

SPIN TRANSFER TORQUES IN INTERACTING WIRES

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Use of a spin polarized current for the manipulation of magnetic domain walls in ferro- magnetic nanowires has been the subject of intensive research for many years. While so far the theoretical investigation has been mostly concerned with the mean-field approximation of the electron-electron correlation, the electrons confined to a quasi-one-dimensional wire be- have fundamentally different from standard Fermi-liquid picture. Our goal is to delve into the full quantum investigation of the electron-electron interaction in low dimensions. To this end, first we establish a reference point by solving the scattering problem of a noninteracting tight-binding Hamiltonian including an s-d interaction that couples the domain wall and the itinerant electrons. With this approach the behaviour of the adiabatic spin transfer torque has been investigated and the existence of a topological torque has been established. Next, we make an inquiry into the full quantum effects of electron-electron correlation on spin densities and consequently on spin transport.

Motivation

One-dimensional (1D) systems (Nanowires):

- Fermi liquid theory breaks down
→ Luttinger liquid theory
- Dominance of e-e correlations
- Pronounced effect of impurities
- DWs are extended magnetic impurities

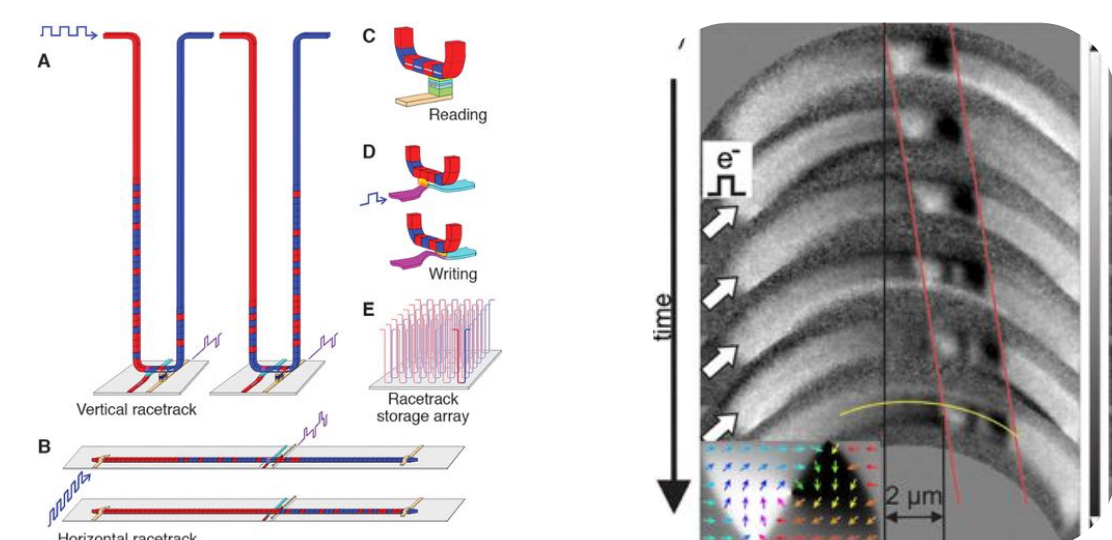
Relevance to applications:

- All electronic (without moving parts), faster, miniaturizable and more energy efficient DW based racetrack memories

Drawbacks?

- High polarized current density needed to move DWs.
- Better understanding of correlations and non-adiabatic spin-torque required

Current induced DW motion and DW based racetrack memories



Left: Proposed racetrack memory: a ferromagnetic nanowire, with data encoded as a pattern of magnetic domains [1].

Right: X-ray photoemission electron microscopy image series of a Py wire. Current pulses were injected. The DW is displaced about 400 nm on average per current pulse [2].

Model and Hamiltonian

Interplay between the electrons and the DW is described by spin-spin coupling in terms of s-d model [3].

$$\mathcal{H} = \sum_{j,\sigma} \left[-t \left(c_{j,\sigma}^\dagger c_{j+1,\sigma} + \text{h.c.} \right) + \mu c_{j,\sigma}^\dagger c_{j,\sigma} - \frac{J_{sd}}{2M_s} \sum_{\sigma'} \left(c_{j,\sigma}^\dagger \sigma_{\sigma,\sigma'} \cdot \mathbf{M}(x_j) c_{j,\sigma'} \right) + U n_{j,\sigma} n_{j+1,\sigma} \right]$$

$c_{j,\sigma}^\dagger$ is the creation operator of a spin σ electron at site j ; t is the hopping, J_{sd} is the s-d exchange interaction and U represents the electron-electron interaction. M_s is the saturation magnetization.

Spin Torques

Different types of torques in DWs and Multilayers (ML)			
Torque	Medium	Due to	Limit
Adiabatic STT [4, 5]	DW, ML	Gradual spin-flip	Adiabatic
Non-adiabatic (Mom. transfer) [6]	DW, ML	Reflection + exchange term	Sharp, interface
Non-adiabatic, due to spin flip relaxation (β term) [7]	Collinear structures	Spin-flip relaxation ($\delta m / \tau_{sf}$)	Interface
Gauge Torque [6, 8]	DW	DW width and misalignment from effective field	Sharp

Torque:

$$\boldsymbol{\tau} = \mathbf{B}_{\text{ex}}(x) \times \mathbf{m}(x) \quad \mathbf{B}_{\text{ex}}(x) = \frac{S J_{sd}}{M_s} \mathbf{M}(x)$$

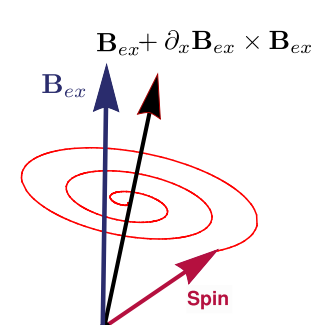
Adiabatic STT: In the plane of incoming and outgoing electron's spin direction.

Gauge STT: Due to the procession of spins around the effective field[9]

$$\mathbf{B}_{\text{eff}}(x) = \mathbf{B}_{\text{ex}}(x) + \partial_x \mathbf{B}_{\text{ex}}(x) \times \mathbf{B}_{\text{ex}}(x)$$

Nonadiabatic STT due to relaxation: Perpendicular to the plane of incoming and outgoing electron's spin direction.

Procession of spin around the effective field, i.e. the sum of the exchange and gauge field.



Collinear magnetization [7]

(Gradient STT: $c_1 = \text{const}$; Zhang-Li STT: $c_1 \propto$ current density):

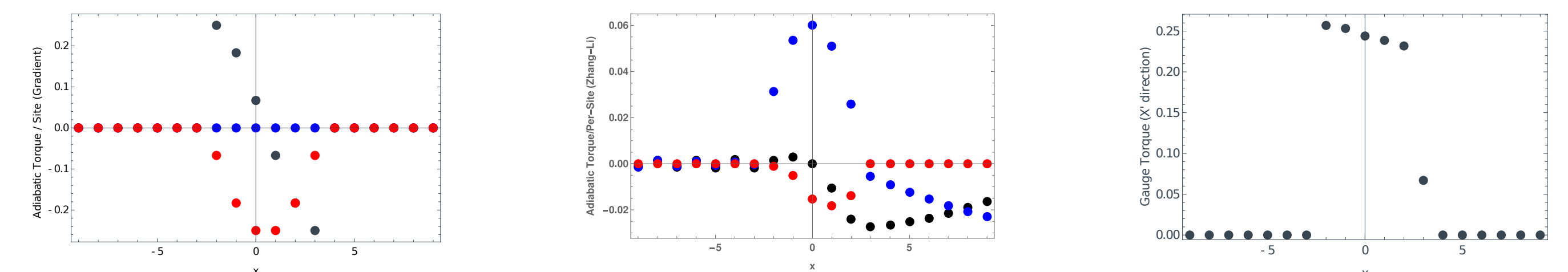
$$\tau(x) = \underline{c_1 \partial_x \mathbf{M}(x)} + c_2 \mathbf{M}(x) \times \partial_x \mathbf{M}(x)$$

Non-collinear magnetization [8]:

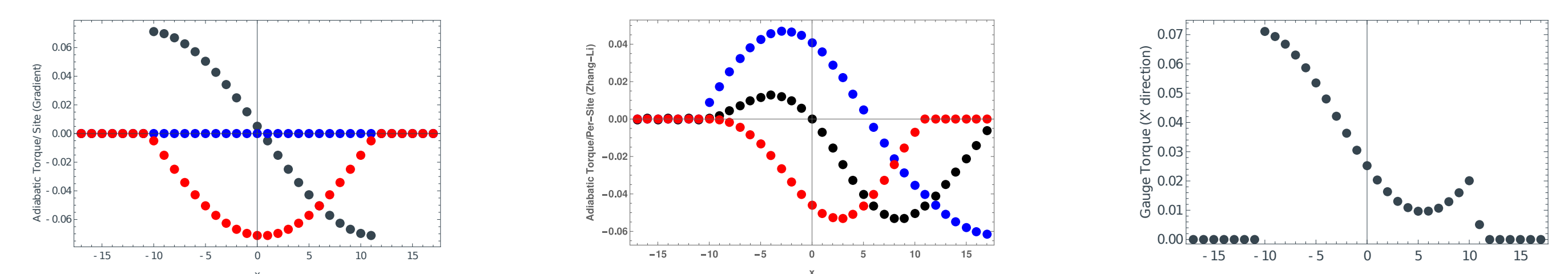
$$\boldsymbol{\tau}(x) = \underline{c_1 \partial_x \mathbf{M}(x)} + \underline{a(x) \hat{\mathbf{x}}'} + b(x) \hat{\mathbf{y}}$$

DW Torques in Non-Interacting Wires

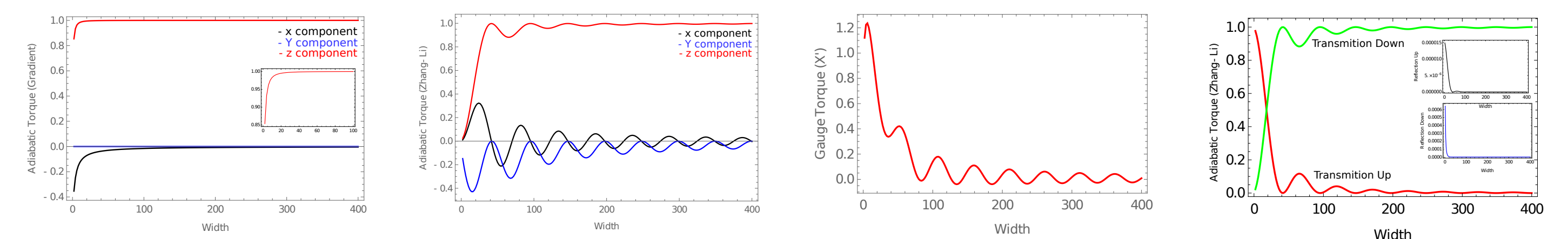
Diagonalizing the noninteracting Hamiltonian ($U = 0$), we investigate the resulting torques on the local magnetic moments.



STT in a DW of width = 5 (sharp limit). Polarization of incoming electron: \uparrow and wave vector of O (lattice site).



STT in a DW of width = 21 (adiabatic limit). Polarization of incoming electron: \uparrow and wave vector of O (lattice site).



These results will be used as a reference point for our future studies into the interacting case.

Luttinger Liquid Theory

The model can be bosonized[10, 3]

The diagonal Hamiltonian:

$$H = H_q + H_{\text{Gsf}}^f + H_{\text{Gsf}}^b + H_{\text{Gp}}^b$$

$$H = \frac{1}{2} \int dx \sum_{i=1,2} \left(\frac{u_i}{g_i} (\partial_x \phi_i)^2 + u_i g_i (\partial_x \tilde{\phi}_i)^2 \right)$$

Spin and charge modes are related to the diagonal modes by a transformation and we use the frame of the local magnetization direction

N.B. this is no longer spin-charge separated!

Scattering terms from the domain wall ($\Theta(x)$ controls the DW profile):

$$\begin{aligned} H_{\text{Gsf}}^f &= - \int dx \frac{k_F + t}{\pi a} \Theta'(x) \sin[\sqrt{2\pi} \tilde{\phi}_s(x)] \cos[k_F x + \sqrt{2\pi} \phi_s(x)] \\ H_{\text{Gsf}}^b &= - \int dx \frac{k_F - t}{\pi a} \Theta'(x) \cos[\sqrt{2\pi} \tilde{\phi}_s(x)] \sin[k_F x + \sqrt{2\pi} \phi_s(x)] \\ H_{\text{Gp}}^b &= - \int dx \frac{t}{4\pi a} \sum_{\sigma} [\Theta'(x)]^2 \sin[2k_F \sigma x + \sqrt{4\pi} \phi_{\sigma}(x)] \end{aligned}$$

We can calculate all spin densities, currents, etc., and the low energy phase diagram[3]:

Backscattering relevance for RG	$\lambda \lesssim \lambda_{\pm}$	$\lambda_+ < \lambda < \lambda_-$	$\lambda_{\pm} < \lambda$
$2 - \gamma_b > 0$	Spin and charge insulator	C1S0	Adiabatic LL
$2 - \gamma_b < 0$	C1S0	C1S0	Adiabatic LL

Outlook

- Non-adiabatic STT due to relaxation can be calculated if we introduce a damping mechanism
- The electrons confined to a 1D wire behave fundamentally different from standard Fermi-liquid quasiparticle picture, so a characteristic change in the spin transfer torque is expected
- To take into account electron-electron correlations, DMRG and bosonization will be used
- DMRG is nearly exact at almost any temperature but is limited by system size
- Field theory provides almost exact analytical results for thermodynamic limit and low T
- Numerical studies will be complemented by bosonization and Luttinger liquid theory
- For more realistic DWs, geometry and system parameters from experimentalists is needed

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