

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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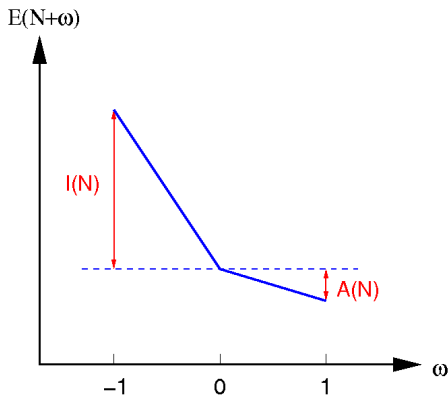
Outline

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

Setting the stage

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))



derivative discontinuity

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

$I(N)$: ionization potential

$A(N)$: electron affinity

N : integer number of electrons

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

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

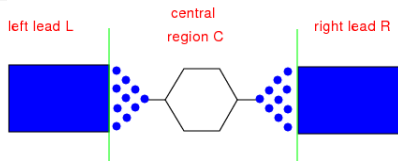
KS discontinuity $\Delta_{KS} = \varepsilon_{LUMO} - \varepsilon_{HOMO}$

xc contribution to discontinuity:

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

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

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

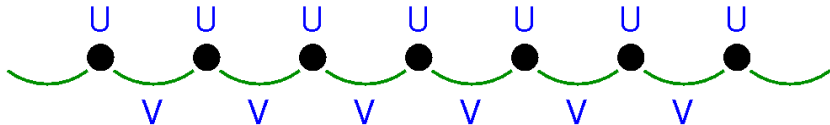
$$[i\partial_t - \hat{H}_{CC}(t)]\psi_{k,C}(t) = \int_0^t d\bar{t} \Sigma_{emb}^R(t, \bar{t})\psi_{k,C}(\bar{t}) + \sum_{\alpha} H_{C\alpha} g_{\alpha}^R(t, 0)\psi_{k,\alpha}(0)$$

where (retarded) embedding self energy Σ_{emb}^R and (retarded) Green function g_{α}^R 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 $e_{xc}^{BA}(n, U)$ from this parametrization

(Static) DFT for the Hubbard model

derivative discontinuity at $n = 1$

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

local approximation:

for non-uniform Hubbard models, i.e., non-constant on-site energies or even different interactions at each site:

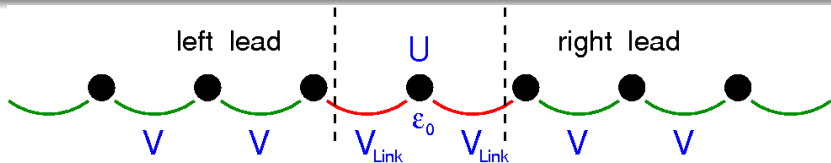
use $e_{xc}^{BA}(n_i, U_i)$ as xc energy at site i (Bethe ansatz LDA, BALDA)

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))

Simple impurity model for transport



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

interested in case of weak links $|V_{\text{Link}}| < |V| \rightarrow$ use U/V_{Link} as parameter in BALDA \rightarrow 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 \rightarrow self-consistency condition for n

$$n = 2 \sum_{\alpha=L,R} \int_{-\infty}^{\infty} \frac{d\omega}{2\pi} f_{\beta}(\omega - W_{\alpha}) \Gamma(\omega - W_{\alpha}) |G(\omega)|^2$$

$$G(\omega) = [\omega - v_{KS}(n) - \Sigma_L(\omega - W_L) - \Sigma_R(\omega - W_R)]^{-1}$$

$$v_{KS}(n) = \varepsilon_0 + \frac{1}{2}Un + v_{xc}^{BALDA}(n)$$

W_{α} : bias in lead α

$f_{\beta}(\omega)$: Fermi function at inverse temperature β

$\Sigma_{\alpha}(\omega)$: embedding self energy for lead α

Landauer formula for steady state current

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

with transmission function

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

→ zero-bias conductance:

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

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!!

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

$$I = \int_{-\infty}^{\infty} \frac{d\omega}{2\pi} T(\omega) [f_{\beta}(\omega - \Delta v_L^{KS}) - f_{\beta}(\omega - \Delta v_R^{KS})]$$

$$\Delta v_{\alpha}^{KS} = v_{\alpha}^{KS}(t = \infty) - v_{\alpha}^{KS}(t = 0)$$

→ zero-bias conductance in TDDFT:

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

with dynamical xc corrections Q_{xc}

Dynamical Coulomb blockade and the derivative discontinuity

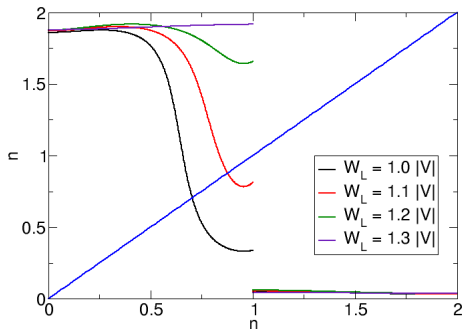
Collaborators:

- 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 self-consistent density for impurity model

l.h.s. and r.h.s. of self-consistency condition for n

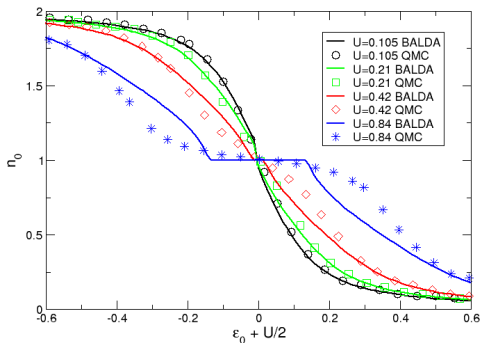


no solution for steady state density for some values of the bias.

to understand physics of this regime \longrightarrow smoothen xc discontinuity

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 ; $V_{\text{Link}} = 0.18$



QMC results from:

X. Wang et al,
PRB **77**, 045119 (2008)

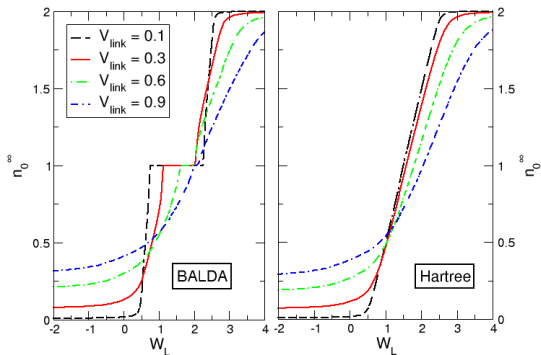
small U :

very good agreement

large U :

reasonable agreement +
Coulomb blockade step

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



BALDA:

step structure

for small V_{Link}

width of step: U

→ Coulomb blockade

Hartree:

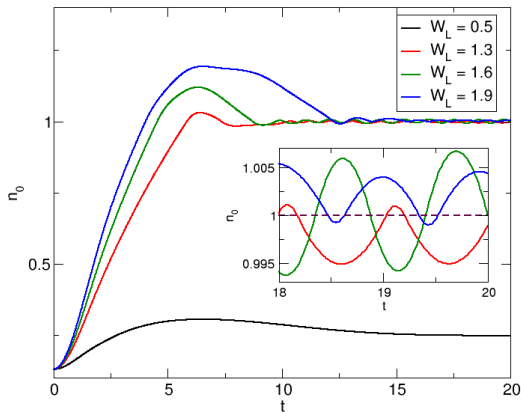
no step structure

→ crucial role of
discontinuity

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

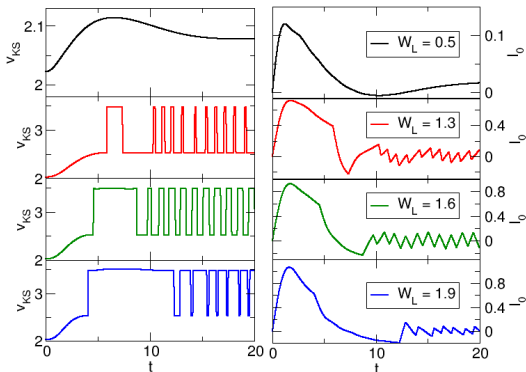
Time-dependent density in presence of discontinuity

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

Time-dependent KS potentials and currents



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

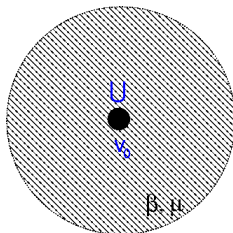
Kondo effect: what TDDFT has to say

Collaborator:

- G. Stefanucci, Univ. Tor Vergata, Rome, Italy

Ref: G. Stefanucci, S. Kurth, PRL **107**, 216401 (2011)

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 $n_s(\tilde{v}_s)$

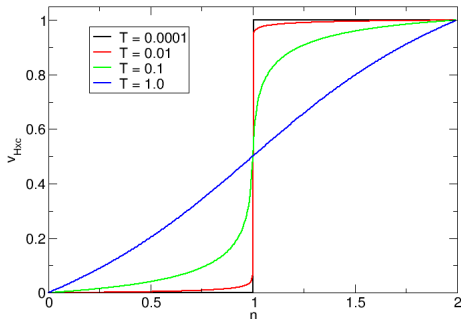
invert analytically $\longrightarrow \tilde{v}_s(n_s)$

Hartree-xc potential:

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

Hartree-xc potential for single-site model

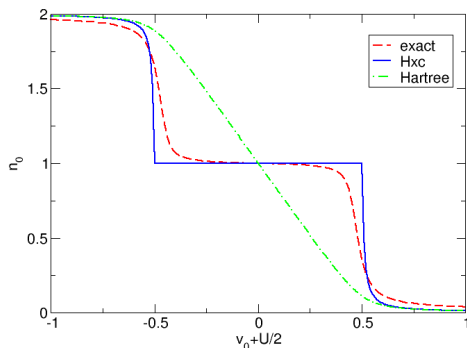
Hartree-xc potential for different temperatures (in units of U)



derivative discontinuity
emerges naturally in the
zero-temperature limit

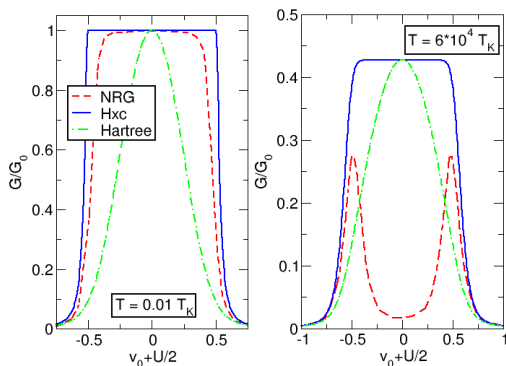
Density at zero temperature compared to exact results

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



$U/\gamma = 100$; γ : DOS
in wide-band limit;
density pinned to 1
due to discontinuity;

Finite temperature conductance with Landauer



$T \ll T_K$: Kondo
plateau in conductance
due to discontinuity;

$T \gg T_K$: plateau *not*
destroyed; no Coulomb
blockade peaks

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

Finite temperature conductance with Landauer

two ways to understand $T = 0$ result:

- Meir-Wingreen formula for conductance:

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

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

- 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

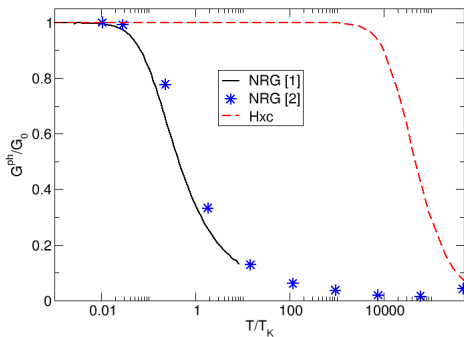
Finite temperature conductance with Landauer

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 temperature conductance with Landauer

note: 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)

→ explicit example for dynamical xc corrections of TDDFT

Related works

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)

Summary

- 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!