> The derivative discontinuity in transport: towards a density functional description of Coulomb blockade and Kondo effect

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Preface: (TD)DFT and strong correlations

- Can one ever hope to describe strongly correlated systems with (TD)DFT?
 Sure! Hohenberg-Kohn, Kohn-Sham, Runge-Gross not only apply to weakly correlated systems
- So what's wrong with our approximations? focus here: many popular functionals (LDA, GGA) don't have derivative discontinuity at integer particle numbers
- What quantities can we hope to get right with (TD)DFT? certainly the density and all quantities which can be expressed in terms of the density
 but beware: many physical quantities of the Kohn-Sham system (e.g., spectral functions) are not supposed to

reproduce those of the physical system

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Preface and Outline

Setting the stage Dynamical Coulomb blockade and the derivative discontinuity Kondo effect: what TDDFT has to say Summary



- Setting the stage
- Derivative discontinuity and time-dependent picture of Coulomb blockade
- The Kondo effect: what TDDFT has to say
- Summary

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Setting the stage

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Derivative discontinuity in static DFT

total energy as function of (fractional) particle number is a series of straight lines (Perdew et al, PRL 49, 1691 (1982))

E(N+ω)



derivative discontinuity

$$\Delta = I(N) - A(N)$$

I(N): ionization potential A(N): electron affinity

N : integer number of electrons

m

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Derivative discontinuity in static DFT (cont.)

for given external potential $v(\mathbf{r})$, extend HK ground state energy functional to non-integer particle numbers:

derivative discontinuity

$$\Delta = \lim_{\omega \to 0} \left(\frac{\delta E_v[n]}{\delta n(\mathbf{r})} \Big|_{N+\omega} - \frac{\delta E_v[n]}{\delta n(\mathbf{r})} \Big|_{N-\omega} \right) = \Delta_{KS} + \Delta_{xc}$$

KS discontinuity $\Delta_{KS} = \varepsilon_{LUMO} - \varepsilon_{HOMO}$

xc contribution to discontinuity:

$$\Delta_{xc} = \lim_{\omega \to 0} \left(\frac{\delta E_{xc}[n]}{\delta n(\mathbf{r})} \bigg|_{N+\omega} - \frac{\delta E_{xc}[n]}{\delta n(\mathbf{r})} \bigg|_{N-\omega} \right)$$

<u>note</u>: for traditional functionals (LDA, GGA): $\Delta_{xc} = 0 !!$

Derivative discontinuity in static DF1 TDDFT for transport (Static) DFT for the Hubbard model A simple impurity model for transport Landauer steady-state approach TDDFT correction to Landauer conductance

TDDFT for transport



TD Kohn-Sham equation for orbitals

$$[i\partial_t - \hat{H}(t)]\psi_k(t) = 0$$

Hamiltonian of extended system L-C-R, no direct hopping between left and right leads

$$\hat{H}(t) = \begin{pmatrix} H_{LL}(t) & H_{LC} & 0 \\ H_{CL} & H_{CC}(t) & H_{CR} \\ 0 & H_{RC} & H_{RR}(t) \end{pmatrix}$$

TDDFT for transport

downfolding of equation of motion for extended orbitals (in region L-C-R) onto equation for orbital projected onto central region only

Equation of motion for orbital projected on central region

$$[i\partial_t - \hat{H}_{CC}(t)]\psi_{k,C}(t) =$$

$$\int_0^t \mathrm{d}\bar{t} \, \Sigma_{emb}^R(t,\bar{t})\psi_{k,C}(\bar{t}) + \sum_\alpha H_{C\alpha}g_\alpha^R(t,0)\psi_{k,\alpha}(0)$$

where (retarded) embedding self energy Σ^R_{emb} and (retarded) Green function g^R_{α} for isolated lead α describe coupling to leads

details in:

S. Kurth, G. Stefanucci, C.-O. Almbladh, A. Rubio, E.K.U. Gross, PRB **72**, 035308 (2005)

(Static) DFT for the Hubbard model



Schönhammer et al, PRB (1995); N.A. Lima et al, PRL (2003): parametrize total energy per site based on exact, Bethe ansatz (BA), solution of uniform Hubbard model with density n:

$$e^{BA}(n,U) = -\frac{2|V|\zeta}{\pi}\sin\left(\frac{\pi n}{\zeta}\right)$$

with parameter $\zeta(U)$ depending on interaction strength U one can extract xc energy $e^{BA}_{xc}(n,U)$ from this parametrization

(Static) DFT for the Hubbard model

derivative discontinuity at n = 1

$$\Delta_{xc} = \lim_{\epsilon \to 0^+} \left[v_{xc}^{BALDA}(n=1+\epsilon) - v_{xc}^{BALDA}(n=1-\epsilon) \right]$$
$$= U - 4|V| \cos\left(\frac{\pi}{\zeta(U)}\right)$$

local approximation:

for non-uniform Hubbard models, i.e., non-constant on-site energies or even different interactions at each site: use $e_{xc}^{BA}(n_i, U_i)$ as xc energy at site i (Bethe ansatz LDA, BALDA)

adiabatic approximation:

time-dependence of TDDFT xc potential at site i through $v_{xc}(i,t) = v_{xc}^{BALDA}(n_i(t))$ (C. Verdozzi, PRL (2008))



one interacting impurity, Hubbard-like on-site interaction U, non-interacting leads, hopping V in leads and hopping $V_{\rm Link}$ from leads to impurity, on-site energy ε_0 at impurity

ε,

interested in case of weak links $|V_{\rm Link}| < |V| \longrightarrow$ use $U/V_{\rm Link}$ as parameter in BALDA \longrightarrow modified discontinuity at impurity

$$\Delta_{xc} = U - 4|V_{\text{Link}}|\cos\left(\frac{\pi}{\zeta}\right)$$

Self-consistency condition for steady state density

Landauer approach:

assume for biased system there exists steady state with density n at impurity \longrightarrow self-consistency condition for n

$$n = 2 \sum_{\alpha = L,R} \int_{-\infty}^{\infty} \frac{\mathrm{d}\omega}{2\pi} f_{\beta}(\omega - W_{\alpha}) \Gamma(\omega - W_{\alpha}) |G(\omega)|^{2}$$
$$G(\omega) = [\omega - v_{KS}(n) - \Sigma_{L}(\omega - W_{L}) - \Sigma_{R}(\omega - W_{R})]^{-1}$$
$$v_{KS}(n) = \varepsilon_{0} + \frac{1}{2} Un + v_{xc}^{BALDA}(n)$$

 W_{α} : bias in lead α $f_{\beta}(\omega)$: Fermi function at inverse temperature β $\Sigma_{\alpha}(\omega)$: embedding self energy for lead α

Derivative discontinuity in static DFT TDDFT for transport (Static) DFT for the Hubbard model A simple impurity model for transport Landauer steady-state approach TDDFT correction to Landauer conductance

Landauer formula for steady state current

$$I = \int_{-\infty}^{\infty} \frac{d\omega}{2\pi} T(\omega) \left[f_{\beta}(\omega - W_L) - f_{\beta}(\omega - W_R) \right]$$

with transmission function

$$T(\omega) = \operatorname{Tr} \left\{ \Gamma_L(\omega - W_L) G(\omega) \Gamma_R(\omega - W_R) G(\omega) \right\}$$

 \longrightarrow zero-bias conductance:

$$G/G_0 = -\int_{-\infty}^{\infty} \frac{\mathrm{d}\omega}{2\pi} T(\omega) \frac{\partial f_\beta(\omega)}{\partial \omega}$$

Landauer formalism originally derived for non-interacting electrons; often used together with *static* DFT for conductance calculations but: static DFT is equilibrium theory, conductance is nonequilibrium property!!

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TDDFT correction to Landauer conductance

TDDFT is a proper non-equilibrium theory; steady state (if achieved) is the long-time limit of a time evolution process where system initially in equilibrium is perturbed by time-dependent bias

$$I = \int_{-\infty}^{\infty} \frac{\mathrm{d}\omega}{2\pi} T(\omega) \left[f_{\beta}(\omega - \Delta v_L^{KS}) - f_{\beta}(\omega - \Delta v_R^{KS}) \right]$$
$$\Delta v_{\alpha}^{KS} = v_{\alpha}^{KS}(t = \infty) - v_{\alpha}^{KS}(t = 0)$$

 \rightarrow zero-bias conductance in TDDFT:

$$G/G_0 = -(1+Q_{xc})\int_{-\infty}^{\infty} \frac{\mathrm{d}\omega}{2\pi}T(\omega)\frac{\partial f_{\beta}(\omega)}{\partial\omega}$$

with dynamical xc corrections Q_{xc}

Steady state picture for discontinuous potential Time-dependent transport: dynamical Coulomb blockade

Dynamical Coulomb blockade and the derivative discontinuity

Collaborators:

- G. Stefanucci, Univ. Tor Vergata, Rome, Italy
- E. Khosravi and E.K.U. Gross, MPI Halle, Germany
- C. Verdozzi, Univ. Lund, Sweden

Ref: S. Kurth, G. Stefanucci, E. Khosravi, C. Verdozzi, E.K.U. Gross, PRL **104**, 236801 (2010); see also: C.A. Ullrich, Physics **3**, 47 (2010)

Steady state picture for discontinuous potential Time-dependent transport: dynamical Coulomb blockade

Steady state self-consistent density for impurity model

l.h.s. and r.h.s. of self-consistency condition for \boldsymbol{n}



no solution for steady state density for some values of the bias.

to understand physics of this regime \longrightarrow smoothen xc discontinuity

Steady state picture for discontinuous potential Time-dependent transport: dynamical Coulomb blockade

Ground state densities in BALDA and QMC

compare BALDA and QMC ground state densities of impurity model as function of the on-site energies ε_0 for different values of the interaction U; $V_{\rm Link} = 0.18$



Steady state picture for discontinuous potential Time-dependent transport: dynamical Coulomb blockade

Steady-state density vs. bias for different $V_{\rm Link}$



BALDA:

step structure for small V_{Link} width of step: U \rightarrow Coulomb blockade <u>Hartree:</u>

no step structure \rightarrow crucial role of discontinuity

<u>note</u>: the role of the discontinuity in steady-state transport has also been discussed in C. Toher et al, PRL 95, 146402 (2005)

Steady state picture for discontinuous potential Time-dependent transport: dynamical Coulomb blockade

Time-dependent density in presence of discontinuity

Fermi energy $\varepsilon_F = 1.5|V|$, on-site energy $\varepsilon_0 = 2|V|$, right bias $W_R = 0$, interaction U = 2|V|, $V_{\text{Link}} = 0.3V$



for bias in step region of steady-state picture: no steady state; evolution towards a dynamic state of oscillating density around integer electron number

Steady state picture for discontinuous potential Time-dependent transport: dynamical Coulomb blockade

Time-dependent KS potentials and currents



in CB region: KS potential rapidly varying; train of rectangular steps; currents: sawtooth-like at impurity;

Finite temperatures: same model, different functional Densities and conductances

Kondo effect: what TDDFT has to say

Collaborator:

- G. Stefanucci, Univ. Tor Vergata, Rome, Italy
- Ref: G. Stefanucci, S. Kurth, PRL 107, 216401 (2011)

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Finite temperatures: same model, different functional Densities and conductances

Single-site model to construct finite temperature functional



4 states in Fock space: $|0\rangle$, $|\uparrow\rangle$, $|\downarrow\rangle$, $|\uparrow\downarrow\rangle$

calculate density $n(v_0 - \mu) = n(\tilde{v}_0)$ invert analytically $\longrightarrow \tilde{v}_0(n)$

non-interacting KS system: density $n_s(\tilde{v}_s)$ invert analytically $\longrightarrow \tilde{v}_s(n_s)$

Hartree-xc potential:

$$v_{Hxc}(n) = \tilde{v}_s(n) - \tilde{v}_0(n)$$

Finite temperatures: same model, different functional Densities and conductances

Hartree-xc potential for single-site model

Hartree-xc potential for different temperatures (in units of U)



derivative discontinuity emerges naturally in the zero-temperature limit

Finite temperatures: same model, different functional Densities and conductances

Density at zero temperature compared to exact results

local approximation: use single-site potential on impurity, vanishing KS potential in leads, should be reasonable approximation for weak coupling



Finite temperatures: same model, different functional Densities and conductances

Finite temperature conductance with Landauer



NRG data from: Izumida, Sakai, J. Phys. Soc. Jpn. (2005)

Finite temperatures: same model, different functional Densities and conductances

Finite temperature conductance with Landauer

two ways to understand T = 0 result:

• Meir-Wingreen formula for conductance:

$$\frac{G}{G_0} = \gamma^2 |\mathcal{G}(\mu)|^2 \frac{\gamma - \operatorname{Im} \Sigma(\mu)}{\gamma}$$

with many-body GF $\mathcal{G}(\omega)$ at impurity and self energy Σ at Fermi energy: $\mathrm{Im}\,\Sigma(\mu)=0\longrightarrow$ can describe conductance by a KS potential $v_s=v_0+\mathrm{Re}\Sigma(\mu)$

• Friedel sum rule (Langreth):

conductance determined by density n_0 on dot: $G = G(n_0)$ if KS potential gives good density \longrightarrow good conductance

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Finite temperatures: same model, different functional Densities and conductances

Finite temperature conductance with Landauer

Warning: KS density of states is *always* a simple Lorentzian, i.e., it has nothing to do with the exact DOS of the model!

However: due to discontinuity, the peak of the Lorentzian is pinned to Fermi energy (like the many-body resonance for the exact DOS) leading to conductance G_0

Finite temperatures: same model, different functional Densities and conductances

Finite temperature conductance with Landauer

<u>note</u>: at particle-hole symmetric point $v_0 = -U/2$ our approximation gives *exact* KS potential $v_s = 0$ for all temperatures



for finite T: Landauer *does not* give correct conductance although static KS potential is exact!

NRG results from [1] T.A. Costi, PRL (2000) [2] Izumida et al, JPPS (2005)

 \longrightarrow explicit example for dynamical xc corrections of TDDFT

Finite temperatures: same model, different functional Densities and conductances



simultaneous works on DFT for Anderson model at zero temperature:

- G. Stefanucci, S. Kurth, PRL 107, 216401 (2011)
- Bergfield, Liu, Burke, Stafford, arXiv:1106.3104 (2011)
- Tröster, Schmitteckert, Evers, arXiv:1106.3669 (2011)

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- Derivative discontinuity in transport
- absence of steady state in CB regime; instead: TD picture of CB as dynamical state of charging and discharging of weakly coupled system
- conductance plateau as function of gate (Kondo) at T=0
- understand in terms of Meir-Wingreen formula and Friedel sum rule
- finite *T*: Landauer not enough; TDDFT dynamical xc corrections essential

If you want to describe strongly correlated systems, the derivative discontinuity is your friend!