + \Delta L(\xi)$

1) choose a good reference system (L_0), which allows for emergent property. We want to leverage the locality of correlations (as known from success of LDA and DMFT) to

renormalized condition. ΔL makes L exact, hence ΔL is not just the interaction, but

3) Use *Diagrammatic Monte Carlo* to evaluate Feynman expansion to high order until

Uniform Electron gas as testbed for method development



with Hubbard-Stratonovich can be transformed to

$$L = \sum_{\mathbf{k}\sigma} \psi_{\mathbf{k}\sigma}^{\dagger} \left(\frac{\partial}{\partial \tau} - \mu - \frac{\hbar^2 \nabla^2}{2m} \right) \psi_{\mathbf{k}\sigma} + \sum_{\mathbf{q}\neq 0} \Phi_{\mathbf{q}}^{\dagger} \frac{q^2}{8\pi} \Phi_{\mathbf{q}} + \frac{i}{\sqrt{2V}} \sum_{\mathbf{q}\neq 0} \rho_{\mathbf{q}} \Phi_{\mathbf{q}}^{\dagger} + \rho_{-\mathbf{q}} \Phi_{\mathbf{q}}$$
$$\phi_{\mathbf{q}}(\mathbf{r}) \qquad \qquad \psi_{\mathbf{k}\sigma}(\mathbf{r})$$

boson that mediates the interaction electron operator

$$\overline{\gamma} \sum_{\mathbf{q}\neq 0} \rho_{\mathbf{q}} \frac{8\pi}{\mathbf{q}^2} \rho_{-\mathbf{q}} \qquad \qquad \frac{2}{|\mathbf{r}-\mathbf{r}'|} \stackrel{\mathsf{FT}}{\to} \frac{8\pi}{\mathbf{q}^2}$$

Uniform Electron gas, a testbed for method development

$$L = L_0 + \Delta L(\xi)$$
$$L_0 = \sum_{\mathbf{k}\sigma} \psi^{\dagger}_{\mathbf{k}\sigma} \left(\frac{\partial}{\partial \tau} - \mu - \frac{\hbar^2 \nabla^2}{2m}\right)$$

 $\Delta L =$

 $\phi_{\mathbf{q}}(\mathbf{r})$

boson that mediates the interaction



 $\psi_{\mathbf{k}\sigma}(\mathbf{r})$

electron operator

VDMC for electron gas

boson that mediates the interaction



electron operator

VDMC for electron gas



Coulomb interaction is static and short ranged



Counter terms make sure that we get the exact answer at large order for any λ

Open question: How to determine parameters λ and v_k



$$\begin{split} (\xi = 1) \end{pmatrix} \psi_{\mathbf{k}\sigma} + \sum_{\mathbf{q} \neq 0} \Phi_{\mathbf{q}}^{\dagger} \frac{q^{2} + \lambda_{\mathbf{q}}}{8\pi} \Phi_{\mathbf{q}} & \text{Kun Chen} \\ \frac{\lambda_{\mathbf{q}}}{8\pi} \Phi_{\mathbf{q}} + \sqrt{\xi} \frac{i}{\sqrt{2V}} \sum_{\mathbf{q} \neq 0} \rho_{\mathbf{q}} \Phi_{\mathbf{q}}^{\dagger} + \rho_{-\mathbf{q}} \Phi_{\mathbf{q}} \\ \frac{\partial_{\mathbf{q}} \Phi_{\mathbf{q}}}{\partial_{\mathbf{q}} (i\omega)} = \frac{1}{i\omega + \mu - \frac{k^{2}}{2m} - v_{\mathbf{k}}} \\ electron \text{ propagator is optimized} \end{split}$$

(DFT KS-potential or DMFT self-energy, etc)

Counter-term makes sure that the exact answer is obtained for any v_k at large p.o.

Screening length

Possible choices for λ :

1)
$$\lambda = -\widetilde{\Pi}_{\mathbf{q}=0,\omega=0}$$
 ho

2)
$$\frac{\lambda}{8\pi} = -\widetilde{\Pi}_{\mathbf{q}=0,\omega=0}^{N=1}$$
 Exact

3)
$$\frac{d\widetilde{\Pi}_{\mathbf{q}\omega=0}}{d\lambda} = 0 \rightarrow \lambda$$
 (born d)

I) Poor convergence and rapid oscillations with orders (approx. 5-times too small) 2) To converge we need to go to order $25=8\pi$! (approx. 5 times too large) 3) The best choice is due to variational perturbation theory, i.e., still quite small perturbation order, but quite monotonic convergence to exact answer.



e principle of smallest sensitivity. rowed from variational perturbation theory)

Example: expansion for polarization



 $\xi\lambda_q$

 \bigcirc

$$\frac{+\lambda}{\pi} - \xi \frac{\lambda}{8\pi} - \xi P_{\mathbf{q}}^0 - O(\xi^2) \cdots \right)^{-1}$$

+ ()+ 2n order correction

+ () ...) + ... 3rd order correction



From sign problem to sign blessing

How to group diagrams to sign-blessed groups?

Symmetry preserved in each group:

Crossing symmetry, spin rotational symmetry, ... At the lowest order leads to "Hugenholtz diagrams"



Ward identity (each MC step is conserving):

Baym-Kadanoff algorithm is used to construct groups of diagrams with consistent internal variables (preserve particle number, energy, momentum **in each MC step**).

Vertex renormalization:

Make sure to combine diagram with the corresponding counter-term that cancels the high-energy contributions





Kun Chen and K. Haule, Nature Communications 10, 3725 (2019).

dielectric constant-direct comparison to DMC





Kun Chen and K. Haule, Nature Communications 10, 3725 (2019).

Calculated values at different densities. VDMC get four significant digits at order N=6.





Kun Chen and K. Haule, Nature Communications 10, 3725 (2019).

Spin-susceptibility & local field correction



LDA excellent approximation up to $k=k_F$. RPA much worse.



effective mass

 $\frac{m}{m^*} = Z$

- Over the last 50 years, the mass in electron gas was controversial, some theories predicting monotonic behavior with density, and other with a turning point.
- in moderately correlated systems.



K. Haule and Kun Chen, Scientific Reports 12, 2294 (2022)

$$Z\left(1+\frac{m}{k_F}\frac{d\Sigma(k_F,\boldsymbol{\omega}=0)}{dk}\right)$$



Important for understanding which method predicts better Bloch bands and bandwidths

Quasiparticle dispersion near the fermi level is defined by effective mass m*/m.

DFT assumes m*/m=1 (non-interacting Kohn-Sham ansatz)

Exact solution (VDMC) remarkably close to m*/m~1. Bounded by vertex corrected perturbation theory using local field factors.

GOWO and QSGW overestimate mass GW underestimates mass

At the uniform density limit, **DFT ansatz is** remarkably accurate, better than GW.

[G0W0] L. Hedin, *Phys. Rev.* 139, A796–A823, (1965). [G+&G-]Simion, G. E. & Giuliani, PRB 77, 035131, (2008). [QSGW] A.Kutepov, G. Kotliar, arXiv: 1702.04548 [GW] K.Van Houcke, et.al., Phys. Rev. B 95, 195131 (2017)

Uniform electron gas: Landau parameters

Landau parameters for UEG.

have never been computed before by controlled method

r _s	Z	m^*/m	F_0^a	F_0^s
1	0.8725(2)	0.955(1)	-0.171(1)	-0.209(5)
2	0.7984(2)	0.943(3)	-0.271(2)	-0.39(1)
3	0.7219(2)	0.965(3)	-0.329(3)	-0.56(1)
4	0.6571(2)	0.996(3)	-0.368(4)	-0.83(2)



compressibility diverges at r_s=5.2, and expansion breaks down



- F_{0} is going critical at $r_{s}=5.2$, where polarization and compressibility diverges.

Polarization also diverges at this point, signaling subtle instability

Bandwidth of Na metal is controversial for 35 years:

-ARPES bandwidth show reduction for 18-25% [1,2] -some GW calculation reproduce reduction [3], most do not. -DMC shows increased bandwidth, not reduced [5].



K. Haule and Kun Chen, Scientific Reports 12, 2294 (2022)

[1] E. Jensen & E.W. Plummer, PRL 55, 1912–1915, (1985).

[2] I.-W. Lyo & E.W. Plummer, PRL 60, 1558–1561, (1988).

[3] J.E. Northrup, M.S. Hybertsen, & S.G. Louie, PRL 59, 819 (1987).

[4] X. Zhu, & A.W. Overhauser, RPB 33, 925(1986).

[5] R. Maezono, M.D. Towler, Y Lee, & R.J. Needs, PRB 68, 165103, (2003).

[6] J. McClain, J. Lischner, T. Watson, D.A. Matthews, E. Ronca, S.G. Louie, T.C. Berkelbach, G. K-L Chan, PRB 93, 235139 (2016)



Exp1: E. Jensen & E.W. Plummer, PRL 55, 1912–1915, (198 Exp2: D. V. Potorochin et.al., arXiv:2112.00422

Real frequency quantities: exchange-correlation kernel

Recently we developed real-frequency diag-MC for uniform electron gas. $\chi(\mathbf{q},\omega) = P_{KS}^{0}(\mathbf{q},\omega) + P_{KS}^{0}(\mathbf{q},\omega)[V_{q} + f_{xc}(\mathbf{q},\omega)]\chi(\mathbf{q},\omega)$ In UEG we compute: $f_{xc}(q,\omega) = \frac{1}{P^0(q,\omega)} - \frac{V_q}{1 - \varepsilon(q,\omega)}$ $f_{xc}(q,\omega)$ on real frequency axis N=1q = 0.1 kFN=2N=2 $Re(f_{xc}[Ry])$ $r_s=2$ 10 **→** N=1 RPA -1010 $m(f_{xc}[Ry])$ -10 $q=0.1 k_{F}$ 50 $r_s=2$ −30 + 0.0 0.2 0.8 1.0 0.0 0.4 0.6 1.2 0.2 8.0 0.4 0.6 1.0 1.2 ω/EF ω/EF

dielectric function on real frequency axis 50



I. S. Tupitsyn, A. M. Tsvelik, R. M. Konik, and N. V. Prokof'ev, PRL 127, 026403 (2021)

Challenging to calculate, but a lot of non-trivial structure below EF. Such change of sign was needed in Si to explain optical data (PRL 102, 11301 (2009)).





Screening in UEG on the two particle level

$$V(r) = \frac{e^2}{4\pi\varepsilon_0 r} e^{-r/\xi} \quad \text{wher}$$

$$\xi = \frac{1}{\sqrt{\lambda}} \approx \frac{1}{\sqrt{E_F}} = \frac{1}{\sqrt{E_F}}$$

Na metal is close to electron gas with r_s~4 and E_F~3eV

$$\xi_{Na} \approx 2r_B \approx 0.8 R_{MT} \approx$$

VDMC:

[1] Kun Chen, K. Haule, Nature Communications 10, 3725 (2019) [2] K. Haule, K. Chen, <u>Scientific Reports</u> **12**, 2294 (2022)



We find the fastest convergence for spin/charge susceptibility when $\lambda/E_F \sim I$



Hund's coupling is very large, because Yukawa screening reduces F₀, but not much F₂, F₄.

Local point of view converging much faster than long-range point of view.

Short range correlations point of view

In realistic solids (ab-initio) we do not have yet such calculations that would add corrections in a controlled systematic way (through counter-terms), nor we have a way to estimate error of such local approximation. Nevertheless, the local DMFT approximation is the first step in this directions, and is already very successful in numerous solids. It allows **high-throughput calculation** of physical properties.

More restrictive than short range interaction, but very good starting point when screening makes interaction short range.

DMFT approximates:

 Φ all local Feynman diagrams (in fully dressed perturbation theory)

 $\Phi[\{G_{ii}\}]$

Similarity with local density approximation:

LDA: $V_{xc}(\mathbf{r}\tau,\mathbf{r}'\tau') = \delta(\mathbf{r}-\mathbf{r}')\delta(\tau-\tau')V_{xc}(\mathbf{r})$

exact in the limit of constant density



$$\{G_{ij}\}] \approx \Phi[\{G_{ii}\}]$$

i is site, or cluster...

can be obtained by solving an auxiliary quantum impurity problem (A.Georges & G. Kotliar, 1992).

> DMFT: $\Sigma_{ij}(\omega) = \delta_{ij}\Sigma(\omega)$ exact in the limit of large connectivity (∞D).

Much less restrictive than LDA. Keeps entanglement between ion and environment