A 2D Simulation Methodology for Thermo-Magnetics Effects on Tunneling Mechanisms of Nano-scaled MOS Devices

by

Gabriela Alejandra Rodríguez Ruiz

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Advisors:

Dr. Librado Arturo Sarmiento Reyes
Dr. Edmundo Antonio Gutiérrez Domínguez

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Abstract

The development of a 2D numerical simulation methodology that accounts for thermal and magnetic effects on the gate tunneling current of nano-scaled MOSFETs, is the main goal of this thesis. The Schrödinger-Poisson coupled equation system is modified to account for the influence of a static magnetic field. The wavefunctions, which are the solution to the Schrödinger-Poisson coupled equation system, and the energy, are then obtained as a function of the magnetic field and temperature. Then, by considering open boundary conditions with the Perfectly Matched Layer-PML method and using the Tsu-Esaki direct tunneling model, the gate tunneling current under the influence of a magnetic field and temperature is calculated. By modifying the source files of the commercial GTS Framework device simulation tool to incorporate the new simulation methodology, the gate tunneling current is computed as a function of electrical bias, temperature and magnetic field. By doing so, and by sweeping the magnetic field from negative to positive values, it is found out that electrical charges tunneling, through the gate oxide, from the semiconductor to the gate terminal, are swept from left to right. Therefore, the proposed simulation methodology, accompanied with experimental results, is a very valuable tool to investigate non-homogeneous space distributed tunneling properties.
Resumen

El desarrollo de una metodología de simulación numérica bi-dimensional que tome en cuenta efectos térmicos y magnéticos sobre la corriente de tunelado de compuerta de MOSFETs nanométricos, es el principal objetivo de esta tesis. El sistema de ecuaciones acopladas de Schrödinger-Poisson se modificó para tomar en cuenta la influencia de un campo magnético estático. Las funciones de onda y las energías, las cuales son la solución del sistema de ecuaciones Schrödinger-Poisson, se obtienen como una función del campo magnético y de la temperatura. Considerando condiciones de frontera abiertas mediante el método *Perfectly Matched Layer* PML y utilizando el modelo de tunelado directo de Tsu-Esaki, la corriente de tunelado de compuerta bajo los efectos del campo magnético aplicado y la temperatura es calculada.

Mediante la incorporación de la nueva metodología de simulación dentro de los archivos fuente de la herramienta comercial de simulación GTS Framework, la corriente de tunelado de compuerta es calculada como una función de la polarización eléctrica, la temperatura y el campo magnético. Mediante este proceso, y haciendo un barrido del campo magnético de valores positivos a negativos, se encontró que el tunelado de cargas eléctricas a través del óxido, desde el semiconductor hasta la compuerta es barrido de derecha a izquierda (de lado de la fuente al lado del drenaje), lo que equivale a hacer un mapeo del espacio o un escaneo de las propiedades de tunelado. Debido a esto, la metodología de simulación propuesta, acompañada con resultados experimentales, es una herramienta muy valiosa para investigar las propiedades de tunelado en espacios distribuidos no homogéneamente.
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Acronyms

CMOS - Complementary Metal-Oxide-Semiconductor.
DIBL - Drain Induced Barried Lowering.
DT - Direct Tunneling.
FDM - Finite Difference Method.
FEM - Finite Element Method.
FET - Field Effect Transistor.
FVM - Finite Volume Method.
FN - Fowler-Nordheim.
GTS - General TCAD Solutions.
IC - Integrated Circuits.
INAOE - Instituto Nacional de Astrofísica, Óptica y Electrónica.
ITRS - International Technology Roadmap for Semiconductors.
MOSFET - Metal-Oxide-Semiconductor Field-Effect Transistor.
PML - Perfectly Matched Layer.
SCE - Short Channel Effects.
TAT - Trap Assisted Tunneling.
TC - Transmission Coefficient.
VSP - Vienna SChrödinger-Poisson.
WKB - Wenztsel-Kramers-Brillouin.
Chapter 1

Introduction

1.1 Motivation

For more than four decades, the semiconductor industry has been committed to reduce the dimensions of the MOSFET with the aim to obtain a better performance, lower cost and higher density integration of integrated circuits. Nowadays, the dimensions of the MOSFET have evolved from micrometer scales to nanometer scales, where for instance the channel length of a MOSFET is in the order of tens of nanometers and the oxide thickness is less than two nanometers [1]. According to the International Technology Roadmap for Semiconductors (ITRS 2012) in 2022 the oxide thickness will be less than 0.9 nm [2].

Other characteristic, that has been applied since many years, is the mechanical strain on nano-scaled MOSFETs. Mechanical strain has been applied into nano-scaled MOSFETs to enhance the average carrier mobility and current drive capability. However, such a mechanical stress is not uniformly distributed on the channel plane, which results in a non-homogeneous channel plane conductive properties (Figure 1.1 and Figure 1.2) [3, 4]. A pure electrical characterization technique for measuring the gate tunneling current $I_g$ only gives the average scalar value of the $I_g$ current at the gate contact, giving no information about the non-homogeneous space distributed conductive properties in nano-scaled MOSFET.

Recently, a new magnetic-electrical combined characterization technique for analyzing electrical and conductive properties of nano-scaled MOSFETs with ultra thin gate oxides
1.1. Motivation

Figure 1.1: Simulation of the electron density distribution of a 35-nm MOSFET. [4]

has been developed at INAOE [5]. This alternative technique is based on the application of a magnetic field $\vec{B}$, which allows the electron flow to be deflected through regions with different conductivities. Then, by measuring the average current, under different magnetic flux magnitudes and directions, the conductivity can be scanned or mapped through different regions of the MOSFET channel [6]. Unlike a pure electrical characterization technique, this new electro-magnetic technique gives more information about the local space distribution of conductive properties of the MOSFET and proves to be effective to detect inhomogeneous tunneling properties along the channel of a MOSFET; a phenomenon that cannot be explained by resorting to purely electrical test techniques.

As the alternative characterization technique makes use of an externally applied magnetic field, there is the need to incorporate the magnetic field variable in a simulation tool. However, until now, there is no simulation tool that allows to incorporate electromagnetic effects in nano-scaled MOSFET simulations. In fact, this is introduced in this work, namely the incorporation of the magnetic field variable is done in addition of the temperature, so thermo-magnetic simulations can be performed in nano-scaled MOSFETs.

In this thesis, a preliminary modeling approach for simulating the electrical behavior of a nano-scaled MOSFET under the influence of an external magnetic field and thermal variations is proposed. As a first developing stage, a 2D modeling approach based on the
1.2 Objectives

The main objective of this thesis is to develop a 2D simulation methodology for analyzing the gate tunneling current of nano-scaled MOSFETs under the influence of a static magnetic field. In order to achieve this general objective, the next particular objectives are proposed:

- Analyze, understand and modify the algorithm for the solution of the Schrödinger-Poisson system.
- Establish the appropriate boundary conditions.
- Incorporate the magnetic field into the model.
- Develop a methodology to compute the gate tunneling current of a MOSFET under the influence of a magnetic field.
- Incorporate the simulation methodology into the GTS Framework device simulator.

Schrödinger-Poisson system that incorporates the magnetic field and allow us to simulate the gate tunneling current of a MOSFET under the influence of a magnetic field and thermal variations is introduced.

Figure 1.2: Simulation of the stress distribution along the channel length L (a) and along the width W (b) directions [3]
1.3 Structure of the thesis

This thesis is organized as follows, chapter 2 introduces the theoretical fundamentals about gate tunneling current in nano-scaled MOSFETs and also shows a review of the state of the art of tunneling current simulation approaches. Chapter 3 presents the simulation methodology proposed in this thesis for simulating the gate tunneling current of nano-scaled MOSFETs under the influence of a magnetic field. Chapter 4 describes the experiments and results obtained with our simulator, and finally, in Chapter 5 the conclusions and future work are shown.
2.1 MOSFET scaling

The Metal-oxide-semiconductor Field-effect transistor (MOSFET) is one of the most used transistor in microelectronics industry, where more than 90% of all digital circuits are designed using the CMOS technology [7].

During the last four decades, the semiconductor industry has worked in the continuous reduction of MOSFET’s dimensions to nanometer scales. The main reasons for pursuing miniaturization are the increment of packing density, chip functionality, lower costs and better performance of transistors (increment of current drive capability and speed). Following the Moore’s law the number of transistors on IC doubles approximately every two years and according to the ITRS 2012 the channel length will be less than 9.0 nm and the oxide thickness will be less than 1.0 nm in 2022 [2]; (Figure 2.1).

Nevertheless, this scaling down process presents some important challenges to device design. The short channel effects (SCEs) and the gate tunneling current are some of the more important effects that reduce the performance of nano-scaled MOSFETs [9].

2.1.1 Short channel effects and gate tunneling current

- **Impact ionization.** The scaling down of MOSFETs dimensions produces high vertical and horizontal electric fields (> $10^6 V/cm$). The rising in electric fields happens because
the terminal voltages are not scaled down to maintain the speed of the MOSFET. Due to this reduction in size but not in voltage, short channel effects appear in nano-scaled MOSFETs. Due to the high intensity of the electric field near to the drain, the electrons have enough kinetic energy to hit a bounded electron and release it to the conduction band; which simultaneously produces an electron-hole pair. This process is called impact ionization and it produces electron-hole pairs in the silicon lattice causing even more ionization in the semiconductor. As a direct consequence, there is an increase in the drain-source current and substrate current (Figure 2.2) [9, 10].
• **Punch-Through and Drain Induced Barried Lowering (DIBL).** Due to the miniaturization of the channel length, the depletion region from drain can reach the source depletion region. The consequence of this, is the reduction of the barrier for electron injection and this effect is known as Punch-Through. In long channel devices, the semiconductor is depleted completely by the gate bias but in short channel devices, the semiconductor is depleted also by the source/drain bias. As the drain bias increases the depletion region is expanded and it can interact with the source/channel junction producing a lowering in the potential barrier (Figure 2.3 and Figure 2.4) [9].

![Figure 2.3: Punch-Through effect](image1)

![Figure 2.4: DIBL effect](image2)

• **Mobility reduction.** As the channel length is reduced, the lateral and vertical magnitude of electric fields are increased. As a result, the carrier velocity can be saturated and it produces a decrease in the charge mobility [10].
2.2 Gate leakage current in MOSFETs

- **Hot electron effect.** Due to the high electric field in nano-scaled MOSFET, electrons can acquire enough energy to pass over or tunneling the oxide barrier. Such electrons can become trapped in the oxide region, changing the threshold voltage of the device (Figure 2.5) [9].

  ![Figure 2.5: Hot electron effect](image)

- **Gate tunneling current.**

  In nano-scaled MOSFETs, quantum effects as channel energy quantization and gate tunneling current dominate the electronic properties of the transistor. Due to the scaling down of MOSFETs dimensions, the oxide thickness is thinner than 2 nm, which causes an important flow of current through the gate. The oxide thickness is too thin that electrons can tunnel through the oxide barrier to the gate electrode. This tunneling current is one of the key leakage currents that contributes most to device degradation, leads to excessive power dissipation and loss of on-current density. Because of that, the reduction of leakage current is the main issue in MOSFET down scaling (Figure 2.6) [7, 9, 10].

  ![Figure 2.6: Gate tunneling current](image)

2.2 Gate leakage current in MOSFETs

In nano-scaled MOSFETs, quantum effects as channel energy quantization and gate tunneling current dominate the electronic properties of the transistor [11]. Due to the aggressive scaling
down of the oxide thickness and the high electric fields produced by the high doping of channel, electrons can tunnel through the oxide barrier producing a high current flow, which results in a high power consumption in the off-state of the transistor. The gate tunneling current is one of the biggest limitations in the attempt to further reduce the size of transistors [12, 13].

This gate leakage current in MOSFETs is produced by different tunneling processes that are bias-dependent, and by thermal stress. The main three tunneling processes are the Fowler-Nordheim (FN), the Trap assisted tunneling (TAT) and the Direct tunneling (DT).

- **Fowler-Nordheim tunneling**

  In nMOS transistors, when a large positive voltage (> 4V) is applied to the gate, the metal side of the band diagram is lowered enough so the potential barrier becomes approximately triangular (Figure 2.7). Due to this triangular potential barrier, electrons can easily tunnel from the conduction band of the semiconductor to the conduction band of the oxide. This tunneling process through an approximately triangular potential barrier is called Fowler-Nordheim tunneling. The simple expression for calculating the FN density current tunneling is

  \[
  J_{FN} = \frac{q^2 m_0}{16\pi^2 \hbar m_{ox} \phi_b} F^2 e^{-\beta/F}.
  \]  

Here \(\hbar\) is the Planck’s reduced constant, \(m_{ox}\) is the electron effective mass in the insulator, \(m_0\) is the electron mass in free space, \(q\) is the electron charge, \(\phi_b\) is the energy barrier height at the semiconductor-oxide interface, \(F\) is the electric field across the oxide, and \(\beta = \frac{4(2m_{ox})^{1/2}}{3q\hbar} \phi_b^{3/2}\).

The Fowler-Nordheim tunneling formula is very accurate for modeling gate tunneling current of MOSFETs with oxide thickness in the order of 7-10 nm [14].

- **Trap assisted tunneling**

  Defects in the insulator produce electronic states in the band gap of the oxide and can split the energy barrier into two parts, (Figure 2.8). These states allow the charge tun-
2.2. Gate leakage current in MOSFETs

During tunneling through thinner energy barriers. Electrons have a higher probability of tunneling in oxides with traps than those that tunnel the entire oxide. The basic model for the current density assisted by traps $J_{TAT}$ is [15]

$$J_{TAT} = q \int \frac{N_T}{\tau_c + \tau_e} \delta x,$$

where $N_T$ is the trap density at a distance $x$ away from the emitting electrode, $\tau_c$ is the characteristic times for carrier capture and $\tau_e$ is the characteristic times for emission by the trap.

- **Direct tunneling**

  In ultra-thin oxide MOSFET (below 5nm), the strength of the electric field at the oxide is so high ($10^7 V/cm$) and the oxide is too thin (< 5nm) that electrons can tunnel in
2.2. Gate leakage current in MOSFETs

a direct way through the barrier of the oxide. In this tunneling process, electrons can transverse the entire width of the potential barrier, even with lower applied voltage, and for this case the oxide potential profile is trapezoidal, (Figure 2.9).

![Figure 2.9: Direct Tunneling (trapezoidal barrier)](image)

The tunneling probability increases dramatically as the oxide thickness is reduced [9]. This tunneling process is called direct tunneling and the electron flux from this tunneling process is the most dominant gate leakage process through oxides below 5nm [15], which is the case of a 28 nm CMOS transistor. Moreover, in order to model this effect, classical physics is not appropriate and quantum physics needs to be considered. The direct tunneling current can be described by the Tsu-Esaki formula [17],

\[
J = \int_{E_{\text{min}}}^{E_{\text{max}}} TC(E_i)S(E_i)dE_x,
\]

where \(S\) represents the number of electrons in an energy level, which are available for tunneling through the barrier potential function, the \(TC\) term is the transmission coefficient, \(E_x\) represents the electron kinetic energy in the direction perpendicular to the barrier.

Due to the impact of the direct tunneling current on the performance of nano-scaled MOSFETs with ultra-thin oxides, in this thesis we are particularly interested in modeling the direct tunneling (DT). In the next section a review of different modeling approaches for direct tunneling currents is introduced.
2.3 Literature Review of direct tunneling modeling

Since the scaling of gate oxide started in the 1970’s, a lot of gate tunneling modeling approaches have been developed. In the 70’s Harrison [16] and Tsu [17] proposed one of the first general direct tunneling current formula. The authors assume that the band structure varies slowly and the potential barrier has a triangular or trapezoidal shape. In order to calculate the transmission coefficient, they use the Wenztsel-Kramers-Brillouin (WKB) method [19]. The WKB method gives an approximation of the solution of the Schrödinger equation by means of an analytical formula, which depends on the width of the barrier. Such approximation method is accurate for triangular potential barriers. After that, Rana et al. [20] solved the self-consistent Schrödinger-Poisson system in one dimension for accumulation layers in MOS devices. The authors consider extended and quasi bound states for tunneling current. Moreover, they compare a semi-classical model with their model, obtaining almost the same current results. Then, Lo et al. [22] considered a tunneling current from the quantized states in the inversion layer of nano-scaled MOSFETs. The authors have calculated the lifetimes of quasi-bounded states and found that more than 80 percent of the total current comes from the lowest two energy states. The authors used the matrix-transfer method for calculating the transmission of the wavefunctions. The matrix-transfer method is used for calculating the transmission coefficient of a potential barrier. The method divides the potential barrier into a number of potential energy steps. By using a piecewise representation of the potential, the wavefunctions are approximated by constant values [23, 24]. In 1998, another work focused in the solution of the model in 1D was developed by Janick and Majkusiak [25]. These authors assumed a linear distribution of the potential (considering a system of decoupled equations Schrödinger-Poisson), rendering an analytical expression for the charge distribution. The Janick and Majkusiak’s work considers a confined triangular well in a closed system. Register et al. [27] introduced an analytical model for direct tunneling gate current in MOSFETs. The analytical expression is based on the WKB method and the electric field on the surface. This method is accurate only for low electric fields ($< 10^6\text{V/cm}$). Mundanai et al. [28] used a WKB approximation and obtained the tunneling current for different dielectrics materials and thicknesses. Lee et al. [30] used a modified WKB method
for calculating the direct tunneling current of MOS structures and investigated the effects on the direct tunneling current with different uniform doping concentrations in the substrate. In 2004, another important work was developed by Driskill [35]. This work computes the charge density and the tunneling current in a 1D quantum well. Moreover, Driskill assumes open boundary conditions, and calculates the tunneling current by means of the transfer matrix method and using finite difference method.

One of the first research groups, which solves the 2D Schrödinger-Poisson system, was the group of Spinelli et al [31] and the group of Trellakis [21]. In 2002, The authors solved a closed 2D Schrödinger-Poisson system using a finite element method. In order to calculate the current, they use a semiclassical approach, adopting a drift-diffusion model. After that, in 2003 the NANOTCAD2D simulator was developed by Curatola et al [32,33]. This simulator solves the 2D Schrödinger-Poisson system by means of the finite difference method. It is a robust simulator that the device can be defined as a closed system (two dimensions confined), semi-open system (one dimension confined and one dimension open), and an open system (all dimensions open); it also has models for ballistic current. Then, another complete work was the one developed by Polizzi and Abdallah [34]. The authors propose a methodology for solving in 1D and 2D the Schrödinger-Poisson system, considering either an open or a closed system. In such a work, the device is subdivided in slices of one dimension and the waves functions are calculated in each slice; at the end of the process, the waves functions are integrated to compute the global quantities, i.e., the charge density. This methodology solves the equations by means of the finite element method. Finally, there is another research group from the Technical University of Vienna, which has developed a simulator called Vienna Schrödinger Poisson Solver (VSP Solver) [37, 40, 46]. The VSP Solver is a robust and complete semiconductor device simulator with more than twenty models to choice, having particularly a special module that can solve the Schrödinger-Poisson system (1D and 2D) for both closed and open systems. The simulator is based on the volume finite method and allows the definition of a variety of structure devices and a wide variety for materials. Table 2.1 shows a review of gate tunneling simulation methodologies for nano-scaled MOS-FETs.
### Table 2.1: Comparison of gate tunneling simulation methodologies for nano-scaled MOSFETs.

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So far, all these are pure electrical models, but only a few simulators account for the effect of the magnetic field, some of them allows to analyze the transistor under the influence of an external magnetic field. There are many works like those described in [47–50, 52], which model a MOSFET under the influence of a magnetic field classically. All these works are based on the drift-diffusion model in which magnetic field dependent terms are added for accounting the action of the Lorentz’ force on the motion of the carriers. Popovic et al. proposed a methodology for calculating the drain current under the influence of a magnetic field. Also
this method is based on the drift-diffusion model and uses the finite element method. The authors assumed a lineal source-drain potential over the channel length. Nathan et al. also used a drift-diffusion model that includes magnetic field terms, but they assumed a constant drain source field over the channel length. They used the finite difference method as a discretization method. Riccobenne et al. ’s model took into account the hole and electron currents. The authors developed their model in a simulation tool called GENSIM and they used a box integration method for the discretization of the system. Rodríguez et al. developed a 2D/3D methodology based on the drift-diffusion approximation, which incorporates magnetic field terms. The authors developed their methodology in the MINIMOS-NT simulation tool and used the finite volume method as discretization scheme. Yosry et al., based on the Rodríguez’s work, developed a compact model (empirical model) for calculating the drain current under the influence of a magnetic field, which depends on geometrical properties. All these mentioned works calculate the drain current (but not the gate tunneling current) under the influence of a magnetic field. However, in order to model the new nano-scaled MOSFETs, models based in quantum mechanics needs to be accounted for.

Until now, there are few simulators that incorporate electromagnetic effects using quantum models, such is the case of Birner et al. [53]. The simulator created by them is called Nextnano and it allows to simulate quantum wires (2D considering a closed system), under the influence of a magnetic field. Although the simulator can compute the wavefunctions and the charge density under the influence of a magnetic field, but it cannot calculate the gate tunneling current. This simulator also is capable of modeling only the charge transport of a transistor without the presence of a magnetic field.

Another interesting work is the one developed by Sudiarta [57], in this work only the Schrödinger equation (not coupled with Poisson equation) is solved under the influence of a magnetic field but considering a closed system. A summary of the methodologies developed for modeling transistors under the influence of a magnetic field is shown in table 2.2.
Table 2.2: Comparison of methodologies for modeling MOSFETs under the influence of a magnetic field.

2.4 Discussion

In spite of the advantages and disadvantages that exhibit the device simulators mentioned above, until now there is no simulation tool that allows to analyze the gate tunneling current of a nano-scaled MOSFET under the influence of a magnetic field. As we mentioned in chapter one, with the emergence of the new electromagnetic characterization techniques, there is a necessity to incorporate thermo-magnetic effects in simulation tools that help us to analyze non uniform conductive properties in nano-scaled MOSFETs.

Taking into account all these properties, in this thesis, in collaboration with the Technical University of Vienna, a simulation methodology for solving the 2D Schrödinger-Poisson system that incorporates the magnetic field and allows to calculate the gate tunneling current of a nano-scaled MOSFET is developed.

In the next chapter the methodology simulation proposed in this work is described.
Chapter 3

Simulation methodology

3.1 Schrödinger-Poisson system

In nano-scaled MOSFETs various quantum effects become dominant over the device performance. For instance, gate-oxide tunneling and channel energy quantization are some quantum mechanical effects that impact the MOSFET performance. Calculation of these quantum effects in nano-scaled transistors are frequently based on the self consistent Schrödinger-Poisson system. [13]-[46].

The wavefunctions and the energies of the system are calculated by solving the time-independent Schrödinger equation,

\[
\begin{pmatrix}
\hat{p}^2
\end{pmatrix}_{2m^*} + V(\vec{r}) \psi(\vec{r}) = E \psi(\vec{r}),
\]

(3.1)

where \( \hat{p} = -i\hbar \nabla \) is the momentum operator, \( \psi \) is the wavefunction, \( E \) is the energy eigenvalue, \( V(\vec{r}) \) is the potential energy and \( m^* \) is the effective mass. The potential \( V(\vec{r}) \) is defined by

\[
V(\vec{r}) = -q\phi(\vec{r}) + \Delta E_{ce} \tag{3.2}
\]
where $\Delta E_{cr}$ is the energy potential profile generated by the bending of the oxide-substrate energy bands, $q$ is the electron charge and $\phi$ is the electrostatic potential.

The relation between the space charge and the electrostatic potential is calculated by means of Poisson equation,

$$\nabla^2 \phi(\vec{r}) = -\frac{q}{\varepsilon} (N_d(\vec{r}) - n(\vec{r})),$$

(3.3)

where $\varepsilon$ is the permittivity, $\phi$ is the electrostatic potential, $N_d(\vec{r})$ is the ionized donor concentration and $n(\vec{r})$ is the electron density distribution. The electron density $n(\vec{r})$ is determined from the solution of the Schrödinger equation and it is given by,

$$n(\vec{r}) = \frac{m^* kT}{2 \pi \hbar^2} \sum_i |\psi_i(\vec{r})|^2 \ln \left(1 + \exp\left(\frac{E_f - E_i}{kT}\right)\right),$$

(3.4)

where $k$ is the Boltzmann constant, $T$ is the temperature, $\hbar$ is the reduced Planck constant, $E_f$ is the fermi level and $E_i$ is the energy level associated to eigenfunction $\psi_i(\vec{r})$. In equation 3.4, the Fermi-Dirac distribution is assumed. This means, that all the particles into a system are considered identical with half-integer spin. Moreover, the Fermi-Dirac distribution allows to consider the many-particle system in terms of a single particle energy states, in which a null interaction between the particles of the system is considered.

A complete description of the charge transport of a nano-scaled MOSFET can be achieved by means of the self consistent solution of the Schrödinger and Poisson coupled equations. The Schrödinger-Poisson system is solved in a consistent way in order to find the potential distribution and the corresponding carrier concentration. The consistent solution of the system is found by solving both equations in a loop until the solution converges. Figure 3.1 shows the flowchart of the Schrödinger-Poisson system solution.
3.2 Schrödinger-Poisson system for a constant magnetic field

In the presence of a constant magnetic field $\vec{B}$, where $\vec{B}$ can be described by a vector potential $\vec{A}$ such that

$$\vec{B} = \nabla \times \vec{A},$$  \hspace{1cm} (3.5)
the kinetic potential term in the Hamiltonian operator is modified [56]. The quantum mechanical momentum operator \( \vec{p} \) becomes \( \vec{p} - q\vec{A}(\vec{r}) \) and the Schrödinger equation for a particle in a magnetic field can be defined as

\[
\left[ \frac{1}{2m} (\vec{p} - q\vec{A})^2 + V(\vec{r}) \right] \psi(\vec{r}) = E \psi(\vec{r}). \tag{3.6}
\]

Suppose we have a surface formed by the \( xy \) plane and a constant magnetic field along the \( z \) direction is applied (Figure 3.2), then a possible choice for \( \vec{A} \) is given by

\[
\vec{A} = (0, Bx, 0) \tag{3.7}
\]

![Figure 3.2: Constant magnetic field applied perpendicular to the surface of the \( xy \) plane.](image)

Any gradient of a scalar potential \( \theta(\vec{r}) \) can be added to \( \vec{A} \) without changing the value of the magnetic field \( \vec{B} \), since \( \nabla \times \nabla \theta(\vec{r}) \).

This invariance of the magnetic field for different choices of \( \theta(\vec{r}) \) is called gauge invariance. Another choice for a vector potential \( \vec{A} \) that generates the same constant magnetic field \( \vec{B} \) along the \( z \)-direction is given by

\[
\vec{A}(\vec{r}) = (-By, 0, 0) \tag{3.8}
\]
In order to incorporate the magnetic field into the Schrödinger equation we modified the general scheme of the solution of the self-consistent loop shown in figure 3.1. The module corresponding to the solution of the Schrödinger equation was changed for the Schrödinger equation that includes the magnetic field; see equation 3.6. The proposed scheme of the self-consistent loop that includes the magnetic field is shown in figure 3.3.

Figure 3.3: Self consistent solution flow chart of the Schrödinger-Poisson system, considering a magnetic field.
3.3 Gate tunneling current

Once the wavefunctions $\psi$ and the energies $E$ are computed, the direct tunneling current can be calculated. The direct tunneling current can be described by the Tsu-Esaki formula [17],

$$J = \int_{E_{\text{min}}}^{E_{\text{max}}} TC(E_x)S(E_x)dE_x,$$

(3.9)

where $S$ represents the supply function, $TC$ represents the transmission coefficient, $E_x$ represents the electron kinetic energy, $E_{\text{min}}$ represents the lower energy and $E_{\text{max}}$ represents the higher value of the discretized energies into the potential well. The supply function is given by

$$S = \ln \left( \frac{1 + \exp \left[ E_{fs} - E/k_B T \right]}{1 + \exp \left[ E_{fg} - E/k_B T \right]} \right),$$

(3.10)

where $E_{fg}$ and $E_{fs}$ are the electron Fermi levels in the gate and semiconductor, respectively, $E_x$ is the electron energy. Moreover, the supply function represents the number of electrons in an energy level, which are available for tunneling through the barrier potential [18].

The transmission coefficient $TC$ represents the probability of a particle tunneling through the barrier potential and it can be calculated as

$$TC = \frac{J_t}{J_i},$$

(3.11)

where the $J_t$ term corresponds to the transmitted current density and $J_i$ corresponds to the incident current density. The current density probability $J$ can be defined as

$$J(\vec{r}) = \frac{q}{2m^*} \left( \psi^* (\vec{p}\psi) - (\vec{p}\psi)^* \psi \right).$$

(3.12)
The simulation methodology proposed in this thesis, for calculating the gate tunneling current under the influence of a magnetic field, is shown in Figure 3.4.

Figure 3.4: Proposed simulation methodology for calculating the gate tunneling current of a nano-scaled MOSFET under the influence of a magnetic field.
3.4 GTS Framework

The Technical University of Vienna in collaboration with Global TCAD Solution Company (http://www.globaltcad.com/en/products/gts-framework.html) have developed a TCAD Framework which includes simulation tools for analyzing 2D/3D semiconductor device structures [60]. The TCAD tool is called GTS Framework and it consists in four simulation tools: GTS Structure, GTS Vision, MINIMOS-NT and VSP tool, Figure 3.5 and Figure 3.6.

![GTS Framework](image)

**Figure 3.5: Main screen of the GTS Framework.**

- **GTS Structure.** This is a graphical simulation tool for definition of 2D/3D semiconductor device structures. With this tool, the meshing, the doping, and the geometry of the device structure can be defined.

- **GTS Vision.** This is a visualization tool, which allows to analyze device structures and simulation results.

- **MINIMOS-NT.** This is a simulation tool for 2D/3D classical modelling of micro-scaled semiconductor device structures.
• **VSP.** The Vienna Schrödinger Poisson (VSP) is a simulation tool for 2D/3D quantum modelling of nano-scaled semiconductor device structures [65–67].

![GTS Structure](image1)

![GTS Vision](image2)

![MINIMOS-NT](image3)

![VSP](image4)

Figure 3.6: Simulation tools inside GTS Framework

### 3.4.1 Vienna Schrödinger Poisson solver VSP

The VSP simulation tool is focused in the solution of the Schrödinger-Poisson equations system. The VSP is developed in C++ programming language under the object oriented paradigm and it uses stable libraries (Blas, Lapack and Arpack) for numerical calculations.
The flowchart of the VSP module is shown in Figure 3.7. The VSP requires two input files: the semiconductor device structure created in the GTS Structure tool (Figure 3.8(a)) and the ipd file in which the initial conditions and models are defined (Figure 3.8(b)).
3.4. GTS Framework

V\textsubscript{INITIAL} is calculated by means of a classical model simulation (drift-diffusion model). This potential energy $V\textsubscript{INITIAL}$ is used in the first iteration of the Schrödinger equation and by means of the Finite Volume Method [63], and the ARPACK numerical library [64], the wavefunctions and energies are calculated. Then, the wavefunctions and energies obtained from the Schrödinger module are used in order to calculate the charge density, which then it is used in the Poisson module to obtain the electrostatic potential $\phi$. Using the electrostatic potential $\phi$ a new potential energy is calculated. If the difference between the new potential energy and the old potential energy is less than a convergence criterion, the iteration loop stops. Otherwise, the new potential energy $V\text{new}$ is used in the Schrödinger equation and the process is repeated until the convergence criterion is reached.

The output of the VSP module are the wavefunctions and the discretized energies which are saved in a .debvz file, see Figure 3.9.

The VSP procedure to solve numerically the differential equations into the Schrödinger-Poisson system is defined by three stages: definition of the boundary conditions, conversion of the differential equations into a system of algebraic equations, and finally the solution of the algebraic equations, see Figure 3.10.
• **Definition of the boundary conditions.** The VSP uses closed boundary conditions, which means that the wavefunctions vanish to zero at the $Si - SiO_2$ interface. These boundary conditions do not allow that the wavefunctions pass through the barrier potential, thus the tunneling current cannot be calculated.

• **Discretization of differential equations.** The VSP tool uses the finite volume method as a discretization method. The finite volume method (FVM) is a method to approximate the solution of a partial differential equation \[61–63\]. The FVM uses the representation of the partial differential equation (3.13) by its integral form as is show in equations 3.14 and 3.15.

\[
  u_{xx}(x) = f(x) \quad (3.13)
\]

\[
  \int_{x_{i-1/2}}^{x_{i+1/2}} u_{xx}(x) \, dx = \int_{x_{i-1/2}}^{x_{i+1/2}} f(x) \, dx \quad (3.14)
\]

\[
  u_x(x_{i+1/2}) - u_x(x_{i-1/2}) = \int_{x_{i-1/2}}^{x_{i+1/2}} f(x) \, dx. \quad (3.15)
\]

The FVM takes into account three points in the mesh $x_{i-1/2}$, $x_i$, $x_{i-1/2}$, and with this information the derivative can be approximated, see Figure 3.11.
3.4. GTS Framework

Figure 3.10: Flowchart of the VSP procedure to solve numerically a partial differential equation.

Figure 3.11: Discretization points using finite volume method.

Then by means of the midpoint integration rule \([61, 62]\),

\[
\int_{x_0}^{x_1} f(x) \, dx \approx MP(f, h) = hf(x_0 + \frac{h}{2}),
\]

(3.16)
3.5 Computer implementation of the proposed simulation methodology into the VSP tool

The right-hand side of equation 3.15 can be approximated by,
\[ \int_{x_{i-1/2}}^{x_{i+1/2}} f(x) \, dx \approx hf(x_i), \quad (3.17) \]
and the left-hand side can be approximated by central finite differences by
\[ u_x(x_{i+1/2}) - u_x(x_{i-1/2}) \approx \frac{u(x_{i+1}) - u(x_i)}{h} - \frac{u(x_i) - u(x_{i-1})}{h}. \quad (3.18) \]
Finally the discretized equations are defined as
\[ u(x_{i+1}) - 2u(x_i) + u(x_{i-1}) = h^2 f(x_i) \quad (3.19) \]

- **Solution of the algebraic equations.** After differential equations are discretized, the set of algebraic equations are solved for each node in the mesh. The system is solved by means of general solution methods (Gauss-Jordan, LU decomposition, Gauss-Seidel). Particularly, the VSP tool uses the ARPACK numerical library to solve the algebraic system [64].

### 3.5 Computer implementation of the proposed simulation methodology into the VSP tool

The Vienna Schrödinger-Poisson tool does not consider magnetic effects on the Schrödinger-Poisson coupled equations system. Moreover, the VSP neither allows to calculate tunneling currents because it considers only closed boundary conditions. In this thesis, we made a collaboration with the Technical University of Vienna and we added a module for calculating the gate tunneling current under the influence of a magnetic field.

The first step in order to incorporate the magnetic field was to define the magnetic vector potential \( \mathbf{A} \). Because in this thesis, we are interested in analyzing the magnetic effects on the gate tunneling current, the magnetic field \( \mathbf{B} \) was applied parallel to the surface of the MOSFET and perpendicular to the gate current lines (Figure 3.12).

Thus, we defined the vector potential \( \mathbf{A} \) such as
\[ \mathbf{A}(\mathbf{r}) = -y e_x B, \quad (3.20) \]
Figure 3.12: magnetic field $B$ was applied parallel to the surface of the MOSFET and perpendicular to the gate current lines.

where $\vec{e}_x$ is the cartesian unit vector along the $z$-direction and $\vec{B}$ is the component of magnetic field, in the $z$-direction.

Moreover, we had to change the Schrödinger equation by adding the vector potential $A$ in the kinetic energy term as is described in equation 3.6. The VSP solves the Schrödinger-Poisson system considering a closed system. A closed system means that all axes are confined and at the boundaries of the device geometry the wavefunctions vanish 3.13.

Figure 3.13: System with closed boundary conditions in which the wavefunctions are transmitted and reflected.

In order to calculate the gate tunneling current in a nano-scaled MOSFET, open boundaries need to be considered. The energies of an open system are complex values $E = E_r + Ei$
where the real part $E_r$ is the resonant energy and the imaginary part $E_i$ can be understood as the decay probability of the electron can have into the domain. In recent works, a method based on absorbing boundary conditions (Perfectly Matched Layer method) has been applied to determine the energies of an open system \([40, 73, 74]\). The PML method adds absorbing layers with non physical meaning which act as artificial open boundary conditions preventing reflections \([3.14]\). When the wavefunction enters to the absorbing layer, the wavefunction is attenuated and it decays exponentially.

Figure 3.14: System with open boundary conditions by adding absorbing layers in which the wavefunctions are transmitted but not reflected.

This process is done by means of replacing coordinates $x$ by stretched coordinates $\tilde{x} = x + if(x)$, where $f(x)$ represents how the imaginary axis has been deformed. Then the derivatives of the system can be expressed as

$$
\delta \tilde{x} = \left(1 + i \frac{\delta f(\tilde{x})}{\delta \tilde{x}} \right) \delta x,
$$

(3.21)

There is a variable change in order to define the equation in terms of the real coordinates $x$ where $s(x) = 1 + i\sigma(x)$, then the derivatives of the equation are changed by

$$
\frac{\delta}{\delta \tilde{x}} = -\frac{1}{s_x} \frac{\delta}{\delta x},
$$

(3.22)
3.5. Computer implementation of the proposed simulation methodology into the VSP tool

In the PML regions where $\sigma$ is equal to zero, the term $s_x$ is equal to 1 and the wavefunction does not change. For the case when $\sigma > 0$ the imaginary part of the wavefunction increases and as a result the wavefunction decays exponentially. This happens because the wavefunction can be seen as $e^{ikx} = e^{ik(Re_x + Im_x)} = e^{ikRe_x - kIm_x}$, thus the solution exponentially decays as $Im_x$ increases (where the equation is evaluated in the positive real and imaginary axis).
3.5. Computer implementation of the proposed simulation methodology into the VSP tool
Chapter 4

Results and discussion

4.1 Magnetic effects on wavefunctions and energies

As a first benchmark device a circular n-type Si nanowire of 30 nm with equivalent oxide thickness $EOT$ of 2.0 nm, $N_D = 1 \cdot 10^{17} \, \text{cm}^{-3}$ and $N_A = 3 \cdot 10^{17} \, \text{cm}^{-3}$ was used for analyzing the magnetic effects on the wavefunctions and energies. For this experiment, closed boundary conditions were used. Thus, the wavefunctions are confined in the silicon region and they vanish to zero at the $Si – SiO_2$ interface. The magnetic field $B$ was applied perpendicular to the $xy$ plane of the nanowire, see Figure 4.1.

![Figure 4.1: Circular nanowire structure](image)

35
4.1. Magnetic effects on wavefunctions and energies

Figures 4.2 and 4.3 show the probability density $|\psi|^2$ of wavefunctions of the second and sixth state respectively, under the influence of a magnetic field $B$ of 0T, 50mT, and 500mT.

When a perpendicular magnetic field is applied, there is an effect called hybridization or mixture of states and the wavefunctions are modified.

This happens because when a magnetic field is applied the Hamiltonian is perturbed and no longer separates the states and they begin to mix.

As a direct consequence the mixing of states results in new hybrid states, which results in an increase of the probability of found electrons where before cannot be found.

The hybridization between two degenerated electronic states produces a repulsion between each other in energy which lifts the degeneracy of the states.

![Figure 4.2: Probability density $|\psi|^2$ of the second level wavefunction of a cross section of a circular nanowire under the influence of a magnetic field applied perpendicular to the $xy$ plane.](image)

Moreover, when hybridization between two degenerated electronic states occurs, the states repulse each other in energy resulting in an energy splitting which is called Zeeman effect [71, 72] (Figure 4.4).
4.1. Magnetic effects on wavefunctions and energies

Figure 4.3: Probability density $|\psi|^2$ of the sixth level wavefunction of a cross section of a circular nanowire under the influence of a magnetic field applied perpendicular to the $xy$ plane.

Figure 4.4: Split energies due to the applied magnetic field.

Furthermore, in Figure 4.5 we see the probability density of the second state $|\psi_2|^2$ and the density probability of the third state $|\psi_3|^2$. 
For this case, both states are degenerated because they have the same energy value. When a perpendicular magnetic field is applied, the second and third states begin to mix and with stronger magnetic fields both states will exhibit a circular or ring shape [72].

Figure 4.5: Hybridization of $\psi_2$ and $\psi_3$ states.
4.2 Magnetic effects on nano-scaled MOSFET

Asymmetrical channel conductivity

In the work’s of P´ ondigo developed at INAOE [6,68], the $I_d - V_g$ characteristic curve of the 28-nm nMOSFET, without applying an external magnetic field, was measured (Figure 4.6). Ideally, the transistor should be off at $V_g = 0.0V$, but we see in Figure 4.6 that there is a leakage current $I_{off}$ of 10pA at $V_d = 0.1V$.

Furthermore, the subthreshold sope $S$ should be close to its ideal 60mV/dec value. The subthreshold slope $S$ is given by

$$S = \left[ \frac{\delta V_g}{\delta \log(I_d)} \right] = \ln(10) \frac{mkT}{q},$$

where $k$ is the Boltzmann constant, $q$ is the electron charge, and $m$ is a numerical factor in the 1 to 2 range.

An external magnetic field was applied perpendicular to the surface of the MOSFET, the leakage current $I_{off}$ the subthreshold slope $S$ were measured as a function of the magnetic field $B$ (Figure 4.7).

We see that there is no parabolic reduction of the $I_{off}$ current and neither in the subthreshold slope $S$ [69]. Under an ideal situation, for negative $B$ field the current lines should be deflected by the Lorentz force to the left side in the x-axis direction, while for positive $B$ field to the right side of the FET channel plane showing a symmetrical behavior, see inset Figure 4.7.

Moreover, they measured the transconductance $g_m$ and the threshold voltage $V_t$ as a function of a magnetic field (Figure 4.8). We see that there is an asymmetry in the transconductance and also in the threshold voltage. There is an asymmetry in the experimental results which cannot be explained with the assumption of an isotropic and homogeneous channel FET conductance [5]. This non-homogeneous conductance can be attributed to the
4.2. Magnetic effects on nano-scaled MOSFET

Figure 4.6: Measurement $I_d - V_g$ characteristic curve of a 28-nm nMOSFET with $B = 0$ mT with $V_{DS} = 0.1$ V.

Figure 4.7: Measurement of the leakage current $I_{off}$ and subthreshold slope $S$ for 28-nm nMOSFET as a function of a magnetic field applied perpendicular to the surface of the MOSFET with $V_{DS} = 0.1$ V.
4.2. Magnetic effects on nano-scaled MOSFET

Figure 4.8: Measurement of the transconductance \( gm \) and subthreshold voltage \( Vt \) for 28-nm nMOSFET as a function of a magnetic field applied perpendicular to the surface of the MOSFET with \( V_{DS} = 0.1 \) V.

asymmetry and non-homogenity caused by the mechanical stress induced in the transistor or by process variations [3].

Magnetic effects on the gate tunneling current

In order to analyze the non-homogenity of the gate tunnel current under the influence of an external magnetic field \( \vec{B} \), we used as a benchmark device a 28-nm n-type Si MOSFET with equivalent oxide thickness \( EOT \) of 2.0 nm, gate width \( W \) of 1.0 \( \mu \)m, \( N_D = 5 \cdot 10^{19} \) cm\(^{-3} \) and \( N_A = 3 \cdot 10^{18} \) cm\(^{-3} \), see Figure 4.9.

We applied an external magnetic field parallel to the surface of the channel (z-axis) and perpendicular to the gate current lines (Figure 4.10). The measurement tunneling current was compared with the results obtained from our simulation tool. Figure 4.10 shows the measured and simulated magnetomodulated gate current \( \Delta I_G = I_{GB \neq 0} - I_{GB = 0} \) under the influence of a static magnetic field \( B = \pm 400\text{mT} \) with \( V_{DS} = 0.5 \) V. There is a good agreement between measurement data and the results obtained with our simulation tool. The error obtained
4.2. Magnetic effects on nano-scaled MOSFET

(a) Doping definition

(b) Grid definition

Figure 4.9: Definition of a nano-scaled 28-nm MOSFET into the GTS Structure tool.

between simulation and measured data could be associated with the fact that a homogeneous and symmetrical oxide is considered in the simulation settings.

Figure 4.10: Magneto-modulated gate current $\Delta I_G$ for 28-nm nMOSFET at $V_{DS} = 0.5$ V in a magnetic field $B = -400$ mT.

As we mentioned before, a strained 28-nm MOSFET may have a non-uniform conductance along the length and width of the channel [3, 6], such a non-uniformity of the conductive properties of the channel can be recreated by means of a non-uniform gate oxide thickness or by means of a non-uniform doping profile. In this work, we recreate such a non-uniformity for both cases.
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Figure 4.11 shows the percentage of change of magnetomodulated gate current (\(\Delta I_G/I_G\)) for three different non uniform oxide profiles. Oxide number one has \(T_{oxL} = 2.0\) nm and \(T_{oxR} = 2.5\) nm, oxide number two has \(T_{oxL} = 2.0\) nm and \(T_{oxR} = 2.6\) nm, and oxide number three has \(T_{oxL} = 2.0\) nm and \(T_{oxR} = 2.7\) nm. We see that there is a higher percentage of change of \(\Delta I_G\) for oxide three which is the most non uniform device structure. In Figure 4.12 we show that the asymmetry of \(\Delta I_g - B\), with respect of the \(\Delta I_G\) when \(\vec{B} = 0\), increases and tends to saturate at large values for the oxide thickness non-uniformity.

Figure 4.11: Simulated percentage of change of magneto-modulated gate current \(\Delta I_G/I_G(\%)\) for 28-nm nMOSFET with \(V_{DS} = 0.0\) V and \(V_G = 1.0\) V.

Moreover, the behavior of the supply function \(S\) and the transmission coefficient \(TC\) for the third oxide profile under the influence of a magnetic field are shown in Figure 4.13. We see that for high values of the magnetic field \(B\), depending on the magnetic field polarity, the transmission coefficient decreases or increases. This happens due to the non uniformity of the oxide, there is a higher transmission probability when electrons are deflected by the Lorentz force to the thinner oxide side. Moreover, the supply function decreases when a magnetic field is applied. A reduction of the supply function implies a decrease of the charge population with an energy range with a transmission probability to tunnel through the gate oxide. Therefore, a reduction of the supply function implies a reduction of the current that tunnels through the gate oxide.
4.2. Magnetic effects on nano-scaled MOSFET

Figure 4.12: Simulated percentage of change of magneto-modulated gate current $\Delta I_G/I_G(\%)$ for 28-nm nMOSFET with an increase in the right side in the x-axis direction (towards the drain side) of the oxide thickness $ToxR$ in a magnetic field $B = -500\text{mT}$.

Figure 4.13: Simulated supply function $Sf$ and transmission coefficient $TC$ for oxide number three under the influence of a magnetic field with $V_{DS} = 0.0\text{V}$ and $V_G = 1.0\text{V}$.

In order to study the effects of a nano-scaled MOSFET having an non homogeneous channel conductive properties, a structure with three different non uniform doping profiles has been used, see Figure 4.14. Doping profile number 1 has $DopL = 2 \cdot 10^{18}\text{ cm}^{-3}$ and
$DopR = 5 \cdot 10^{18}$ cm$^{-3}$, doping profile number 2 has $DopL = 1 \cdot 10^{18}$ cm$^{-3}$ and $DopR = 5 \cdot 10^{18}$ cm$^{-3}$, and doping profile number 3 has $DopL = 9 \cdot 10^{17}$ cm$^{-3}$ and $DopR = 5 \cdot 10^{18}$ cm$^{-3}$.

Figure 4.14: Non uniform doping distribution profiles in the channel for 28-nm nMOSFET.

Figure 4.15 shows the percentage of change of magnetomodulated gate current ($\Delta I_G/I_G(\%)$) for the defined non uniform doping profiles.

In Figure 4.15, we see that there is a higher percentage of change of $\Delta I_G$ when a magnetic field with positive polarity is applied. With a positive $B$ field the current lines are deflected by the Lorentz force to the right side in the x-axis direction (towards the drain side) where there is a larger doping profile. A larger doping profile implies an increase of the threshold voltage which results in a lower inversion layer charge population. Therefore, the supply function decreases and as a direct consequence the tunneling current decreases. The most non-uniform doping profile number three shows the most pronounced asymmetrical magnetomodulated gate current $\Delta I_G$.

Figure 4.16 shows the relation between the non-uniformity of the semiconductor doping profile $\Delta Doping = DopR - DopL$ and the magnetomodulated gate current $\Delta I_G$. The highest doping profile asymmetry results in the largest asymmetrical magnetomodulated gate current $\Delta I_G$. 
4.2. Magnetic effects on nano-scaled MOSFET

Figure 4.15: Simulated percentage of change of magneto-modulated gate current $\Delta I_G/I_G(\%)$ for 28-nm nMOSFET for three different channel doping profiles in a magnetic field $B = \pm 400$ mT with $V_{DS} = 0.0$ V and $V_G = 1.0$ V.

Figure 4.16: Simulated percentage of change of magneto-modulated gate current $\Delta I_G/I_G(\%)$ for 28-nm nMOSFET with channel doping profile variations in a magnetic field $B = \pm 400$ mT with $V_{DS} = 0.0$ V and $V_G = 1.0$ V.

Thermal effects on the gate tunneling current

In order to study the effects of the temperature on the gate tunneling current of a nano-scaled MOSFET, Figure 4.17 shows the relation between temperature and gate tunneling current.
4.2. Magnetic effects on nano-scaled MOSFET

$I_G$ from simulated and experimental data. Moreover, Figure 4.18 shows the same relation $I_g - Temp$ in logarithmic scaled. For this experiment, we used a as a benchmark device a 28-nm n-type Si MOSFET with equivalent uniform oxide thickness $EOT$ of 2.0 nm, gate width $W$ of 1.0 $\mu$m, $N_D = 5 \cdot 10^{19}$ cm$^{-3}$ and $N_A = 3 \cdot 10^{18}$ cm$^{-3}$. For the simulation device, we assumed a uniform oxide and a uniform doping profile along the channel device.

Figure 4.17: Ig-Vg characteristics of 28-nm nMOSFET under the influence of thermal variations at $V_{DS} = 0.5$ V.

Figure 4.18: Semi-logarithmic Ig-Vg characteristics of 28-nm nMOSFET under the influence of thermal variations at $V_{DS} = 0.5$ V.
There is a good agreement between measurement data and the results obtained with our simulation tool. For the case of \(V_g < 0.2\) the simulation tool sets \(I_g = 0.0\). Because of that, the error obtained considers only the values of \(I_g\) for \(V_g \geq 0.2V\).

The tunneling current rises as there is an increase in temperature. This happens because temperature affects the behavior of the supply function \(S\) and the transmission coefficient \(TC\); see Figure 4.19. With temperature, both the supply function and the transmission coefficient increase [70], and so does the gate tunneling current. For the case of the supply function with more temperature there are more electrons available to tunnel through the barrier potential. While for the case of the transmission coefficient with more temperature the effective mass is changed and there is an increment in the transmission coefficient.

Figure 4.19: Simulated supply function and transmission coefficient of 28-nm nMOSFET under the influence of thermal variations at \(V_{DS} = 0.5\) V.

**Simulation tool performance**

In order to analyze the performance of our simulation tool, the runtimes and percentual errors of our methodology were computed. Figure 4.20 shows the runtime of our methodology with
respect to the grid size definition for one voltage step $V_g$. We see that as it was expected, with a finer grid size the runtime increases. Moreover, we saw that for our particular device size, the device can be divided into a grid size lower than 0.5 nm the potential energy converges.

Figure 4.20: Runtime dependence on the grid size definition.

Furthermore, we analyzed the runtime of our methodology with respect to the number of the eigenvalues calculated, see Figure 4.21. It is important to say that only the ten first eigenvalues have a real contribution to the gate current [40].

We also tested the runtime when a magnetic field is applied and we see that there is a difference of 4.453 seconds approximately for considering the magnetic field in our methodology.

The average percentual error between measurement data and simulation results is shown in Figure 4.22. We can see that the maximum error is obtained with a $V_g = 0.2V$ and the minimum error is obtained with a $V_g = 0.7V$. 


4.2. Magnetic effects on nano-scaled MOSFET

Figure 4.21: Runtime dependence on the number of eigenvalues computed.

Figure 4.22: Percentual average error simulations.
Chapter 5

Conclusions and future work

A 2D simulation methodology for analyzing thermo-magnetics effects on nano-scaled MOS transistors has been introduced. This has been achieved by incorporating the magnetic field and temperature as additional variables into the solution of the Schrödinger-Poisson coupled system using the Vienna Schrödinger-Poisson solver (VSP) [46]. This implementation into the GTS Framework [60] simulation tool allows us to calculate the gate tunneling current in nano-scaled MOSFET devices under the influence of thermo-magnetics effects.

The calculation of the gate tunneling current has been performed using the Tsu-Esaki model and considering open boundary conditions by using the Perfectly Matched Layer method.

An external magnetic field was applied perpendicular to the gate tunneling current of the MOSFET. With the application of the magnetic field, it has been shown a mix of the states of the wavefunctions and the splitting of the energy levels of the system, which have an impact in the gate tunneling current of the MOS device. Moreover, the transmission coefficient and the supply function were calculated in function of the magnetic field and temperature variations.

The non-uniformity of the channel in real strain nMOS devices was reproduced by means of non-uniform oxide profiles and non-uniform doping profiles. By using the magnetic field we were able to scan the non-uniformity of the tunneling properties along the channel.

The proposed simulation methodology has shown a good agreement with experimental results. Therefore, the proposed simulation methodology, is a very valuable tool to investigate
non-homogeneous space distributed tunneling properties and other termo-magnetics effects on the gate tunneling current.

As a future work, in order to do more robust the gate tunneling model, more gate tunneling mechanisms need to take into account; for instance trap assisted tunneling.

Another future work can be focused on studying the magnetic field effects having different orientations and directions. Until now, the VSP can handle 2D structures; however, in order to consider the application of magnetic fields in different directions, the VSP tool needs to handle with 3D structures. With the inclusion of this characteristic, device structures like FINFETs could be analyzed.

Finally, another future work is the implementation of a drain current model which incorporates the thermo-magnetic field effects. With this extension of the simulation tool, thermo-magnetic effects on the drain current by means of applying an external magnetic field perpendicular to the surface of the MOSFET could be done.
Bibliography


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