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Current distribution in metallic multilayers from resistance measurements

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The in-plane current profile within multilayers of generic structure Ta/Pt/(CoNi)/Pt/Ta is investigated. A large set of samples where the thickness of each layer is systematically varied was grown, followed by the measurement of the sheet resistance of each sample. The data are analyzed by a series of increasingly elaborate models, from a macroscopic engineering approach to mesoscopic transport theory. Non-negligible variations of the estimated repartition of current between the layers are found. The importance of having additional structural data is highlighted.

I. INTRODUCTION

From its very beginnings, spintronics is relying on metallic and semiconductor multilayers. The samples where giant magnetoresistance (GMR) was discovered were indeed (Fe/Cr) multilayers [1, 2], with current flowing in the layers’ plane (CIP geometry). The spin-transfer torque (STT) effect [3, 4], when used in the CIP geometry to drive magnetic domain walls [5–8], also involves metallic or semiconductor multilayers with magnetic and non-magnetic parts. In such samples, only the total injected current is known, and its distribution between the various layers is an open question: how much of the current is diverted by the buffer layer, or by the spacer layer (case of spin-valve nanostrips [9], or of synthetic antiferromagnetic nanostrips [10]) affects the computed efficiency of the STT-induced domain wall motion. The current in the non-magnetic layers also gives rise to a non-compensated Oersted field in the magnetic layer, which affects the structure and dynamics of the domain walls [11]. The spin-orbit torques (SOT), the latest family of current-induced torques in the CIP geometry [12–15], also occur in such multilayer samples. Here again, the distribution of the current between the various layers is of crucial importance.

Presently there exists no direct method to observe how current flows in the different layers of the multilayer sample, hence indirect methods must be used. The most employed one consists in using an effective conductivity for each layer, applying the parallel conductors rule of macroscopic electrical engineering to evaluate how much of the current flows in each layer. This effective conductivity is either assumed, or in some cases obtained by growing a series of samples with changing thickness of the layer(s) [15], again using the rules of macroscopic electrical engineering to obtain this conductivity.

However, the increase of the apparent resistivity of a metal film when its thickness decreases below the mean free path of electrons has been discussed as early as 1901, by J.J. Thomson [16]. The development of this idea has led to the so-called Fuchs-Sondheimer semi-classical model [17, 18], which has been further developed and refined. In this paper, from an extensive series of metallic magnetic multilayer samples where all thicknesses were varied, we quantitatively compare the samples conductivities with a number of models, with the goal of estimating the current distribution among the layers. The samples are emblematic of present studies of STT and SOT causing, for example, current-induced domain wall motion.

II. SAMPLES DESCRIPTION

The stack structure of the multilayer reference sample used in this work is Ta 3/ Pt 1.6/ [Co 0.3/ Ni 0.6]x4/ Co 0.3/ Pt 1.6/ Ta 3, where thicknesses are given in
nanometers. The stacks were deposited by dc magnetron sputtering onto a thermally oxidized Si substrate.

In order to estimate the current profile across the thickness, the thickness of each layer involved was varied around its reference value, the other layers having their reference thickness. This leads to a total of 42 samples, which were deposited consecutively with no change of the deposition parameters. The sheet resistance of each sample was measured, just after deposition, by the four-probe technique, at 5 locations across the wafer, and the average of these measurements was taken. The 5 values were extremely close (less than 1% difference), showing the uniformity of the film properties. The reference sample was therefore fabricated 5 times. Slightly different resistance values were measured for this sample (from 58.2 to 64.3 Ω/sq, with no trend in time), the average value being 61.2 Ω/sq and the standard deviation 2.3 Ω/sq, 3.7% of the mean value. These variations can only be attributed to slow fluctuations of the deposition conditions, the noise with respect to the trend for each thickness series being at most 1 Ω/sq. To remove this noise in the raw data, the resistances of each thickness series were multiplied by a series-specific factor such that the average value being 61.2 Ω/sq and the standard deviation showing the uniformity of the film properties. The reference thickness series results and assuming that no current flows in Ta is not enough to solve the problem, as this leads to $G_{Ta} = 23.9$ mSsq.

On the other hand, model 2 tries to fit globally all data, according to

$$G_{Ta} = \sum_i \sigma_i d_i. \quad (1)$$

The results of this model are shown in Fig. 1 by the dashed lines. Negative resistivities are obtained for the Ta underlayer thickness series (see Tab. I). This again proves that one cannot treat the electrical conduction in such multilayers made of nanometer-thick films by macroscopic engineering arguments, a fact known for a long time for single layers. Indeed, the well-known Fuchs-Sondheimer model [17–20] with surface scattering leads to a reduction of current density close to the surfaces, on a scale given by the electron mean free path $\lambda$ (for example, with no specular scattering at both surfaces, the apparent conductivity of a single layer is reduced by a factor 2 for a thickness $d = 0.46\lambda$).

IV. FUCHS-SONDHEIMER MODEL

As the samples are multilayers, we use the multilayer formalism described by Barnáš et al. [21] of the Fuchs-Sondheimer (FS) model.

The detailed derivation of the model can be found in the aforementioned publications, so here we present only the resultant expressions. The distribution function for electrons in a given layer can be written in the form:

$$f(z,v) = f_0(v) + g(z,v),$$

where $f_0(v)$ is the equilibrium distribution function and $g(z,v)$ corresponds to the contribution induced by the external electric field. The $z$-direction is perpendicular to the interfaces (Fig. 2). By solving the Boltzmann equation, one obtains the general expression for $g_\pm(z,v)$ ($g_+$ corresponding to the electrons moving in the positive $z$ direction, $g_-$ to the electrons moving in the negative $z$ direction), $d$ being the layer thickness:

$$g_+(z,v) = -\frac{eE\tau}{m} \frac{\partial f_0(v)}{\partial v_x} \left[ 1 - F_+(v) \exp \left( \frac{-z}{\tau |v_z|} \right) \right], \quad \text{and}$$

$$g_-(z,v) = -\frac{eE\tau}{m} \frac{\partial f_0(v)}{\partial v_x} \left[ 1 - F_-(v) \exp \left( \frac{-(d-z)}{\tau |v_z|} \right) \right],$$

where $F_\pm(v)$ are functions to be determined from the boundary conditions. The symbols $e$ ($e > 0$) and $m$ are the electron charge and effective mass, $\tau$ is the relaxation time, $v_x$ and $v_z$ the components of the velocity vector in $x$- and $z$-directions, respectively.
Figure 1: Measured sheet conductances (dots) vs. thickness of each layer (a-e), followed by the thickness-resolved conductivity plot for the reference sample, as derived from the models (f). The fit of each thickness series by a linear law (model 1) is represented by solid lines, whereas the global fit according to Eq. (1) (model 2) is drawn with dashed lines. No structural information is taken into account here, so that the models used are called 1a resp. 2a. The global rms error is 0.493 mS.sq when fitting each series independently, increasing to 1.621 mS.sq for the global fit. In order to compare the slopes, all panels (a-e) have the same scales.

Figure 2: Scheme of the sample structure with the various parameters of the Fuchs-Sondheimer model applied to analyze the conductivity data.

A. Direct application to the data

We impose the Fuchs boundary conditions at the bottom and top interface using coefficients $p_{\text{bot}}$ and $p_{\text{top}}$ corresponding to the specularity factors [19]. For the inner interfaces, similarly to Ref. 21, we introduce coefficients of specular transmission $T$ and reflection $R$ which differ for each interface (Fig. 2), but we neglect any angular dependence of these coefficients, an approximation at the heart of the FS model. We also neglect the refraction effects that may occur when the Fermi velocities in adjacent materials are different, and we assume the same transmission and reflection coefficients for electrons incident on the interface from the top and bottom. For simplicity, we also neglect any spin dependency of the coefficients, no magnetic field effects being investigated. For the (Co/Ni) multilayer, given the small thickness of the individual layers, the atomic proximity of Co and Ni, their good alloying properties, and the observed weak variation of resistivity with alloy concentration [22], it is treated as a single layer (alloy).

Application of these boundary conditions leads to a set of $N$ equations of $N$ variables, where $N$ is twice the number of layers in the sample (for every layer we have $F_{\pm}^\beta$). These equations are the spin-independent versions of the ones shown in Ref. [21]. We introduce $\beta$, the angle between the $z$ axis and the velocity vector $\mathbf{v}$, and $\lambda = v_F \tau$ the electron mean free path, $v_F$ being the Fermi velocity. Note that all mentioned parameters [$\lambda, v_F, \tau, m$] will generally differ for each layer. In correspondence with [21], we define $y_{AB} = \sigma_B / \sigma_A$, where $\sigma_A$ (\sigma_B) is the bulk conductivity of material A (B). These equations are solved numerically for each $\beta$ to obtain $F_{\pm}^\beta(\beta)$.

The final formula for the apparent conductivity of a
Figure 3: Fuchs-Sondheimer fit of all the data, with assumptions described in the text, and without any input from the structural characterization. The resulting rms error is 0.978 mS.sq (model 3a).

Figure 4: Fuchs-Sondheimer fit of all the data, taking into account a 2.0 nm oxidized Ta cap thickness. The corresponding rms error is 0.829 mS.sq (model 3b).

The single layer is:

\[
\sigma_i = \sigma_0,i - \frac{3}{4} \sigma_0,i \lambda_i \int_0^{\pi/2} d\beta \sin^3 \beta \cos \beta \left[ F_{i,+}(\beta) + F_{i,-}(\beta) \right] \left(1 - \exp \left(-\frac{d_i}{\lambda_i \cos \beta} \right) \right),
\]  

(4)
where $\sigma_{0,i}$ and $d_i$ are the bulk conductivity and thickness of layer number $i$, respectively. By Eq. (1), we obtain the sheet conductance of the multilayer sample, which is compared to the experimental results.

The total number of parameters entering the fit is 20 (Fig. 2), and it is further reduced. First, following Barnaś [21], we neglect reflection at the inner boundaries, the work functions being very close for all metal layers. Second, we fix the $\rho_0 \lambda_i$ product to the reference values for each layer [23, 24]. The values adopted for the $\rho_0 \lambda$ product (in $\Omega\cdot \text{m}^2$) were, from bottom (Ta underlayer) to top (Ta cap) 0.74, 2.35, 1.26, 2.35, and 6.0; see next section for justification of the choices for the Ta values. Additionally, upper boundaries for the mean-free path $\lambda$, equivalent to lower boundaries for the resistivity $\rho$ given by the best samples values, were set at 0.3, 22, 18, 22, and 10 nm, respectively. Third, we set $p_{\text{bot}} = p_{\text{op}} = 0$ on the outer boundaries, as they correspond to layers carrying little current so that the fit error depends weakly on them. After this reduction we are left with 9 parameters for all 42 samples. We fit the mean free paths of all the layers and transmission coefficients at all four inner interfaces.

The best fit of all data by the FS model (model 3) is shown in Fig. 3, together with the corresponding computed current profile across the thickness, for the reference sample. The rms error of this fit is 0.978 mS.sq; it is larger than that of the non-physical fit of each thickness series (model 1), but much lower than the error of the engineering-like independent layers model (model 2). The obtained parameters are given in Tab. I, and the current partition between the different layers as computed by the various models are compared in Tab. II.

B. Use of structural data

On close inspection, the acquired data appear strange at two instances. First, in the Ta cap series, one notices that the first three points show nearly the same conductivities. A cross-sectional transmission electron microscopy (TEM) image of the reference sample, shown in Fig. 5, reveals that the Ta cap layer is partly oxidized, with about 2 nm Ta consumed. Thus, the various models’ fits were performed again under this assumption (see Tab. I for the corresponding data). This variant of the models is indicated by the suffix ‘b’, suffix ‘a’ indicating that the models are applied to the data with nominal thicknesses. For the independent layers model, the rms error decreased to 0.423 mS.sq when fitting each series separately, whereas in the case of the global fit the relative decrease was smaller, the error reaching 1.555 mS.sq. For the FS model, as shown in Fig. 4, the quality of the fit improves markedly.

The TEM image also reveals that the crystalline structures of the two Ta layers are different, the underlayer showing no sign of crystallinity whereas the cap layer does. This is reflected in the fitted values of the FS model with the product $\rho_0 \lambda$ fixed according to the literature values for the two phases of tantalum [24].

Second, the platinum underlayer series, when compared to the FS fit, [Fig. 4(b)] shows a break for the three lowest thicknesses (0.4-1.2 nm). This is tentatively attributed to the formation of textured crystallites as the Pt underlayer starts growing on the amorphous Ta underlayer. These crystallites were directly imaged in TEM (Fig. 6) for the reference sample, giving a typical lateral size of 10 nm. The next section describes the modeling of the additional electron scattering at the grain boundaries.

V. EXTENDED FUCHS-SONDHEIMER MODEL

The FS model alone is not sufficient for the reproduction of experimental data. One reason for that is the polycrystalline structure of the layers in the sample, revealed e.g. by TEM imaging both in transverse cut (Fig. 5) and in plane view (Fig. 6). The boundaries between crystallites introduce an additional scattering of the electrons that is not covered by the FS model. This phenomenon was addressed by Mayadas and Shatzkes [25].

The FS model also does not account for roughness and thickness fluctuation of the layers, that naturally occur in the samples. Several models have been proposed to account for these phenomena; this will be studied in a second part.

A. Mayadas-Shatzkes model

The model by Mayadas et al. [25, 26] introduces grain boundary scattering into the FS model. For bulk samples or thick films, grain boundaries have little effect on the resistivity of metals as the grain size is much larger than the mean free path, however this is not the case for thin films, where the distance between grain boundaries is much smaller.

The modification of the FS model to account for the grain boundary scattering is simple to implement (for full derivation see Ref. [25]), the relaxation time being redefined to

$$\frac{1}{\tau^2} = \frac{1}{\tau} \left( 1 + \frac{\alpha v_F}{v_x} \right) = \frac{1}{\tau} \left( 1 + \frac{\alpha}{\sin \beta \cos \phi} \right) := \frac{M(\beta, \phi)}{\tau},$$

where $\phi$ is the angle between the $x$ axis and the projection of the velocity vector $v$ into the $xy$ plane. The parameter $\alpha$ carries the information about the grains and is defined as

$$\alpha = \frac{\lambda}{D (1 - R_{\text{grain}})},$$

with $D$ the typical grain diameter and $R_{\text{grain}}$ the grain boundary reflection coefficient.

The final formula for conductivity of a single polycrystalline layer thus reads
Figure 5: High magnification cross-sectional TEM image of the reference sample.

Figure 6: Dark-field TEM plane view of the reference sample, obtained by filtering out the (220) diffraction for Pt, Co and Ni (see diffraction pattern in inset).

\[
\sigma = \frac{3 \sigma_0}{\pi d} \int_0^{\pi/2} \int_0^{\pi/2} \sin^3 \beta \cos \beta \cos^2 \phi \cos \phi \cos^2 \beta \sin^3 \beta \left(2d - \lambda \cos \beta \frac{M(\beta, \phi)}{M(\beta, \phi)} \left[F_+(\beta, \phi) + F_-(\beta, \phi)\right] \left(1 - \exp \left(-\frac{d M(\beta, \phi)}{\lambda \cos \beta}\right)\right)\right)
\]

For the fits, the \(\rho\lambda\) products and zero reflectivities are kept as before to make results comparable, but \(R_{\text{grain}}\) is included as an additional fit parameter, resulting in 10 free parameters. The cross-sectional TEM image (Fig. 5) gives information on the grain structure. The Ta underlayer is found amorphous, thus no grains are considered in the model (technically, \(R_{\text{grain}} = 0\) is assumed for the Ta underlayer). For the Ta cap layer, results are weakly
Table I: Models used to fit the resistance data, their rms errors and parameters (layers are numbered from bottom to top, so 1 is the Ta underlayer etc.). Model 1 considers independent layers, fitting each thickness series separately. Model 2 also considers independent layers, but fits all thickness series together. Model 3 is the Fuchs-Sondheimer model. Model 4 is the Mayadas-Shatzkes model, with 2.0 nm oxidized Ta cap, the grain size varying in proportion to the fcc layers thickness (model 4), or to the Pt underlayer thickness (model 4*). Suffix ‘a’ indicates that the nominal thicknesses are considered, whereas suffix ‘b’ means that up to 2.0 nm of Ta cap was considered oxidized.

| Model | error (mS sq) | $\rho_1$ | $\rho_2$ | $\rho_3$ | $\rho_4$ | $\rho_5$ | $\lambda_1$ | $\lambda_2$ | $\lambda_3$ | $\lambda_4$ | $T_{12}$ | $T_{23}$ | $T_{34}$ | $T_{45}$ | $R_{\text{grain}}$ |
|-------|---------------|----------|----------|----------|----------|----------|-------------|-------------|-------------|-------------|---------|---------|---------|---------|-------------|
| 1a    | 0.49          | 82.6     | 34.3     | 27.1     | 32.9     | 79.3     | 0.26        | 10.3        | 5.00        | 14.4        | 7.95    | 0       | 0       | 0       | 0           |
| 1b    | 0.42          | 82.6     | 34.3     | 27.1     | 32.9     | 62.2     | 0.26        | 10.3        | 5.00        | 14.4        | 7.95    | 0       | 0       | 0       | 0           |
| 2a    | 1.62          | -8300    | 54.8     | 36.9     | 46.8     | 2590     | 0.3         | 11.3        | 5.40        | 12.7        | 10.0    | 0       | 0       | 0       | 0.78        |
| 2b    | 1.56          | -807     | 56.8     | 38.3     | 48.2     | 101      | 0.3         | 11.3        | 5.40        | 12.7        | 10.0    | 0       | 0       | 0       | 0.78        |
| 3a    | 0.98          | 287      | 22.7     | 25.2     | 16.4     | 75.5     | 0.3         | 7.6         | 7.0         | 19.9        | 10.0    | 0       | 0       | 0       | 0.97        |
| 3b    | 0.83          | 247      | 20.8     | 23.3     | 18.5     | 60.0     | 0.3         | 7.9         | 10.4        | 16.7        | 10.0    | 0.24    | 1.0     | 1.0     | 0.57        |
| 4b    | 0.71          | 247      | 31.1     | 18.0     | 11.8     | 60.0     | 0.3         | 7.6         | 7.0         | 19.9        | 10.0    | 0       | 0       | 0       | 0.97        |
| 4*b  | 0.67          | 247      | 29.7     | 12.1     | 14.1     | 60.0     | 0.3         | 7.9         | 10.4        | 16.7        | 10.0    | 0.24    | 1.0     | 1.0     | 0.57        |

Figure 7: Mayadas-Shatzkes fit, with assumptions described in the text, taking into account that 2.0 nm of the Ta cap were oxidized, the grain size varying in proportion of the Pt underlayer thickness (model 4b). The resulting rms error is 0.705 mS sq. The curve downwards bend at low platinum thickness in panel (b) directly depicts the assumption of model 4; it is absent in the other variant (model 4*, not shown).

sensitive to grains, and the cross-sectional TEM image does not reveal them. No grain effect is therefore assumed for this layer ($R_{\text{grain}} = 0$). For the two platinum and the (Co/Ni) layers, that all grow with an fcc (111) texture (Fig. 6), cylindrical grains are considered, with the same reflection coefficient $R_{\text{grain}}$.

In the absence of systematic TEM images, the variation of the size of the grains as a function of the thickness of these three layers has to be modeled. The following results are based on speculations about the grain sizes. A study having observed that platinum grains grow linearly with thickness below 50 nm [27], two models (at least) can be considered: the grain size is proportional to the Pt underlayer thickness (model 4), or to the total fcc stack thickness (model 4*). In both cases, the proportionality constant is such that, for the reference thickness, the grain size matches the observed value of 10 nm.

Figure 7 shows the results for model 4. The obtained
Table II: Current distribution in the sample calculated by the various models ($I_n$ is the proportion of the total current that flows in layer $n$, in percent). See Tab. I for the definition of models, their fitted parameters, and the residual error of each fit. Model 0 is the uniform distribution of current throughout the sample, the variant (model 0*) assuming that no current flows in the tantalum, because of its very high resistivity in the amorphous phase.

| Model | nominal thicknesses (models ‘a’) | w/ oxidized Ta cap (models ‘b’) |
|-------|----------------------------------|---------------------------------|
|       | $I_1$ $I_2$ $I_3$ $I_4$ $I_5$   | $I_1$ $I_2$ $I_3$ $I_4$ $I_5$ |
| 0     | 23 12 30 12 23                   | 23 14 35 14 9                   |
| 0*    | 0 23 55 23 0                     | 0 23 55 23 0                     |
| 1     | 12 15 46 16 12                   | 12 16 49 17 6                   |
| 2     | -0.2 17 62 20 0.7                | -2 17 60 20 6                   |
| 3     | 6 12 58 13 11                   | 7 12 61 14 6                   |
| 4     | 7 9 63 14 6                    | 7 14 56 16 7                   |
| 4*    |                                  |                                 |

The alternative grain size hypothesis (model 4*) delivers a slightly better fit, and redistributes the current more evenly among the fcc layers, due to the fact that the fitted internal specularity factors have reached unity. In addition, the grain reflection coefficient is found much larger. This sensitivity of the fitting might just signal the too large number of free parameters. Globally however, models 4 and 4* predict similar current partitions.

B. Thickness averaging

All the above modeling was assuming layers that are perfect in the $z$ direction, despite the common observation that some sample roughness exists and that the layer thicknesses fluctuate from place to place (see for example the cross-sectional TEM image in Fig. 5). In the literature, it has been proposed to account for this fact either by modifying the specularity parameter [28–30], or by averaging over a thickness distribution [31]. The latter model, considering only one-dimensional thickness fluctuations, is clearly oversimplified. Two-dimensional fluctuations in conductor networks have been deeply studied more recently, and an effective medium approximation (EMA) has been shown to describe well the regime beyond the percolation transition [32]. For averaging in two dimensions, the EMA formula is an implicit equation:

$$\langle \frac{\sigma - \sigma_{EMA}}{\sigma + \sigma_{EMX}} \rangle = 0,$$

where the brackets denote the averaging over the probability distribution $\rho(\sigma)d\sigma$.

This effect has been implemented for all models, assuming that the fcc grains, being slightly disoriented as proved by the diffraction image Fig. 6, can grow at different speeds with the largest speed when the (111) plane lies perfectly horizontal. To model this, a normal distribution of the growth rate was taken, with a standard deviation expressed as a percentage $p$ as variable parameter. This corresponds to recent detailed studies in magnetic ultrathin films of skyrmions morphology and pinning [33], and of domain wall dynamics [34]. Typical values of $p$ extracted from these works are $p = 1 - 3\%$. To compute the average in Eq. (8), for each data point, the model values for all the fcc thicknesses modified according to the same value of the normal probability law were used for integration, repeating the average calculation for 10 values of $\sigma_{EMA}$ close to the model value in order to interpolate the solution of the EMA equation.

It was found that the EMA refinement led to negligible fitting error improvements, and thus negligible changes of the fitting parameters, even if values of $p$ as large as 5 % were considered. The mathematical reason for this is simple: as in Eq. (8) the low conductances get a larger weight than the large conductances, the bias of the EMA averaging is towards lower conductances, with shifts that increase as the thickness variation is larger. This results in a systematic negative curvature of the conductance versus thickness curves, quite the opposite of the experimental trend.

VI. CONCLUSION

The schematic types of models for the current distribution are: uniform current through the conductive layers (model 0), constant conductivity for each layer, obtained from the corresponding thickness series (model 1), and Fuchs-Sondheimer global model (model 3). We have shown that increasingly better fits of the data are obtained by considering more and more elaborate models (designated by higher and higher model numbers), taking into account additional structural data.
The remarks emerging from this work are.

- Comparing model 1 to 3, we see that the former underestimates the current in the CoNi multilayer, overestimates it in the Pt layers, and largely overestimates it for the Ta layers. This was to be expected, due to mean-free path effects.
- Model 2 is too rarely employed, which is a pity as this model shows directly the danger of model 1.
- The Fuchs-Sondheimer model (model 3) provides results that are not bracketed by the simplest models (models 0 and 1), as might have been assumed. These models should therefore be taken as indicative only.
- The Mayadas-Shatzkes model (model 4) does improve the fit error, but does not modify much the current distribution.

On this basis, the simplest way to improve model 1 seems to perform a Fuchs-Sondheimer calculation with zero specularities coefficients at the interfaces, using reasonable values for the intrinsic parameters.

The work reported shows that collecting as much as possible structural data is desirable. From breaks of slope in a thickness series one might guess that another phenomenon is taking place, but it is always better to have an independent observation of it. The effect of including structural information is not negligible, compare for example the current in the Ta cap layer as deduced from models 3a and 3b.

Concerning non-uniformities, which are often invoked, we have found that for these samples at least they play a negligible role. This conclusion was reached using an analytical averaging formula, simpler that numerical models proposed previously [35]. This conclusion opposes the one reached some time ago by Hoffmann et al. [36], probably due to the better quality of the films prepared nowadays.

One feature of the data that remains unclear is the influence of the Ta underlayer thickness on the sample conductivity. The observed increase of sheet conductance with thickness of this layer is indeed too large, so that model 1 predicts a large percentage of current in that layer. One may suspect that this is a texture effect. The role of a Ta underlayer on the subsequent growth of Pt, followed by Co, has been indeed recently highlighted by measurements of the interfacial Dzyaloshinskii-Moriya interaction [37], an anti-symmetric exchange term arising at the Co/Pt interface, which is anticipated to depend strongly on the interface quality [38]. To test this hypothesis, TEM studies of the sample as the Ta underlayer thickness is varied would be decisive.

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