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Modelling and Simulation Study of NMOS Si Nanowire Transistors

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June 2018
Dedication

To my brave sons Mustafa and Ali
Abstract

Nanowire transistors (NWTs) represent a potential alternative to Silicon FinFET technology in the 5nm CMOS technology generation and beyond. Their gate length can be scaled beyond the limitations of FinFET gate length scaling to maintain superior off-state leakage current and performance thanks to better electrostatic control through the semiconductor nanowire channels by gate-all-around (GAA) architecture. Furthermore, it is possible to stack nanowires to enhance the drive current per footprint. Based on these considerations, vertically-stacked lateral NWTs have been included in the latest edition of the International Technology Roadmap for Semiconductors (ITRS) to allow for further performance enhancement and gate pitch scaling, which are key criteria of merit for the new CMOS technology generation. However, electrostatic confinement and the transport behaviour in these devices are more complex, especially in or beyond the 5nm CMOS technology generation.

At the heart of this thesis is the model-based research of aggressively-scaled NWTs suitable for implementation in or beyond the 5nm CMOS technology generation, including their physical and operational limitations and intrinsic parameter fluctuations. The Ensemble Monte Carlo approach with Poisson-Schrödinger (PS) quantum corrections was adopted for the purpose of predictive performance evaluation of NWTs. The ratio of the major to the minor ellipsoidal cross-section axis (cross-sectional aspect ratio - AR) has been identified as a significant contributing factor in device performance. Until now, semiconductor industry players have carried out experimental research on NWTs with two different cross-sections: circular cylinder (or elliptical) NWTs and nanosheet (or nanoslab) NWTs. Each version has its own benefits and drawbacks; however, the key difference between these two versions is the cross-sectional AR. Several critical design questions, including the optimal NWT cross-sectional aspect ratio, remain unanswered. To answer these questions, the AR of a GAA NWT has been investigated in detail in this research maintaining the cross-sectional area constant. Signatures of isotropic charge distributions within Si NWTs were
observed, exhibiting the same attributes as the golden ratio (Phi), the significance of which is well-known in the fields of art and architecture.

To address the gap in the existing literature, which largely explores NWT scaling using single-channel simulation, thorough simulations of multiple channels vertically-stacked NWTs have been carried out with different cross-sectional shapes and channel lengths. Contact resistance, non-equilibrium transport and quantum confinement effects have been taken into account during the simulations in order to realistically access performance and scalability.

Finally, the individual and combined effects of key statistical variability (SV) sources on threshold voltage ($V_T$), subthreshold slope ($SS$), ON-current ($I_{on}$) and drain-induced barrier lowering (DIBL) have been simulated and discussed. The results indicate that the variability of NWTs is impacted by device architecture and dimensions, with a significant reduction in SV found in NWTs with optimal aspect ratios. Furthermore, a reduction in the variability of the threshold voltage has been observed in vertically-stacked NWTs due to the cancelling-out of variability in double and triple lateral channel NWTs.
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I am also grateful to CEA-Leti for the training course "Reliability & Defects in Advanced CMOS Technologies".

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During my PhD journey, I met many people in Glasgow who participated in the success of this study. Thank you to all of you.

Thank you,
Author's Declaration

I declare that, except where explicit reference is made to the contribution of others, that this thesis is the result of my own work and has not been submitted for any other degree at the University of Glasgow or any other institution.

Signature

Talib Mahmood Ali Al-Ameri
June 2018


Participations

I. Contributing as a reviewer for 7 journals (2017-2018):
   • IEEE Transactions on Electron Devices.
   • Applied Sciences (MDPI).
   • Journal of Computational Electronics (Springer).
   • Materials (MDPI).
   • Electronics (MDPI).
   • IEEE Transactions on Nanotechnology.

   https://publons.com/author/983777/talib-al-ameri#profile

II. Participating in Technical Program Committee (TPC) for the 13th IEEE Nanotechnology Materials and Devices Conference (NMDC 2018, USA).

III. CEA-Leti training course (Reliability & Defects in Advanced CMOS Technologies), 2018, France.
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<td>Two dimensions</td>
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<td>3D</td>
<td>Three dimensions</td>
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<tr>
<td>CMOS</td>
<td>Complementary Metal Oxide Semiconductor</td>
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<tr>
<td>DD</td>
<td>Drift-Diffusion</td>
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<tr>
<td>DG</td>
<td>Density Gradient</td>
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<tr>
<td>DIBL</td>
<td>Drain Induced Barrier Lowering</td>
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<td>EOT</td>
<td>Equivalent Oxide Thickness</td>
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<tr>
<td>FDUTB</td>
<td>Fully Depleted Ultra-Thin Body</td>
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<tr>
<td>FinFET</td>
<td>Fin Field Effect Transistor</td>
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<tr>
<td>GAA</td>
<td>Gate All Around</td>
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<td>GSS</td>
<td>Gold Standard Simulations</td>
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<tr>
<td>IC</td>
<td>Integrated Circuit</td>
</tr>
<tr>
<td>ITRS</td>
<td>International Technology Roadmap for Semiconductors</td>
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<tr>
<td>$I_{on}$</td>
<td>On-current</td>
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<td>$I_{off}$</td>
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<td>$Si$</td>
<td>Silicon</td>
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<td>$SiO_2$</td>
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<td>$Si_3N_4$</td>
<td>Silicon oxynitride</td>
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<td>Gate dielectric thickness</td>
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<td>$T$</td>
<td>Temperature</td>
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<tr>
<td>$T_{Si}$</td>
<td>Silicon body thickness</td>
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<td>Carrier mobility</td>
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<td>$\sigma V_T$</td>
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<td>$\sigma I_{on}$</td>
<td>On-current standard deviation</td>
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<td>$\sigma DIBL$</td>
<td>DIBL standard deviation</td>
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<tr>
<td>LER</td>
<td>Line Edge Roughness</td>
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<td>MC</td>
<td>Monte Carlo</td>
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<td>Term</td>
<td>Definition</td>
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<tr>
<td>MGG</td>
<td>Metal Gate Granularity</td>
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<td>MOS</td>
<td>Metal Oxide Semiconductor Field Effect Transistor</td>
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<tr>
<td>RDD</td>
<td>Random Dopant Discrete</td>
</tr>
<tr>
<td>SCE</td>
<td>Short Channel Effect</td>
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<tr>
<td>S/D</td>
<td>Source/Drain</td>
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<td>SS</td>
<td>Sub-threshold Slope</td>
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<td>SOI</td>
<td>Silicon on Insulator</td>
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<tr>
<td>TCAD</td>
<td>Technology Computer Aided Design</td>
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<tr>
<td>c</td>
<td>Light speed</td>
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<td>Recombination rate</td>
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<tr>
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<td>BTE</td>
<td>Boltzmann transport equation</td>
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<tr>
<td>⟨τ⟩</td>
<td>Average momentum relaxation time</td>
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<tr>
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<td>q</td>
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<td>RTA</td>
<td>Relaxation time approximation</td>
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<tr>
<td>ψ_qm</td>
<td>Quantum correction</td>
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<td>Quantum correction force</td>
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<tr>
<td>r</td>
<td>Fitting parameter</td>
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<td>b_n</td>
<td>Quasi-Fermi level</td>
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<td>PS</td>
<td>Poisson-Schrödinger</td>
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<tr>
<td>NBC</td>
<td>Neumann boundary conditions</td>
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<td>n(0)</td>
<td>Electron density at the interface</td>
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<td>x_p</td>
<td>Penetration depth</td>
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<tr>
<td>WKB</td>
<td>Wentzel-Kramers-Brillouin</td>
</tr>
<tr>
<td>Symbol</td>
<td>Description</td>
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<tr>
<td>$m_{ox}$</td>
<td>oxide’s electron effective mass</td>
</tr>
<tr>
<td>$\Phi_B$</td>
<td>oxide’s potential</td>
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<tr>
<td>SOR</td>
<td>successive over relaxation</td>
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<tr>
<td>BICGSTAB</td>
<td>biconjugate gradient-stable</td>
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<td>$r_c$</td>
<td>cut-off radius</td>
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<tr>
<td>$\mu_L$</td>
<td>perpendicular field-dependent mobility</td>
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<td>$E_c$</td>
<td>critical electric field</td>
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<td>$\alpha$</td>
<td>secondary fitting parameter</td>
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<td>$S_G$</td>
<td>Gaussian autocorrelation function</td>
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Chapter 1. Introduction

1.1 Research Motivation

Silicon Complementary Metal-Oxide Semiconductor (CMOS) technology and corresponding Integrated Circuits (IC) have constantly improved in terms of functionality and performance over many decades. Metal Oxide Semiconductor Field Effect Transistors (MOSFET) is at the heart of CMOS technology and IC, with more than one billion transistors integrated today on a single IC chip. Through the advancement in IC manufacturing technology, Silicon chip manufacturers are able to deliver 7nm FinFET CMOS technology generation with 19nm channel length transistors this year. With the reduction of gate length in the FinFET CMOS technology, it is necessary to shrink the fin width in order to maintain electrostatic control and to deliver the required contact pitch scaling. In order to maintain good electrostatic control and to minimise the short-channel effects (SCE) simultaneously increasing the performance, the development of tall and narrow fins has been crucial.

The current focus of the advanced CMOS technology development has been the introduction of 7nm FinFETs CMOS technology this year by the world leading foundry TSMC [1]–[3]. Yet the FinFET size must be reduced further in order to continue scaling [4]. The issue involved reducing the FinFET fin width while sustaining or even increasing the fin height, however, is the mechanical fin integrity, the increase in process variability and the subsequent increase in the statistical variability of the transistors [5]. It is highly challenging to control the channel geometry and shape when narrowing and elongating the fin for the purpose of maximising drive current. Although the increase in SCE control and threshold voltage ($V_T$) that arises during the scaling of the fin width ($W_{\text{fin}}$) leads to a reduction in leakage current ($I_{\text{off}}$), scaling of the $W_{\text{fin}}$ below 4nm results in a significant $V_T$ increase and $I_{\text{on}}$ saturation and even reduction. The reason for this is the growing contribution of quantum confinement effect to the charge distribution, scattering and carrier transport, suggesting that
scaling below a specific gate length and channel cross-section is challenging to achieve in practice. This highlights the issues involved with the extreme scaling of FinFET transistors.

Silicon Nanowire Transistors (NWTs) represent a potential alternative to Si FinFET technology, perhaps taking the lead compared with the exploration of other performance improvement options investigating alternative channel materials [6]. Here, the gate length can be scaled beyond the minimum FinFET channel length scaling whilst managing to keep the off-state leakage minimal, due to the greater electrostatic control over semiconducting nanowire channels offered by the gate-all-around (GAA) NWT architecture. Researchers have confirmed the advantages of the GAA transistors when dealing with short-channel effects, highlighting GAA transistors as the most effective option for ultimate channel length scaling. In principle, the engineering and optimisation of GAA NTW transistors requires an ideal balance between leakage current and device performance. In earlier research [7]–[9], it has been proven that NWT mobile charge and gate capacitance are significantly affected by the cross-sectional nanowire shape, with the greatest mobile charge per gate voltage observed in a <110> channel-orientation NWT with an elongated elliptical (shape) cross-section, and the longer diameter running parallel to the Si wafer surface. Additionally, it has been shown that the device performance is significantly influenced by the major-minor cross-sectional axis ratio, or aspect ratio (AR). Experimentally, NWTs are largely available with a circular or elliptical shape [10], [11] or as nanosheet/nanoslab FETs geometries [12]. There are benefits and drawbacks to each of the above types, but the cross-sectional AR is one of the main design features determining NWTs performance. At present, there is no clear answer to the question of what the ideal cross-sectional AR in NWTs is. Other key design issues including the use of multiple NWT channels, which also remains poorly understood.

Given the above points, the current research is mainly focused on the simulation-based design optimisation of NWTs suitable for the 5nm Si CMOS technology generation. This includes optimisation of the NWTs cross section to achieve an optimal quantum mobile charge to gate capacitance ratio [7], [9]. Here, the NWT’s intrinsic speed [13] is measured through intrinsic delay (τ). The simulations also take into account contact resistance, non-
equilibrium transport and quantum confinement effects in order to achieve optimisation based on realistic predictive Technology Computer Aided Design (TCAD) simulations.

The increase in the saturation current \( I_{\text{sat}} \) needed for the 5nm CMOS node has been thoroughly evaluated in the simulations in order to ensure that the performance improvement expected by the industry in this technology generation is attainable. In recent research the adoption of a replacement metal gate (RMG) technique has proved effective in GAA stacked-NWTs. However, these devices, which are similar to RMG FinFETs, come with certain technical requirements. Research demonstrates that the 3D vertically-stacked channels bring high drive current at the optimal layout footprint of GAA NWTs. Compared to vertical NWTs, which are associated with more technological challenges, horizontal GAA NW devices have the advantage that they can be manufactured without varying the current FinFET technology too much. Therefore, many semiconductor manufacturers are at present considering the use of GAA stacked-NWT architecture for extreme CMOS scaling. Nanowire stacking may be necessary to improve per-footprint drive current. Given these points, the last 2015 edition of the International Roadmap for Semiconductor (ITRS) incorporates vertically-stacked horizontal nanowire GAA transistors in order to bring the contacted gate pitch down to less than 40nm by 2021. This is one of the major viable approaches in terms of increasing the CMOS device density beyond the 7nm CMOS technology. The current study therefore explores vertically-stacked NWTs in line with their significance for future CMOS developments in the semiconductor industry.

In order for the chipmaker to make the decision to implement a particular technology, complex research is needed where predictive simulations play an extremely important role. The simulation and evaluation of the statistical variability introduced by the discreteness of charge and granularity of matter is key in the evaluation of future potential technologies. The study of the statistical transistor parameter distributions requires 3D simulation of large statistical samples of microscopically different transistors. Metal gate granularity (MGG), line edge roughness (LER) and random discrete dopants (RDD) are among the key sources of statistical variability and have been explored by many researchers. In the context of the current study, the exploration of the statistical variability in stacked NWTs has been carried out with the most advanced TCAD simulation tools available.
1.2 **Research Aim and Objectives**

The current study aims to investigate the scaling of silicon NWT MOSFETs taking into consideration quantum confinement effects, performance and statistical variability in order to evaluate their applicability for the 5nm CMOS technology generation and beyond. Therefore, the objectives of this research are:

1. To design a realistic, highly-scaled Si NWT in line with new generations of technological advancement and requirements. This entails:
   - Designing the device structure according to the ITRS criteria, and following up research results;
   - Exploring the ways in which NWT gate capacitance, transport charge, subthreshold slope (SS) and drain-induced barrier lowering (DIBL) are impacted by the cross-sectional nanowire shape.
   - Identifying the ideal cross-sectional shape and cross-sectional AR;
   - Applying the ITRS criteria and up-to-date research findings in order to optimise nanowire configuration.
   - Thoroughly investigating quantum confinement effects in <5nm-CMOS technology compatible NWTs with different cross-sections.

2. Performing Poisson–Schrödinger (PS) and density gradient (DG) based quantum corrections in order to determine the suitability of the DG approach to NWT design and simulation.

3. To utilise the experimental design for predictive understanding of the impact of source/drain doping, spacer and gate lengths and trade-offs between $I_{on}$, $I_{off}$, DIBL and SS in order to optimise nanowire configuration.

4. To determine: a) whether a strained single-channel silicon NWT can be used to achieve the 5nm technology node performance target; and b) the number of lateral channels required in a single device if more than one channel is needed to achieve the performance target.
5. To address the gap in the existing literature, which explores scaling using only single-channel NWT simulation, through the simulation of NWTs with numerous lateral channels and channel cross-sections and lengths. Contact resistance, non-equilibrium transport and quantum confinement effects are taken into account during the simulation in order to achieve predictable simulations and realistic conclusions.

6. To carry out a study of the NWT statistical variability using 3D simulation of large statistical samples employing the key statistical variability sources (e.g., RDD, LER, MGG).

7. To analyse the effects of statistical variability on threshold voltage $V_T$, on-current $I_{on}$, and DIBL.

8. To compare the performance of vertically-stacked NWTs to that of single-channel NWTs with the incorporation of sources of statistical variability.

1.3 Thesis Outline

This chapter has presented an introduction to the research topic, highlighting the significance of NWT scaling to future developments within the industry. The chapter began with a discussion of the motivation for the research, with the research aim and objectives outlined thereafter.

Chapter 2 presents a review of the existing research and current issues associated with MOSFET scaling and short-channel effects. This chapter also discusses the physics behind the topic along with the scaling theory. It introduces multi-gate MOSFET architectures, including FinFET and GAA NWTs. The key variability sources are then discussed outlining their effects on the performance of devices and circuits.
The research methodology is presented in Chapter 3, with a discussion of the main simulation approaches and tools, and the physical mechanisms relevant to the exploration of multi-gate CMOS devices, used in this research. This section specifically evaluates the drift-diffusion (DD) and Monte Carlo (MC) models, outlining their advantages and limitations in terms of studying quantum mechanical effects and non-equilibrium transport in nano-scaled devices. The reasons behind the adoption of a calibrated DD-based quantum-corrected 3D Monte Carlo method are then explained.

Chapter 4 begins with a description of the NWT design adopted in this study, along with a discussion of key design considerations such as the main NWT structure design parameters, the structure editor and the doping profiles used. Following this, the design specifics for the 5nm CMOS compatible NWT are then outlined, with the simulation methodology and density gradient calibration also being discussed. The chapter ends with an exploration of the impact of quantum confinement on the electrostatic integrity (e.g., SS and DIBL) of the nanowire based on an analysis of quantum confinement effects on channel gate capacitance and transport charge. The optimal cross-sectional AR is then investigated.

There are four sections presented in Chapter 5. In the first section, non-equilibrium transport in single mono-channel NWT simulation is addressed. In the second section, multi-lateral channel 5nm CMOS compatible NWT simulation is explored, with contact resistance, non-equilibrium transport and quantum confinement effects being considered in the 3D MC simulation model. The remainder of the chapter then focuses specifically on two questions, in relation to the research objectives. The first question is whether the semiconductor industry target can be met in highly-scaled 5nm CMOS technology compliant single-channel silicon NWTs. The second question addresses the alternative if the target cannot be met using single-channel NWTs, i.e. elaborating how many lateral channels are needed within a single NWT to achieve this target.

The introduction of variability sources in the DD-based simulator is then presented in Chapter 6, with an outline of the approaches used for this purpose. Here, calibration of the DD model is achieved with the MC and Poisson-Schrödinger approaches. This allows for the exploration of the impact of the key sources of variability on NWTs. The simulation of
Chapter 1. Introduction

Thousands of devices with random discrete dopants (RDD), metal gate granularity (MGG) and line edge roughness (LER) is then discussed, with the results presented and analysed.

Chapter 7 is the final chapter of this dissertation. This chapter reviews the general findings and conclusions from the research presented in the thesis. It also provides directions for further research in the vibrant area of NWT simulations.
Chapter 2. Background

2.1 Introduction:

Over the last fifty years, silicon-based microelectronics has transformed our lives. The requirement for improved computing technology at lower costs has driven unrelenting CMOS scaling. The development of integrated circuits towards the end of the 1950s first revealed the potential for employing transistors in practically every type of electronic circuit [14]. The first MOS transistor was patented in 1928 by Lilienfeld [15] [16]. Subsequently, production of the first metal-oxide semiconductor field-effect transistor (MOSFET) in 1960 by Kahng and Atalla permitted the cost-effective integration of multiple transistors with interconnects on a single silicon chip [14], [17]. This was followed in 1965 with the postulation of Moore’s Law. Gordon Moore made the momentous observation that the number of components in integrated circuits had increased by a factor of approximately two per year [18] and predicted that this trend would continue in the future. Still standing strong half a century later, Moore’s Law has held its ground despite the frequent challenges it has faced over the decades. Industry response has been to attempt to reach the predicted target set by Moore’s Law, for fear that this target will be reached by competitors by any means. Moore’s Law with regard to the number of transistors in microprocessors is depicted in Figure 2-1, whilst Figure 2-2 depict Moore’s Law with respect to the number of transistors [19] and the processor area for Intel’s microprocessors, correspondingly [19]–[21].

A modern microprocessor contains a few billion transistors. Nowadays Moore’s Law is slowed down by doubling the number of transistors on the chip every three years instead of the original two years because the scaling of the transistors’ dimensions has become more challenging [22]. Transistor scaling has a number of benefits over and above increasing the on-chip transistor density [23]. For example, the delay of the logic gates is reduced and the operating frequency of the transistors is increased by a factor of 1/L_{g} (where L_{g} = transistor gate length) which permits faster circuit operation. For an equivalent degree of functionality, the chip area is reduced by a factor of 1/(L_{g})^{2} and this permits an increased number of dies
to be produced on a single wafer thereby reducing production costs. In addition, as the die size is smaller, the quantity of defects per die is also reduced, resulting in an increased manufacturing yield and productivity [24]. The active switching power per area stays steady, with technology scaling permitting the circuits to operate at reduced power or permitting the circuits to have greater functionality at fixed power. Whilst planar bulk silicon MOSFETs have remained the backbone of the semiconductor industry in attaining constant scaling, the bulk planar FET struggles to deliver scaling benefits beyond the 32/28nm CMOS technology generation [25].

From the early 1990s the semiconductor industry and academia have worked together to forecast the industry’s future. These efforts have been formalised in an international organisation - the International Technology Roadmap for Semiconductors (ITRS) [26] which remained in existence until 2015. The ITRS used to produce bi-annual reports which contained predictions, recommendations and guidelines for the semiconductor industry. The ITRS reports outlined the advancements of the technology, design tools, equipment and metrology tools that need to be developed in order to maintain the exponential evolution of the semiconductor chips needed to sustain Moore’s Law.

The backbone technology of the semiconductor industry is silicon Complementary Metal Oxide Semiconductor (CMOS) technology, with the fundamental unit of CMOS chips being the MOS Field Effect Transistor, the MOSFET. In order to keep up with the frenetic development speed prescribed by Moore’s Law, transistor linear dimensions have decreased by 70%, initially every two years, and since the beginning of this century every three years the transistor and the chip areas have been reduced by 50% in every new technology generation. The sub-micron dimension milestone was passed in the first half of the 1980s, and this year (2018) regular mass production of the 7nm FinFET CMOS technology will be in place with transistors with a 19nm gate length.

Although initially integrated circuits and the corresponding transistors were being manufactured on “bulk” silicon wafers by the end of the 1990s, it was realised that major performance enhancements could be achieved by introducing a novel type of substrate, known as Silicon-On-Insulator (SOI) substrate. Here, transistors are produced in a thin
silicon layer located on the top of a silicon dioxide layer. SOI technology offers performance enhancements in both circuit speed and power reduction. At the beginning of the 2000s, IBM commenced production of microprocessors utilising SOI substrates on an industrial scale. These SOI devices provided benefits of reduced parasitic capacitances and enhanced the drive current.

**Figure 2-1 Moore’s Law with regard to count of transistors and year of introductions [19].**

**Figure 2-2 Moore’s Law with regard to the number of transistors of transistors in Intel’s microprocessors [19-21].**
2.2 Bulk MOS transistor scaling

A number of technical challenges rendered Si bulk MOS transistor scaling impossible in the technology generation of below 22/20-nm. In a long channel, bulk MOSFETs’ transistor operation can be described using the gradual channel approximation treating the lateral and vertical components of the electrical field in the 2D transistor cross section separately: (1) a gate-controlled vertical electric field is responsible for the charge formation in the channel, and (2) a drain-controlled lateral field governs the charge transport [27]. In long channel MOSFETs the threshold voltage $V_T$ is independent of the drain voltage. The application of the gate voltage reduces a potential barrier between the source and the drain at the interface between the silicon and the gate insulator and permits electrons to flow from the source to the drain. During normal transistor operation, basic thermionic emission limits the subthreshold slope (SS) to a minimum of 60 mV/decade at 300K. Degraded 2-D electrostatic integrity at short gate lengths increases the SS above 60 mV/decade resulting in increased off-state leakage current at identical values of $V_T$. In practice, the potential barrier at the source is regulated by the gate, in addition to the drain via their respective capacitive coupling to the charge in the channel [28]. Reducing gate length, the drain influence increases. Consequently, it becomes more difficult for the gate to regulate the channel barrier and turn off the channel. The 2-D short channel effects (SCE) have multiple manifestations. First, they result in a reduction of the threshold voltage with the reduction of the gate length ($V_T$ roll-off). Secondly the $V_T$ is reduced with the increase in the drain voltage (which is also termed ‘drain induced barrier lowering’, or DIBL, at the interface) Thirdly the SCEs degrade the subthreshold swing. Taken together with the SCE, they result in an increase of off-state static leakage power.

MOSFET design requires careful offsetting between drive current, short channel effects and power consumption. The on-state current ($I_{on}$) of a MOSFET can be approximated by

$$I_{on} = WQ_{inv}(V_{GG}) \times \nu(V_{DD}) \approx W C_G(V_{GG} - V_T) \times \nu(V_{DD})$$  \hspace{1cm} (2-1)
where $W$ represents the device’s width, $V_T$ represents threshold voltage, $Q_{inv}$ represents the inversion charge density at the maximum potential barrier near the source and $v$ represents the velocity close to the source region (injection velocity) which is dependent on the drain bias $V_{DD}$. The power consumption (PC) can be estimated by

$$P_{dis} = Dynamic \ power + Static \ power$$

(2-2)

$$Dynamic \ power = \alpha f C_{load} V_{DD}^2$$

(2-3)

$$Static \ power = V_{DD} \left( I_{leak} + I_{th} 10^{-\left(\frac{V_T}{SS}\right)} \right)$$

(2-4)

where $\alpha$ is the activity factor, $f$ operating frequency, and $SS$ is the sub-threshold slope, $I_{leak}$ represents the total leakage current from gate, the junctions and the band-to-band tunnelling, and $I_{th}$ is the drain current at $V_T$. In order to maintain low power consumption, lower $V_{DD}$ and leakage current, higher $V_T$ and a steeper $SS$ is required according to equations (2-3 and 2-4). On the other hand, large gate capacitance, low $V_T$ and high injection velocity are required to attain high performance in terms of saturation current.

To date, device engineers have tried to reduce SCE in short gate length devices using several techniques. The first technique is by minimising the gate oxide thickness to enhance gate control over the channel. The second technique is by reducing the source/drain junction depth (particularly close to the gate edge, where the source/drain areas are known as ‘extensions’) in order to minimise the drain coupling the mobile charge in the channel and its impact on the source barrier height. The third technique is by raising the degree of channel doping to restrain the electric field lines which commence from the source and propagate towards the drain and condoling the depletion and the inversion charge under the gate. In contemporary bulk MOSFETs, bespoke channel doping is achieved by utilising complex vertical and horizontal profiles in order to reduce the SCE. The creation of complex high doping profiles in the channel incurs extra costs and also leads to a reduction in the transistor performance (speed) together with an increased static leakage current due to enhanced band-
to-band tunnelling and gate induced drain leakage (GIDL). When the gate oxide becomes 2nm thick, quantum mechanical tunnelling leads to an increase in the gate leakage current. For oxides thinner than 2nm, the direct-tunnelling gate leakage current sharply increases (~3X for each 1 Å reduction in oxide thickness). This gate leakage increases the standby power consumption and in addition can interfere with proper logic gate operation. As a consequence, most foundries now have replaced the conventional silicon dioxide (SiO$_2$) as a gate dielectric with high permittivity (high-k) gate dielectrics [29], predominantly hafnia (HfO$_2$). This results in high gate capacitance with physically thick insulators with very low probability of tunnelling. Nevertheless, the introduction of such new materials is not without challenges and achieving the desired results with no associated losses in mobility and reliability is currently a field of intensive further research.

Another challenge is the resistance of the source/drain regions. For example, when the source/drain junction depths are reduced to control SCE, there is a need to increase doping levels in order to maintain constant sheet resistance. The upper limit for the solid solubility of dopants is around $10^{20}$ cm$^{-3}$ depending on the dopant space. Consequently, the reductions in junction depth lead to raised series resistance reducing the performance of the transistor. Simultaneously, from a technological perspective, the formation of ultra-shallow junctions which does not diffuse deeper following doping activation annealing, required to achieve low resistivity [28], [30] becomes difficult. With increasing doping density in the channel for SCE suppression, the carrier mobility is degraded as a consequence of increased ionized impurity scattering. Furthermore, the subthreshold slope deteriorates as a consequence of increased depletion capacitance which impedes the control of the surface potential by the gate voltage. Due to very high channel doping close to the source/drain extensions, an additional static leakage mechanism, band-to-band tunnelling (BTBT), becomes significant. Lastly, with reducing channel volume in extremely scaled transistors, the stochastic positioning of discrete dopant atoms leads to random inter-device variations.

The requirement to increase drive currents with transistor scaling and to reduce the supply voltage is associated with an exponential increase in the static, off-state leakage of the transistor. Although the active power density of the chip has grown steadily with the gate length scaling, the static power density has increased much more rapidly. The active power
is a consequence of the dissipative current flow through the complementary CMOS transistor pairs during logic switching. The sub-threshold, static or standby power is dissipated even when the integrated circuit is inactive. The subthreshold leakage was the major leakage mechanisms for long channel transistors. For contemporary nano-scaled bulk MOSFETs BTBT, GIDL and gate leakage dominate the static power dissipation. Whilst static power dissipation was a relatively minor issue a few decades ago, in the latest 28nm and 20nm bulk technology generations it compares in magnitude to the active power. Control and suppression of static power became a major issue for continued gate length scaling in bulk CMOS technology marking eventually its end. The needed higher channel doping and halo implant required to regulate SCE also increases the source/drain to bulk parasitic capacitances slowing the speed in the latest generations’ bulk MOSFETs.

Due to the deficiencies in performance, insufficient control of the CER and exploding power consumption, traditional bulk MOSFETs reached the end of their scaling and useful life with the 20nm bulk CMOS technology which has been introduced into production by only one foundry: TSMC. New device architectures aiming to replace the conventional bulk CMOS in combination with a new channel and gate stack materials have been driving semiconductor research over the past two decades [31], [32].

![Figure 2-3 Typical PDSOI (a) and(b) FDSOI structures [32].](image)
2.3 **SOI MOSFETs**

The first alternative to the bulk MOSFET adopted commercially is the SOI transistor. As depicted in Figure 2-3, two types of SOI transistors exist: partially depleted (PD) and fully depleted (FD). In the PD SOI MOSFETs, a stratum of insulating SiO$_2$ divides the top device-containing layer from the bulk Si underneath. The PD MOSFET has a similar design and similar dimensions to a bulk MOSFET [28]. When in the OFF state, the depletion depth beneath the gate is below the depth of the top silicon layer [33]. The PD MOSFET delivers reduced parasitic Source/Drain capacitance in an inverter circuit, reducing the propagation delay and increasing switching speeds. The potential of the floating body is controlled dynamically by capacitive coupling of the range of electrodes associated with this layer. Therefore, a floating charge can build up in this area, leading to modification of the floating-body charge and the historical effects of transistor characteristics [34]–[39]. Such floating body voltage can alter the threshold voltage of the device, potentially resulting in serious discrepancies between two identical transistors [40]–[42].

PD MOSFET techniques have been successfully applied to high volume production, although this encounters the same challenges associated with scaling as the bulk MOSFET and therefore this is not a scalable technology with a long-term future [43].

A further issue with SOI transistors is self-heating [44], [45]. In SOI circuits, the active transistors are positioned on top of a thermally insulating layer of silicon oxide. When the circuit is in operation, the power generated in the active region cannot be easily dissipated through the silicon substrate. Consequently, the temperature of the silicon body increases resulting in a reduction in mobility and in drive current.

In the fully Depleted (FDSOI) MOSFET the top silicon film is thinned down to such an extent that it becomes entirely depleted in the OFF state (see Figure 2-3). Elimination of the partially depleted region in the FDSOI MOSFET enables suppression of the floating body effects. The reduction of the silicon film thickness also allows better control of the short channel effects and therefore tolerates a reduction in the channel doping densities. In addition, the vertical electric field is reduced for the same channel carrier concentration.
Therefore, the inversion layer mobility is enhanced without detrimentally affecting the OFF-state current. Hence, by increasing the buried oxide thickness, optimal subthreshold slopes of 60 mV/decade can be attained. However, this also increases the drain control on the source-channel barrier via the buried oxide. Employment of a thin buried oxide can overcome this problem through the termination of the drain field lines on the back substrate, but at the cost of a degraded sub-threshold slope [34], [46]–[50].

2.4 Multi-Gate MOSFETs

The channel gate control is improved by the structural positioning of the gate close to the channel. Improved gate control may also be achieved by utilising more gates. Multi-gate MOSFETs can be considered an extension of the fully-depleted SOI MOSFETs, but with a greater number of gates around the thin silicon body. Multi-gate MOSFETs provide greater electrostatic control of the inversion channel [51].

The improved electrostatic control in the multi-gate MOSFET minimises the detrimental short channel effects (SCE) and improves the scalability of multi-gate FETs compared to planar bulk MOSFETs. To achieve this, planar double-gate [52], fully Depleted Lean-channel Transistor (DELTA) [53], vertical double-gate (FinFET) [54]–[56], omega-shaped gate (Ω-gate) [57], [58], Φ-FET[59], and Gate-All-Around (GAA) nanowire [59]–[63] have been proposed by researchers over the last fifteen years.

The double-gate MOSFET has a symmetrical device structure where the channel is controlled by a gate on either side of the Si layer. Better gate control allows further reduction of the Si film thickness for a similar gate length compared with FD-SOI at an identical OFF-state current. Appropriate device design enables volume inversion in the thin Si film, where the majority of the inversion charge resides at the Si film centre, and the vertical field results in greatly enhanced mobility. Due to better electrostatic integrity delivered by the two gates, the channel length can be reduced further, compared with FD-SOI.
There are three techniques to fabricate a double-gate transistor. The first possibility is structuring the device laterally with two opposing single gates on the top and on the bottom as shown in Figure 2-4 (a). The second possibility is to structure the device perpendicularly to the Si substrate with the current flowing perpendicular to the Si surface as shown in Figure 2-4 (c). The third way is to structure the device with the channel and gate vertically to the Si surface but with the current flowing parallel to the surface, as shown in Figure 2-4 (b). The third structure is identical with the FinFET concept [64]. Improved structures like the tri-gate MOSFET have been employed to improve the electrostatic control of the gate permitting additional gate length reduction. Section 2.5 gives a brief overview of the scaling theory and how multi-gates improve the SCE.

Figure 2-4 Double gate FET transistor structure to improve performance whilst reducing $I_{off}$. (a) double gate FET with a channel in the plane of the wafer, with the first gate over it and the second gate beneath. The channel is vertical on the wafer’s surface with one gate on each side (b) and (c). Two structures of the vertical channel have the current direction parallel to the FinFET (b), or parallel to the wafer surface (c)[64].
2.5 Scaling Theory of MOSFET

SOI-MOSFET scaling behaviour and electrostatics is controlled by the channel surface potential which leads to the injection of carriers from the source to the drain terminal. The distribution of the electrical potential in the channel of an FD-SOI-MOSFET is given by the solution of the Poisson's equation in the depletion approximation [65], [66].

\[
\frac{d^2\psi(x,y,z)}{dx^2} + \frac{d^2\psi(x,y,z)}{dy^2} + \frac{d^2\psi(x,y,z)}{dz^2} = \frac{qN_a}{\varepsilon_{Si}}
\]  

(2-5)

Which can be rewritten as [66]:

\[
\frac{dE_x(x,y,z)}{dx} + \frac{dE_y(x,y,z)}{dy} + \frac{dE_z(x,y,z)}{dz} = C
\]

(2-6)

This form of the equation indicates that, at arbitrary \((x, y, z)\) point in the device’s channel, the total change of the electrical field components remains equivalent to a constant \(C\). Therefore, if any one of the components of the electric field increases, the sum of the other two must decrease. The \(x\)-component of the electric field \(E_x\) represents the encroachment of the drain electric field on the channel region, and thereby is responsible for the short-channel effects. The impact of \(E_x\) on a potential barrier near the source positioned at coordinates \((x, y, z)\) can be minimised either by lengthening the channel, or by increasing the electric field inserted by the top/bottom gates, along the direction \(y\), or the lateral gates along the direction \(z\) of the channel. This can be achieved by minimising the silicon film thickness, and/or the fin thickness. Furthermore, an increase of \(\frac{dE_y(x,y,z)}{dy} + \frac{dE_z(x,y,z)}{dz}\) and, therefore, an improved regulation of the channel by the gates and less short-channel effects, may also be attained by raising the number of gates: \(\frac{dE_y(x,y,z)}{dy}\) which can be achieved by the existence of a top and bottom gate instead of merely a single gate, with \(\frac{dE_z(x,y,z)}{dz}\) further raised by the existence of a lateral gate [66][65].
Poisson’s equation can be used to obtained the electrostatic potential in the channel, using closed form parabolic expression proposed by Young [67]:

$$\psi(x,y) = c_0(x)y^0 + c_1(x)y^1 + c_2(x)y^2$$  \hspace{1cm} (2-7)

where x and y represent directions along and perpendicular to the direction of current respectively. The surface potential ($\psi_s$) for the variety of channel geometries, can be represented by the 1D Poisson’s equation. It can be written simply as:

$$\frac{d^2 \psi_s(y)}{dy^2} + \frac{\psi_s(y) - \psi_{gate} - \psi_{built}}{\lambda^2} = \frac{q(\rho \mp N)}{\varepsilon_{Si}}$$  \hspace{1cm} (2-8)

where $\psi_s$, $\psi_{built}$, $\rho$, $N$ and $\varepsilon_{si}$ represent the gate and built in potentials, mobile and channel doping densities and channel permittivity electric field. The 2nd order differential equation (2-8) has a solution expressed in an exponential form:

$$\psi_s(y) \propto e^{-\frac{y}{\lambda}}$$  \hspace{1cm} (2-9)

where $\lambda$ represents the so-called natural length (or screening length), i.e. the length scaling the channel potential variation. The natural length depends on the device geometry. The analytical solutions for single gate [67], [68], double gate, and gate-all-around structures are expressed by equations (2-10), (2-11) and (2-12), respectively [51]:

$$\lambda_{single} = \sqrt{\frac{\varepsilon_{si}}{\varepsilon_{ox}}} \frac{t_{si}t_{ox}}{\varepsilon_{ox}}$$  \hspace{1cm} (2-10)
\[
\lambda_{\text{double}} = \sqrt{\frac{\varepsilon_{si}}{2\varepsilon_{ox}}} \left(1 + \frac{\varepsilon_{ox}t_{sl}}{4\varepsilon_{si}t_{ox}}\right)t_{sl}t_{ox}
\]

\[
\lambda_{\text{GAA}} = \sqrt{\frac{2\varepsilon_{si}t_{sl}}{16\varepsilon_{ox}}} \left[\ln\left(1 + \frac{2t_{ox}}{t_{sl}}\right) + \frac{t_{si}t_{ox}}{16\varepsilon_{ox}}\right]
\]

The purpose of scaling is to achieve optimum performance from each MOSFET generation in order to maximise the ON current \(I_{on}\) minimising the short-channel effects the drain-induced barrier lowering (DIBL), and the off-current. Preserving the electrostatic integrity has allowed scaling to progress by reducing gate length, gate width, oxide thickness and improving the drain current. The constraints limiting the scaling include thermodynamic constraints restricting active doping concentration in source and drain; physical constraints due to SCE, tunnelling through gate oxide, band-to-band tunnelling and GIDL; statistical constraints which are associated with transistor parameter fluctuation due to discreteness of charge and matter; and economic constraints associated with the manufacturing costs per transistor.

The limits of conventional bulk MOSFET scaling including dimensions and voltage scaling have been reached at the 28/20nm bulk CMOS technology generations. Researchers and manufacturers are now focused on overcoming the scaling limitations and extending the life of Moore’s law. There is potential for employing novel channel materials offering high mobility and potential for higher performance including strained Si, Si/SiGe III-V heterostructures. The OFF-state gate leakage has been minimised by the introduction new gate stacks with high-K dielectric, such as HfO\(_2\) and new gate materials, including metal gates, and fully silicide gate. A second avenue to overcome scaling limitations involves the development of novel device architecture, to include SOI, double gate, trigate, and gate all-
around GAA transistors enhancing the electrostatic integrity and also relaxing the limit of the oxide scaling.

2.6 FinFETs

The fully depleted double gate SOI structure (or delta FET) was initially experimentally suggested by Hisamato et. al. in 1989 [69]. Chenming Hu and his team reported the idea of FinFET in 1999 and UTB-SOI (FD SOI) in 2000 [70]. The fundamental concept underlying both innovations is a very thin body (approximately 10nm or less), enabling the gate to be coupled strongly to the channel with no leakage path a great distance from the gate. As a result, the gate is able to control the leakage current much better. FinFETs possess include the formation of a vertical channel known as the Fin, so the channel width of a FinFET is defined in terms of fin height resulting in channel ‘width quantization’ (single, double or multiple fins). Therefore, the current in the FinFETs can be increased by increasing the fin numbers retaining the improved gate control over the channel charge. Fin height determines the structural stability of the FinFET: the smaller the Fin height the more stable the structure is.

FinFETs can be fabricated on bulk silicon wafers where all fins share a common silicon substrate. In this case shallow trench insolation and punch-through doping are used to isolate the individual fins. In the case of SOI FinFETs the fins are naturally isolated by the buried oxide layer. Figure 2-5 schematically compares bulk and SOI FinFETs.

The first generation of multi-gate transistors with three-sided gate control has been manufactured by Intel at their 22nm node with 90nm fin pitch and 34nm fin height featuring a 3rd-generation high-k metal-gate technology and a 5th generation of channel strain techniques. The lower threshold voltage combined with the strain enhancement have lead to a 13% and 27% increase in \( I_{d_{\text{sat}}} \) in N and P channel FinFETs respectively at 0.8V and 10nA/um leakage current compared with the 28nm CMOS technology [71], [72].
The fin width-height ratio determines the robustness of the FinFET structure. In order to create more robust device, the fin Si has been etched with multiple angles to construct the first FinFET generation shown on Figure 2-6. The non-rectangular fin structure leads to non-uniform mobile charges distribution and somewhat compromises the device performance. The height and angle of the fin of the first generation FinFET have been considered as two of the most important parameters in fabrication optimizing processes [73]. However, the fabrication of the fin leads to significant challenges with respect to gate length scaling.

The second generation of FinFET in the 14 nm CMOS node of Intel reduced the fin pitch and gate pitch by 0.7 (42nm, 70nm respectively). The $I_{\text{dsat}}$ improvement is 15% for NMOS and 41% for PMOS compared to the Intel 22nm technology [71]. The height of fin has been increased from 34nm to 42nm [74].

The possibility of improving the performance of the second generation FinFET is reported in [75] which records the first 14nm strained Ge P-channel FinFETs offering higher performance P-channel transistors while sharing the same technology platform. This PMOS FinFET with 45nm and 100nm fin and gate pitches respectively offers $I_{\text{sat}}=1.2\text{mA/um}$. However, replacing the Si with high mobility materials is not without challenges [76]: The first challenge is the trade-off between the transistor performance and the OFF state current. Secondly the scalability should be economically affordable and co-integration with Si

![Figure 2-5 Three-dimensional schematic representation for the SOI and bulk FinFET.](image)
transistors will be necessary, therefore these materials need to be integrated on a Silicon platform. The mismatch between Si and Ge is approximately 4%, and for III-V is approximately 12% in the case of InAs. Therefore, achieving active semiconductors layers with significant low defectivity is one of the important concerns. Finally, finding a native gate stack with high-k dielectric and low interface defects could be something of a show-stopper [77].

![Figure 2-6 Three-dimensional schematic representation for the SOI FinFET. The fin diverges from the vertical by 8° [73].](image)

Since fin-width scaling below 8nm is critical [74], Zheng et. al. proposed the inserted-oxide FinFET (also called iFinFET) to facilitate gate-length scaling below 10 nm while mitigating the need to form very-high fin aspect-ratio (>10:1 height: width) fin [78]. Since the benefits of the iFinFET stem from the improved capacitance coupling between the gate and the segmented channel regions, it is beneficial to consider high-k oxide materials like HfO$_2$ or Si$_3$N$_4$ as inserted iFinFET dielectrics. The drawback of inserting higher-k material to form iFinFET is the higher capacitance between the channel, drain, and source. As a result, the channel regions have compromised electrostatic integrity.

Scallop-shaped FinFETs (S-FinFETs) are another structural modification merging the advantages of the conventional Si GAA NWT and FinFET. The proposed S-FinFETs demonstrate superior electrostatic integrity in the channels compared with classic bulk-Si FinFETs or tri-gate FETs due to the configuration of quasi-surrounding gate electrodes on
scalloping fins by a specific Si etching process. S-FinFETs could extend general FinFET technology into the sub-10-nm node [79]. Although, S-FinFETs improve the drain-induced barrier lowering and subthreshold swing, they do not yield a significant improvement to the drain saturation current - \( I_{\text{dsat}} \).

The contact all around CAA T-FinFET structure has been proposed for 10nm CMOS. It has better device performance including immunity from SCEs, bulk stressor selectivity, lower source/drain resistance and is very effective in series resistance reduction. However, controlling process variability of such strictures is challenging for 10nm CMOS technology and beyond [80].

The 10nm Intel CMOS technology features in the 3rd generation of FinFET and the 7th generation of strained silicon. The 10nm FinFET technology features rectangular 7nm x 46nm fin shape. An important innovation of the 10nm FinFET technology is the low dielectric constant of the spacers that leads to approximately a 10% reduction in parasitic capacitances. The reduction of the contact resistance together with the reduction of the gate pitch provides boost to the saturation current [55].

Scaling down the fin width beyond 7nm is expected to improve the gate control. X. He et al. [81] presented an experimental study and a simulation study of the impact of the ultimate fin width scaling up to 1.6 nm on the FinFET performance. The results show improvement in DIBL and SS due to better gate control. However, the DIBL and SS decrease with fin width until it reaches a critical point (\( W_c \approx 4\text{nm} \)). When fin width becomes less than 4nm the DIBL and the SS deteriorates due to punch-through in the region below the fin. The FinFET performance (\( I_{\text{on}}/I_{\text{off}} \)) retreated when the fin width becomes less than 5nm as shown in Figure 2-7.
Figure 2-7 Impact of fin width on FinFET $I_{on}/I_{off}$ performance. TCAD simulations have been shown that DIBL, SS, and gate length ($L_G$) are optimised at $W_c\approx 4\text{nm}$. This critical fin width has been selected as reference point refer to (100% performance). Here, performance $I_{on}/I_{off}$ is improved as the fin width is scaled up to 5.5nm but deteriorates swiftly with fin width [81].
2.7 Nanowire transistor (NWT)

The relentless pursuit of high-performance and low-power transistors with ever increasing integration density has driven CMOS technology to the ultimate nanoscale dimensions. Table 2-1 highlights the relations between Si thickness for a given gate length (Lg) with acceptable electrostatic integrity for a range of structures including planar ultra-thin-body SOI, double-gate FinFET, tri-gate FET and gate-all-around nanowire transistors.

<table>
<thead>
<tr>
<th>Structure</th>
<th>Silicon thickens</th>
</tr>
</thead>
<tbody>
<tr>
<td>planar ultra-thin-body SOI</td>
<td>$\cong \frac{1}{3} L_g$</td>
</tr>
<tr>
<td>double-gate FinFET</td>
<td>$\cong \frac{2}{3} L_g$</td>
</tr>
<tr>
<td>Tri-gate FET FinFET</td>
<td>$\cong L_g$</td>
</tr>
<tr>
<td>gate-all around nanowire FET</td>
<td>$\leq 2 L_g$</td>
</tr>
</tbody>
</table>

Table 2-1 The association between Si thickness for a given gate length (Lg) and acceptable electrostatics for a range of Si MOSFET structures

Multi-gate MOSFET have superior electrostatic integrity compared with conventional planar bulk MOSFETs providing a way forward for extended transistor scaling delivering higher performance, lower supply voltage and reduced threshold voltage $V_T$ variability due to tolerance to low channel doping. Amongst all multi-gate CMOS transistors, the silicon nanowire transistor (NWT) provides the route to ultimate CMOS scaling based on its superior electrostatic integrity. These silicon-based NWTs, have an additional benefit - silicon was and has remained the workhorse of the semiconductor industry over more than forty years. It is, therefore, a very attractive candidate for extending the CMOS scaling at or beyond the 5 nm CMOS technology. The gate-all-around (GAA) structure offers optimum capacitive coupling between the gate and the channel allowing the scaling to channel lengths that are unattainable for the alternative single and multi-channel transistor architectures [66].
Nanowire transistors (NWT) could be faster than traditional bulk, SOI and FinFET transistors and could be scaled to smaller channel lengths. NWTs are in an active stage of research and development by semiconductor manufacturers and research groups around the globe as promising candidates for sustaining Moore’s Law. A variety of factors have led to the explosion of research into NWTs. Firstly, semiconductor nanowires can easily be manufactured on a large scale with reproducible electronic characteristics needed for large-scale integrated systems. Secondly, compared with “top–down” nanofabricated device architectures, “bottom–up” synthesised compounds nanowires provide good control for at least one of the critical device dimensions, the channel width, which is at or beyond the limits of the conventional lithography. Furthermore, the crystalline structure, smooth surfaces and the capability of producing radial and axial nanowire heterostructures can reduce scattering and permit higher carrier mobility, when compared with nanofabricated samples of similar dimensions. Lastly, because the nanowire body diameter can be controlled well below 4 nm, the electrical integrity of nanowire-based electronics can be sustained even if the gate length is aggressively scaled, an accomplishment that has become increasingly difficult for conventional top-down fabricated FETs.

Figure 2-8 A Uniform NWT with four elliptical cross-sections (18.8nm×20.2nm), (16.5nm×13.8nm), (10.2nm×10.6nm), and (6.3nm×5.0nm) [82].
Bangsaruntip et al. have developed a fabrication process to fabricate a highly uniform NWT with an elliptical cross-section (see Figure 2-8). Several NWTs with different cross-sectional areas (190π-15.75π) have been made in order to find the impact of NWT diameter on SCE [82]. Characterisation results have shown that a good electrostatic control and satisfactory drive currents, for example the drive current of an elliptical NWT with the cross-section (9nm×13.6nm) is 0.95 mA/μm. There is a significant dependency of the SCE against wire size [82].

Over the past three years many scenarios have been proposed to scale NWTs. R. Kim et al., present an NWT performance guideline based on atomistic quantum transport simulation including strain effects for Ge, Si, GaAs, and InAs NWT with a 13nm gate length. Results have shown that a Ge NWT with improved source and drain design could deliver better current versus delay performance whilst III-V NWTs have shown themselves to be superior for power reduction capacitance [83]. Y. Lee et al., presented an experimental way to scale the Ge and Ge_{0.9}Si_{0.1} NWT with an unusual diamond-shape [84]. An experimental study endorses simulations on industry-related solutions for FinFET and NWT based on different semiconductors such as Si, Ge, and III-V. A substantial impact on self-heating effects was observed by converting from FinFET to NWT. However, modifying the S/D doping or the material of the gate had a little impact on self-heating effects [85]. Another comprehensive study based on state-of-the-art physical models has examined the performance, reliability limits, and variability of 10 nm technology and beyond based on NWT. There have been indication of degradation of ON current of NWT technology compared to FinFETs [86]. Bangsaruntip et al. (IBM) have demonstrated for the first time that Si NWT can be integrated to density targets equivalent to CMOS scaling target of the 10 nm technology. This scaled NWT achieves a good performance (The I_{on} at V_{DD} = 1.0 V 0.976 mA/μm) [87]. The perpendicular stacked NWT architecture, designed for sub-7nm CMOS technology, can offer considerable benefits such as low off-state current thanks to gate all around electrostatic control, whilst this structure drive high on-state current thanks to 3-D vertically stacked channels [88]. The stacked NWT structure has been utilised to construct a functioning ring oscillator [89]. Nanosheet devices are a special case of NWT considered at
the 7nm design rules as a replacement for FinFETs, enabling further scaling down to the 5nm and 3nm technology nodes.

With a relaxed pitch, it is possible to match the effective width of ultra-scaled FinFETs leading, however to a 30% increase in width when broad nanosheets were considered. The vertical stacked nanosheets offer flexible design choices for performance and management of the power due to the acceptable $W_{\text{eff}}$. It has excellent electrostatics and dynamic performance compared with aggressively scaled FinFETs [12]. Figure 2-9 compares the 2D cross-section of a FinFET, lateral stacked NWT, and vertically stacked nanosheets.

Nanowire FETs are fabricable in two configurations; horizontal and vertical. The horizontal NWTs configuration is utilised in plain 2D layouts similar to FinFETs in which the available space for contact and gate placement will be very tight. Therefore, the scaling the NWTs beyond 3nm is a major challenge, as in the end the horizontal configuration will reach physical limits. The vertical configuration of NWTs turns the layout configuration from a 2D to a 3D layout (see Figure 2-10). Here, the gate length can be extended without
occupying a wide area on the wafer. This technology requires intensive research of process design co-optimization.

Figure 2-10 Schematics of vertical NWTs with an array of 3×3 NWTs.

2.8 Statistical Variability

One major challenge with the further scaling of traditional MOSFET is the statistical variability in the transistor characteristics associated with the discreteness of charge and granularity of matter. Transistor sizes are now quantifiable in atomic-scale units and, consequently, self-averaging of atomic scale fluctuations and imperfections in the transistor structure is no longer occurs. MOSFET manufacturing techniques cannot be controlled accurately on atomic scales, resulting for example in a random number and position of individual dopant atoms controlling the transistor performance. This means that variability in MOSFET performance happens as a consequence of factors such as random dopant placement, atomic scale interface roughness and gate morphology, discrete charges at the interface and in the gate oxides and others. This is highlighted by the fact that, merely tens of atoms within the channel determine the responses of a sub-30 nm gate length MOSFET.

Off-state current (leakage current) and $V_T$ variability have been significantly enhanced by the introduction of high-k metal gates. Introduction of these new materials leads to additional variability sources which should also be taken into account. These include variations in the high-k composition and metal gate granularity resulting in local variations
of the gate work function. Intrinsic parameter fluctuations already represent a significant challenge for the semiconductor industry and the forecast is that they will continue to be an important issue in the future.

Both systematic and statistical sources of variation are present at the device level. Systematic variation represents a sub-class of process variability, which is layout dependent, and is a result of strain and lithography including proximity effects, (e.g. imperfections in lithography processes) which are predominantly deterministic and predictable, however complex. Consequently, they are controllable to a certain extent with compensation techniques available at the circuit design stage. As an example, distortions which take place during the photo-lithography process can be corrected by employment of the Optical Proximity Correction (OPC). Furthermore, strain, which is commonly employed to improve transistor performance, leads to both microscopic statistical variations and larger scale deterministic variations which are predominantly due to the effects of the circuitry layout. In this instance, the layout dependent variations could be more significant than the local fluctuations and consequently, strain can be effectively regarded as a systematic variability source and its impact could be examined using TCAD tools. A more serious challenge, however, are entirely statistical intrinsic parameter fluctuations and their inherent stochasticity. Statistical fluctuations are caused by the fundamental discreteness of charge and granularity of matter which are impossible to accurately control (e.g. the arrangement of dopant atoms in a device). These variability sources are responsible for in excess of 50% of the total variability in the 45/40 nm technology generation and have a much more pronounced effect at the 32nm technology. It is therefore clear that statistical fluctuations are of key significance to CMOS scaling and integration in the future.

Random discrete dopants (RDDs) in the channel and source/drain regions are the predominant source of statistical variability in modern bulk MOSFETs, in the 45/40 nm and 32/28 nm technology generations. Whilst RDDs are the predominant source of statistical variability, the contribution of line edge roughness (LER) increases in significance because LER scaling currently lags behind ITRS requirements. Whilst innovative device architectures such as SOI and multi-gate MOS FETs tolerate low channel doping, resulting in reduced RDD variability, they are liable to LER effects. Line edge roughness is a
consequence of the molecular morphology of the photo-resist employed in the lithography process. Its primary influence on MOSFET operation is connected with local variations in the gate length where the transistor has non-regular gate edges along the channel width. The significance of LER will increase as a source of variability as transistor dimensions decrease. LER introduces serious variability in subthreshold current and DIBL in addition to the threshold voltage variability. A further concern with LER is the degradation in $I_{on}/I_{off}$ ratio, resulting from enhanced short channel effects which can lead to significant degradation of both device and circuit output.

Metal Gate Granularity (MGG) is an issue mainly connected with the “gate first” process technology, whereby the gate metal is deposited prior to any high temperature annealing procedure. During high temperature processing, the nominally amorphous metal gate material becomes poly-crystalline. This results in the formation of grains with different crystallographic orientation and differing metal work function. Within the case of polysilicon gates, the interfaces between grains cause Fermi level pinning and doping non-uniformity as a consequence of rapid diffusion along grain boundaries. Both effects introduce significant variability in the performance of the devices. The effect of MGG on device performance is closely correlated with the metal grain size, in respect to the overall gate size. In the case of metal gate, large grain size with respect to overall gate size, has a bimodal effect on transistor performance, whereby differing device instances are controlled by

![Figure 2-11](image-url) *Figure 2-11 The main statistical variability sources (RDD, MGG, and LER) in Si NWT.*
different grain work functions. Interestingly, small grain size, relative to overall gate area, causes self-averaging of the grain work functions and has a smaller influence on overall device performance. In the gate last processes, where the gate is deposited post annealing (and other high-temperature processes), the gate granularity could be significantly reduced. Figure 2-11 shows the fluctuation of the electric potential of three stacked NWTs suffering from the main sources of statistical variability.

2.9 Summary

This chapter describes the challenges that need to be addressed in the scaling of conventional MOSFETs and presents the technological innovations that have been proposed. The theory of MOSFET scaling is also outlined.

The relentless requirement for high-performance and low power devices with density of integration drives CMOS technology to ultimate nanoscale dimensions. Nanowire MOSFETs are among the candidates to extend CMOS downscaling to ultimate limits, eventually replacing the triple gate FinFET architectures after the 7nm CMOS technology generation. The nanowire gate-all-around configuration has the best immunity against short channel effects (SCE) allowing ultimate transistor scaling. Since statistical variability is becoming a serious scaling constraint factor, the sources of statistical variability have been discussed in detail.

To summarise, the fast progression in nanofabrication technology has created possibilities for the applications of silicon nanowire transistors in the future CMOS technology generations. As a result, more profound understanding of NWT physics and the development of TCAD (Technology Computer Aided Design) tools for NWT design is becoming increasingly important. The aim of this thesis is to comprehensively investigate TCAD simulations in order to examine the physics of silicon nanowire transistor devices. With the advanced modelling tools employed in this study, the ultimate performance limits of SNWTs will be evaluated and important challenges in NWT device design will be addressed.
Chapter 3. Simulation Methodology

3.1 Introduction

Semiconductor device design and optimisation depend greatly on the accurate simulation of the device’s physical behaviour and electrical characteristics. By modelling a semiconductor device’s behaviour as part of the CMOS Technology Computer Aided Design (TCAD) process, we can assess its key parameters and operation and understand how to improve its functionality and performance. Evaluating device performance using back of an envelope calculation becomes practically impossible with the scaling-down of the devices, due to the three-dimensional structure and the sophisticated physics mechanisms governing the nanoscale device operation. In order to effectively assess the operation of 3D MOSFETs, it is essential to carry out 3D numerical simulation of the device’s characteristics and behaviours. The modelling of new CMOS transistors has been performed using a number of different approaches, such as Nonequilibrium Green Functions (NEGF), Monte-Carlo (MC), hydrodynamic, and drift-diffusion (DD) methods. However, since the computational intensity associated with the physically more accurate methods is extremely high, the choice of simulation techniques is always a trade-off between accuracy and computational efficiency. Therefore, a compromise the solution will be to calibrate relatively simple, computationally efficient simulation such as DD on models provide an accurate physical information on the quantum mechanical phenomena and can be appropriately deal with non-equilibrium transports occurring in the device.

In this study the self-consistent Poisson- Schrödinger solver was adopted for studying the impact of the quantum confinement on the mobile charge and capacitance of NWTs with different cross section. Although the PS quantum corrections provide very accurate quantum charge distribution in the channel of the simulated NWTs, the large number of cross-sectional solutions of the Schrödinger equation significantly slows the simulations and reducing efficiency and productivity. Therefore, we calibrate the DG quantum corrections to the PS charge distribution and then use the DG simulations Figure 3-1 shows a simplified
calibration flowchart of DD to Poisson- Schrödinger simulation. This approach has been widely employed in chapter 4 to investigate the quantum confinement effect. We also employed the 3D ensemble MC simulation approach with accurate quantum corrections for the predictive simulation of nanowire transistor (NWT) charge transport. For the simulation of large multichannel NWTs and for the simulation of statistical variability we use drift diffusion simulations thoroughly calibrated to the results of the Monte Carlo simulations. Figure 3-2 presents an overview of the calibration flowchart of DD to quantum corrected 3D Monte Carlo simulation. This approach has been widely employed to investigate performance and statistical variability of NWTs in chapter 5 and chapter 6 respectively.

The following section of this chapter (section 3-2) will concentrate on basic aspects of classical DD approach and its unsuitability to deal with physical phenomena occurs at in NWTs.

Section 3-4 present relevant modifications of the basic DD, namely, density gradient calibrations. The detailed description of Schrödinger and Monte Carlo is presented in sections 3-3 and 3-5 respectively. Further improvement to DD can be achieved by a large variety of mobility models. Section 3-7 illustrates the most efficient mobility models and how can be used to calibrate DD to accurate models such as quantum corrected Monti Carlo. The conclusions are drawn in the final Section.
Figure 3-1 The calibration flowchart of DD to Poisson-Schrödinger simulation.

Figure 3-2 The calibration flowchart of DD to quantum corrected 3D Monte Carlo.
3.2 Drift Diffusion

Near equilibrium device physics and behaviour are primarily modelled using the drift diffusion (DD) approach. The DD method is suitable when the devices are of an adequate size to discount quantum effects, and for long-channel devices that are not rapidly switched. Whilst one of the main benefits of the model is its straightforward implementation, resulting in widespread adoption, it is incapable of capturing nonequilibrium transport effects. The DD method works based on the assumption that carriers are in thermal equilibrium, and although field-dependent diffusivities and mobilities can be used, the assumption is that these parameters can react immediately to any electric field fluctuation. This assumption is inaccurate for smaller devices, wherein the transport variables are not directly correlated to the local electric field. Additionally, the classic DD method cannot capture velocity overshoot [90] and other non-local effects. Furthermore, the DD model can provide highly inaccurate representations of charge velocity distribution, potential configuration, and charge density in aggressively scaled transistors. Thus, the DD approach is often unsuitable for exploring the in-depth physics of short-channel devices during simulation [91], [92].

The DD method is based the Poisson’s equation (a fundamental electrostatics equation based on the Maxwell equation), current transport equations for electrons and holes in drift-diffusion approximation (derived from Boltzmann’s transport equation) implemented in the current continuity equation. The above equations are solved self-consistently using numerical techniques. The DD equations are outlined in further detail in the following sections.

3.2.1 Maxwell's equations

The differential-form of Maxwell's equations are;

I. Gauss's law of electrostatics:

\[ \nabla \cdot D = \rho , \quad \nabla \cdot E = -\frac{\rho}{\varepsilon_0} \]  \hspace{1cm} (3-1)

II. Gauss's law for magnetism:

\[ \nabla \cdot B = 0 \]  \hspace{1cm} (3-2)

III. Faraday's law of induction:
\[ \nabla \times E = -\frac{\partial B}{\partial t} \]  

(3-3)

IV. Ampère's circuital law:

\[ \nabla \times E = \varepsilon_o \left( J + \varepsilon_o \frac{\partial E}{\partial t} \right) \]  

(3-4)

The wave equation is one of the key equations used in the field of electromagnetics. The wave equation is derived from Ampere’s law and Faraday’s law, demonstrating that all waves travel at the speed of light:

\[ \nabla^2 E = \mu \varepsilon \frac{\partial^2 E}{\partial t^2} \]  

(3-5)

The wave equation expresses the relationship between electrostatic potential and space charge, or between a given electron density distribution and the relevant electric field. Here, \( E \) is vector. For instance, we can say that field \( E \) is travelling in \( z \)-direction, with no variation in the \( x \)- and \( y \)-direction (i.e., zero partial derivatives in the \( z \)- and \( y \)-direction). In this case, Equation (3-5) can be reduced to a simplified scaler wave equation:

\[ \nabla^2 E_x = \frac{\partial^2 E_x}{\partial z^2} = \mu \varepsilon \frac{\partial^2 E_x}{\partial t^2} \]  

(3-6)

Equation (3-6) is transferable to \( f(z + ct) \) form where be simplified by “chain rule”

\[ c_o = \frac{1}{\sqrt{\mu \varepsilon}} \]  

(3-7)

\[ \lambda = \frac{c}{f} \]  

(3-8)

A quasi-stationary condition is assumed for the electric field given that standard device dimensions are significantly lower than \( \lambda \). This can be expressed in the form of a scalar potential field gradient:
3.2.2 Poisson’s equation

Poisson’s equation is derived from the aforementioned Gauss’s law of electrostatics in combination with Equation (3-9):

$$\nabla^2 \psi = -\frac{\rho}{\varepsilon_0 \varepsilon_r} \quad (3-10)$$

In semiconductors, space charge density $\rho$ consists of mobile charges (determined by electrons and holes) and fixed charges (determined by ionized donors and acceptors):

$$\nabla^2 \psi = -\frac{q}{\varepsilon_0 \varepsilon_r} (p - n + N_D^+ - N_A^-) \quad (3-11)$$

where $\rho$ is charge density; $\varepsilon$ is permittivity; $\psi$ is electrical potential; $p$ and $n$ are electron and hole densities, respectively; and $N_D^+$ and $N_A^-$ are donor and acceptor densities. If only electrons, are considered in unipolar MOSFET operation Poisson’s equation is presented as follows:

$$\nabla^2 \psi = -\frac{1}{\varepsilon} (-n + N_D^+ - N_A^-) \quad (3-12)$$

The Poisson’s equation is non-linear as a result of the carrier concentrations’ dependency on electrostatic potential.
3.2.3 **Current continuity equations**

The following equation is derived based on the divergence of Ampere’s circuital law:

\[
\nabla \cdot (\nabla \times H) = \nabla \cdot J + \nabla \frac{\partial D}{\partial t} = 0 \tag{3-13}
\]

In semiconductors, the sum of electron and hole current densities, \(J_n\) and \(J_p\), produce total current density \(J\):

\[
\nabla J_n - q \frac{\partial n}{\partial t} = qR \tag{3-14}
\]

\[
\nabla J_p + q \frac{\partial p}{\partial t} = -qR \tag{3-15}
\]

The electron dynamics can be expressed using transport equations, such as the Boltzmann transport equation (BTE), which capture the dynamics of electron density distribution resulting from driving forces such as the electron density gradient or external electric field. In drift-diffusion approximation the electron current density is given by

\[
J_n = -q n \mu_n \nabla \Psi + q D_n \nabla n \tag{3-16}
\]

where \(J_n\) is electron current density, \(n\) is electron density, \(\mu_n\) is electron mobility, and \(D_n\) is the electron diffusion coefficient.

Maxwell’s equations can be used to obtain the continuity equation, addressing carrier generation, recombination and other time-based phenomena. It is effectively an equation that expresses current conservation, as a key feature of nature:
\[
\frac{\partial n}{\partial t} = \frac{1}{q} \nabla J_n - R_n + G_n
\]  

(3-17)

where \( G_n \) is the electron generation rate, and \( R_n \) is the recombination rate. When generation or recombination are absent, \( \nabla J_n = 0 \), denoting a constant current density in the case of a constant cross-section region.

3.2.4 Transport equations

The classical transport description in semiconductor devices is based on the Boltzmann transport equation (BTE). BTE is an integral-differential equation derived from classical mechanics and statistical dynamics laws. This is represented as in [93]:

\[
\frac{\partial f}{\partial t} + v \cdot \nabla_r f + qE \cdot \nabla_p f = \left( \frac{\partial f}{\partial t} \right)_{\text{coll}}
\]  

(3-18)

where \( f(r, p, t) \) is the single particle distribution function, \( v \) is the group velocity of electrons, and \( E \) is the applied electric field.

\[
\mu = q \langle \tau \rangle m^{-1}
\]  

(3-19)

As shown in (3-19) the mobility value is evidently impacted directly by the \( \langle \tau \rangle \) value, where \( \langle \tau \rangle \) is the average momentum relaxation time, and \( m \) is the effective mass tensor. Therefore, mobility can be estimated more efficiently when scattering mechanisms are captured. A diagonal effective mass tensor with even diagonal elements is seen in Si. Thus, a scalar \( \mu \) is used to express mobility.

Electron motion can be determined by numerous scattering mechanisms in a real Si sample. Matthiessen’s rule can be adopted to combine the mobilities associated with different
scattering mechanisms based on the assumption that the scattering mechanisms are statistically independent:

\[
\frac{1}{\tau} = \frac{1}{\tau_1} + \frac{1}{\tau_2} + \ldots + \frac{1}{\tau_n}
\]  

(3-20)

where \( n \) is the independent scattering mechanisms. Thus, the following equation expresses overall mobility:

\[
\frac{1}{\mu} = \frac{m^*}{q\tau} = \frac{1}{\mu_1} + \frac{1}{\mu_2} + \ldots + \frac{1}{\mu_n}
\]  

(3-21)

A gradient of material properties, temperature or carrier concentrations can result in carrier transport in semiconductors, together with the application of an electric field. The following simplification of the BTE in is adopted in order to arrive at an approximated expression for the current density, in response of concentration gradients and electric field. The BTE collision term (on the right side) can be expressed using the following equation based on the relaxation time approximation (RTA):

\[
\left( \frac{\partial f}{\partial t} \right)_{\text{coll}} = \frac{f - f_0}{\tau}
\]  

(3-22)

where \((f_0)\) denotes the equilibrium distribution function. The following equation can be used to express 1D BTE under the assumption of steady state conditions:

\[
v_{xc} \frac{\partial f}{\partial t} + \frac{qE}{m} \frac{\partial f}{\partial v_x} = \frac{f - f_0}{\tau}
\]  

(3-23)
The following equation is derived through the multiplication of (3-23) with $v_x$, integrated across the 3D velocity space:

$$\int v_x^2 \frac{\partial f}{\partial x} d^3v + \frac{qE}{m} \int v_x \frac{\partial f}{\partial v_x} d^3v = \frac{\int v_x f_0 d^3v \int v_x f d^3v}{\tau}$$  \hspace{1cm} (3-24)

The right-hand side of the BTE is then expressed as follows, with the first integral on the right-hand side disappearing as a result of the symmetrical equilibrium function [94]:

$$J_x = -q \int v_x f d^3v$$  \hspace{1cm} (3-25)

$$J_x = q \frac{q \tau E}{m} \int v_x \frac{\partial f}{\partial v_x} d^3v - q \tau \frac{d}{dx} \int v_x^2 f d^3v$$  \hspace{1cm} (3-26)

$$\int dv_y \int dv_z \int v_x \frac{\partial f}{\partial v_x} dv_x$$

$$= \int dv_y \int dv_z [v_x f]_{-\infty}^{+\infty} - \int f d^3v$$  \hspace{1cm} (3-27)

$$= -n \int v_x^2 f d^3v = n(v_x^2)$$

Current density is expressed as follows, with mobility $\mu$ and the $\langle v_x^2 \rangle = \frac{kBT}{m}$ average value incorporated:

$$J_n = q \mu_n \left[ nE + \frac{kT}{q} \nabla n \right]$$  \hspace{1cm} (3-28)

Hole current density is expressed similarly:
\[
J_p = -q\mu_p \left[ pE - \frac{kT}{q} \nabla p \right]
\] (3-29)

The essential DD-based simulation equations are then assembled including the Poisson equation (3-12), the current continuity equations (3-17), and the current expression in drift-diffusion approximation (3-28) and (3-29):

\[
J_n = J_{\text{diffusion}} + J_{\text{drift}} = qD_n \nabla n - q\mu n \nabla \psi
\] (3-30)

\[
J_p = J_{\text{diffusion}} + J_{\text{drift}} = qD_p \nabla p - q\mu p \nabla \psi
\] (3-31)

where \(\mu_n\) and \(\mu_p\) are electron and hole mobility, respectively, and \(D_n\) and \(D_p\) are the diffusion coefficients relevant to each type of carriers. The Einstein relation links carrier mobility with the carrier diffusion coefficient or near to thermal equilibrium. In the case of nondegenerate semiconductors, \(D = \mu \frac{kT}{q}\): where \(k\) is Boltzmann’s constant, \(T\) is absolute temperature, and \(q\) is single electron charge.

In summary the semiconductors’ electrical behaviour is effectively described through the self-consistent solution of the DD set of equations including the Poisson’s equation and the current continuity equation in drift-diffusion (DD) approximation. In the next sub-section (3-2-5) we will discuss how the above equations are consequently solved. However, nanoscale MOSFETs demands for the inclusion of quantum effects and as the device dimension shrinkage, classical DD cannot be appropriately capture quantum effects without corrections [95]. The classical DD does not deal with non-equilibrium transport.
3.2.5 **Self-consistent calculation based on the Gummel’s method**

In most cases, electron device simulation requires the solution of a self-consistent system of equations describing the devices’ charge distribution and the electrostatic potential $\psi$. Self-consistency occurs when flow of charge in the device is in equilibrium with the potential distribution in the device, which in part is affected by the moving charges. Here, charge density $\rho$ adjusts when $\psi$ is altered which in turns changes the potential, ‘self-consistency’ is reached when the charge and the potential reach equilibrium condition.

The achievement of self-consistency is essential for correctly obtaining the simulated devices’ field distribution and current flow. In order to achieve this, we solve Poisson’s equation and the current continuity equation simultaneously, which enables to determine the correct mobile charge distribution and corresponding current (3-11).

The numerical solution of the DD set of equation is based on the self-consistent solution of the electron density ($n$) and the electrostatic potential ($\psi$) as two unknown variables. Both variables are position dependent. In the case of a MOSFET in absence of generation and recombination the current continuity equation results in constant current density ($J$) along the channel. The self-consistency in the Gummel approach is obtained by alternated solutions of the Poisson and the current continuity equation. Typically, the solution for the electron density $n$ from the current continuity equation is based on the previous solution for the electrostatic potential $\psi$. An updated $\psi$ distribution is than obtained by applying the new carrier concentration distribution $n$ when solving again the Poisson’s equation. Convergence is achieved if the new $\psi$ distribution is sufficiently close to the previous $\psi$ distribution. If not, the iterations between the solution of the current continuity equation and the Poisson equation continue. [96], [92].

The electrical characteristics of a single CMOS transistor can be simulated using the GSS TCAD device simulator, GARAND. This is a DD simulator based on finite volume discretisation of the Poisson and the current continuity equations. The self-consistency in Garand is achieved using Gummel iterations (Figure 3-3). Additionally, GARAND also considers the quantum mechanical confinement effects using the density gradient quantum
corrections approach. GARAND is a 3D simulator and arbitrary-shaped 3D MOSFET structures can be simulated using GARAND.

3.2.6 Limitations of classical Drift-Diffusion approach

The DD approach has been adopted widely by the industry and was accurate in the simulation of relatively large devices in above 100nm technology generations. The main benefits of the DD approach in 2D and 3D simulation environment, its simplicity and its computational efficiency. The DD method assumes local relation between carrier velocity and the electric field, incapable to describe non-local effects based on the fact that the velocity and energy at particular point depends on the field distribution along the particle trajectory leading to the current position [90]. The DD method applied to contemporary and future CMOS transistors is most suitable in the subthreshold region of the device operation where the electrostatics plays a dominant role, and where there is a weak coupling between the current continuity and the Poisson equation solution. Here, the channel contains only a small amount of mobile charge due to the high source-to-drain potential barrier. Gate voltage, drain voltage together with line-edge roughness (LER), interface roughness, differences in dopant placement, and other barrier fluctuation sources determine the barrier landscape [97]–[99]. It is well understood that DD is inaccurate in capturing on-current ($I_{on}$) and the on-current variability in contemporary MOSFETS above the threshold level [100].

The assumptions that form the basis of DD equations are violated as the devices are scaled to nanometre dimensions, resulting in a non-equilibrium and non-local transport problem. Here, prior scattering along the carrier trajectories must be considered. The traditional DD approach assumes that the carriers are in thermal equilibrium with the crystal lattice, discounting the carrier heating and the non-equilibrium transport. The hot carriers arise due to an increase in the electric field caused by the device scaling. Therefore, the DD approach runs out of steam as the devices are scaled to nanometre dimensions. Non-local effects are not represented by the DD simulation, since it only captures instant local velocity modifications related to the local electric field. In small devices, field changes occur quickly, with $\mu_n$ and $D_n$ no longer related directly to the local electric field, and with carrier history now a determining the velocity and the energy at particular point in the simulation domain.
In particular the DD simulation approach does not capture the carrier velocity overshoot which is non-equilibrium transport phenomenon [101], arising as a result of the carriers moving with velocity higher than the saturation velocity in the case of small devices with rapidly varying electric field in the channel. The spatial velocity overshoot is a result of the faster response in average carrier velocity with the average kinetic energy lagging when carriers move from a low field to a high field region in rapidly field changing conditions. Since the DD approach is based on a simple field-dependent mobility model that is connected to the electric field at a local level, the DD approach does not capture the velocity overshoot [102]. Previous research indicates that velocity overshoot has beneficial impact on the transistor performance, increasing drive current, transconductance and reducing circuit switching time [103]. Higher order approximation to the BTE, where thermal equilibrium is not assumed, including the energy transport equations and hydrodynamic set of equations [104] can capture overshoot effects at greater computational expense. The need for more accurate BTE solutions has been highlighted due the deficiencies of the DD and the hydrodynamic approach in the simulation of extremely-scaled devices [104].

With the transistor scaling to nanometre dimensions there is a greater need to represent accurately the quantum effects in the simulations. In particular in NWT FETs, the accurate simulation of mobile charge and gate capacitance require quantum mechanical treatment. Quantum corrections can be introduced into DD simulations. Methods based on the solution of the Schrödinger equation are adopted in the research presented in this thesis. The effective quantum potential extracted from the quantum solution is combined with the classical potential (obtained by solving Poisson’s equation) providing the driving potential that is used in the current continuity equation in the DD simulation, or for determining the particles’ driving force in the MC simulations. A 2D Poisson-Schrödinger solver used in the planes normal to the transport direction to provide the quantum charge distribution and the associated effective quantum potential. Next section (section 3-3) will describe Poisson-Schrödinger simulation.
3.3 Poisson-Schrödinger

The simulation of aggressively-scaled MOSFET requires the accurate inclusion of quantum effects. Contemporary MOSFETs have a channel length close to the mean carrier free path, with an oxide layer approximately three Si lattice interatomic atomic layers thick. In order to improve the electrostatic control in NWTs, the gate completely surrounds the nanowire channel on top of the gate oxide layer. Quantum confinement starts to play important role when the nanowire diameter becomes smaller than 8nm. The self-consistent solution of the Schrödinger and the Poisson equations in the 2D cross sectional plane of the NWT can provide accurate description of the quantisation effects in the NWT channel. The potential well in the NWTs is narrow leading to strong quantisation effects including sub band splitting and affecting the shape of the carrier distribution in the channel, the ground state (the lowest permitted electron energy level in the quantum well) is shifted above the bottom of the conduction band resulting in threshold voltage shift. The 1D quantum transport direction along the channel, surrounded by the gate, is perpendicular to the confinement planes (2D) [105].

The quantum mechanical confinement has a significant impact on charge distribution. In classical DD simulations the maximum of the inversion layer charge is at the interface, with the charge density exponentially decreasing away from the interface. In reality the quantum mechanical charge distribution peaks at a distance away from the interface. In NWTs the quantum mechanical charge distribution results in volume inversion with the current flowing dominantly in the middle of the NWT channel.

In GARAND the first-order quantum mechanical effects are taken into account using density gradient quantum corrections. However, in this study a more sophisticated quantum mechanical model is adopted, based on the self-consistent Poisson-Schrödinger (PS) equation solutions. This offers more accurate description of the quantum mechanical effects, at the expense of lower computational efficiency and greater computational times. Similarly, to the DG simulation approach, the more accurate PS based quantum corrections are implemented in the DD simulations using an iterative convergence scheme. The iterations start with classical DD simulation. Once convergence is achieved, the DD solution is used
to determine the quasi-Fermi level, used as reference in the PS simulations and the initial shape of the confinement potential in each discretisation plane normal to the transport direction. The 2D quantum mechanical charge distribution in each 2D discretisation plane is then obtained by solving self consistently the Poisson-Schrödinger equation in each discretization plane. Figure 3-4 presents a flowchart of this simulation.

A quantum correction term is derived from the quantum mechanical charge distribution obtained from the Schrödinger solution, and this term applied to the current density equation in the next iteration of the DD system solution. The iterations between the DD and the PS solutions are repeated until convergence. Figure 3-5 illustrates the secondary DD loop. The same quantum correction term can also be used in simulations carried out using the Monte Carlo method.

The Poisson and Schrödinger equations are solved in the 2D discretisation plane providing the correct charge distribution and gate capacitance in the channel of the NWTs. The Poisson equation (3-12) in this case includes the doping charge and the quantum mechanical charge distribution acquired from the solution of the Schrödinger equation. The Schrödinger equation in tensor effective mass approximation is given in the form:

\[
-\frac{\hbar^2}{2} \nabla \frac{1}{m_{ij}} \nabla \psi = (\psi - E)\psi 
\]

(3-32)

The wavefunction is set to zero through the default setting of Dirichlet boundary conditions to the outer bounds of the Schrödinger domain. In order to prevent the wavefunction entering metallic regions, metals are also subject to the same boundary condition. The LAPACK libraries’ numerical eigenvalue solver is applied to calculate the eigenvalues and the wave functions. The solutions of the Schrödinger equation provide the wave functions in each valley in the band structure within each of the 2d discretisation cross-sections. The total mobile charge distribution is obtained as a sum of the charge distribution in each calculated from the corresponding wave function distributions. The new potential value is then solved.
3.4 Density gradient calibration

Whilst the accuracy in simulating the NWTs’ quantum charge distribution using non-equilibrium Green’s functions or direct Poisson/Schrödinger equation solutions in cross-sectional discretisation planes is high, this comes at great computational expense, yielding lower productivity, reduced efficiency, and slower simulation speed. Computational efficiency is of great importance when evaluating many design options, and it is therefore essential to use efficient methods capturing first-order quantum mechanical effects in order to support the computer-aided design of upcoming devices. This is particularly important when dealing with statistical variability which requires the 3D simulations of large statistical samples of macroscopically identical but microscopically different transistors.

A further quantum correction term ($\psi_{qm}$), which corresponds to the second-derivative of the carrier density’s square root [7], [106], [105], is introduced by the density gradient formalism in the extended version of the BTE:

$$\frac{\partial f}{\partial t} + v \cdot \nabla_r f + \frac{1}{\hbar} F_{qc} \cdot \nabla_p f = \left( \frac{\partial f}{\partial t} \right)_{coll}$$

where $F_{qc}$ is the quantum correction force derived from the sum of the classical and the quantum correction potentials expressed as:

$$F_{qc} = -\nabla \left( \psi_c - \frac{\hbar^2}{12m^*} \nabla^2 \ln(n) \right)$$

where $\psi_c$ is classical potential, $m^*$ is effective mass, and $n$ is electron concentration. The quantum correction based on the second derivative of the carrier concentration is derived through series expansion and taking into account the lowest-order element. Introducing the new quantum correction to the current expression in DD approximation to the following quantum corrected current equation:
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\[ J_n = q D_n \nabla n - q \mu_n n \nabla \psi + 2 \mu_n \nabla \left( b_n \frac{\nabla^2 \sqrt{n}}{\sqrt{n}} \right) \]  

(3-35)

\[ b_n = \frac{\hbar^2}{4q m r} \]

where \( r \) is a fitting parameter taking values between 1 and 3, based on the number of occupied sub-bands.

In many cases, first-order quantum corrections are implemented through a so-called effective quantum potential. Here, a quantum correction term is applied to adjust the electrostatic potential used in traditional simulation methods. This allows the necessary quantum mechanical effects to be modelled, therefore maintaining the stability of the simulator and its computational efficiency.

As noted, a further the quantum correction term \( \psi_{qm} \), which corresponds to the second-derivative of the carrier density’s square root, can be obtained using the density gradient (DG) formalism. The effective quantum potential \( \psi_{qm} \) is related to the quasi-Fermi level \( \Phi_n \) by the following equation:

\[ \psi_{qm} = 2 b_n \frac{\nabla^2 \sqrt{n}}{\sqrt{n}} = \Phi_n - \Psi + \frac{k_B T}{q} \ln \left( \frac{n}{n_i} \right) \]  

(3-36)

\[ b_n = \frac{\hbar^2}{12q m n} \]

where \( b_n \) is a parameter, which can be calibrated to replicate the electron distribution obtained usually from self-consistent Poisson-Schrödinger (PS) simulations. Figure 3-3 presents a standard DD simulation flowchart incorporating density gradient quantum corrections.
Effective mass is used as a fitting parameter, with the density gradient equation calibration 3-38 solved numerically and calibrated against the results of Poisson-Schrödinger solution. With the assumption that the density gradient effective mass is anisotropic, with different effective mass components of the principle direction (m_x, m_y and m_z) the DG equation can be written in the following form:

\[
\psi_{qm} = 2b_n \frac{\nabla^2 \sqrt{n}}{\sqrt{n}} = \Phi_n - \psi + \frac{k_B T}{q} \ln \left( \frac{n}{n_i} \right) \\
= \phi_n - \psi + \frac{k_B T}{q} \ln \left( \frac{S^2}{n_i} \right) 
\]  

\[(3-38)\]

where \( b_n = \hbar^2 / 12 q \), and \( S = \sqrt{n} \). The charge distributions derived from the Poisson-Schrödinger solutions along the NWT cross-section’s major and minor axes are used as the basis for calibrating the conduction band density gradient effective masses in the silicon and in the oxide.

To secure charge neutrality being maintaining taking into account the impact of the electrostatic potential on the electron injection, Neumann boundary conditions (NBCs) are adopted in the source and drain when implementing NEGF simulation approach and other quantum mechanical transport methods. Whilst most DD simulators use the Dirichlet boundary conditions for the potential at the ohmic S/D contacts, NBCs are also employed in GARAND to allow for the consistent application of the DG quantum corrections. NBCs are well suited for DG quantum corrections because they allow the correct quantum mechanical charge distribution to be maintained in the source/drain regions. This is especially important
in the case of NWTs, FinFETs, and FDSOI transistors. Where implementations of the DG approach, care should be taken when applying the quantum potential boundary conditions at the interface between the semiconductor and the oxide. The approach described in [107] adopted in GARAND to take into account the wave function penetration in the oxide. Using this approach, the gradient of the electron concentration perpendicular to the semiconductor/oxide interface is influenced by the effective masses in the two media. The calculation of electron density penetration, as a function of distance \( x \) from the semiconductor/oxide interface, is as described by the following expression:

\[
n(x) = n(0)e^{\frac{-2x}{x_p}}
\]  

(3-39)

where \( n(0) \) is electron density at the interface, and \( x_p \) is penetration depth, derived from the Wentzel-Kramers-Brillouin (WKB) equation:

\[
x_p = -\frac{\hbar}{\sqrt{2m_{ox}\Phi_B}}
\]  

(3-40)

where \( m_{ox} \) is the oxide’s electron effective mass, and \( \Phi_B \) is the oxide’s potential barrier. Based on Eqn. (3-37), the component \( b_n \nabla \sqrt{n} \) set normally to the semiconductor/oxide interface can be expressed as:

\[
n.b_n \nabla_p \sqrt{n} = -\left(\frac{b_{ox}}{x_p}\right)\sqrt{n}
\]  

(3-41)

where \( b_{ox} \) is expressed as:
\[ b_{ox} = -\frac{\hbar^2}{12q m_{ox}} \]  

 Normally, density gradient effective masses would be calibrated in respect of the results achieved from NEGF, Poisson-Schrödinger simulations.

The DG quantum corrections are essential when carrying out ‘atomistic’ simulations including random discrete dopants (RDD). In classical DD ‘atomistic’ simulations deep Coulomb well generated by each discrete dopant in the simulation can trap carriers resulting in an increase of the resistance of the RDD region. The trapping of majority carriers results in an artificial reduction of the current. At the same time, the trapping of the minority carrier can affect the shape and the width of the depletion region. In classical DD simulations these effects are dependent on the mesh spacing, and they can be completely eliminated by the incorporation of the density gradient quantum corrections in the DD simulation approach.

The GSS Enigma framework allows automatic calibration of the DG parameters to the solution of the Poisson-Schrödinger equation capturing also the orientation dependence of the DG parameters. Furthermore, artificial S/D tunneling is avoided by setting large density gradient effective mass along the transport direction allowing quantum confinement to be applied correctly in the direction perpendicular to the transport.
Figure 3-3 Simulation flowchart for Drift Diffusion based a modified Gummel algorithm [8].
Figure 3-4 Flow Diagram of the 2D Poisson-Schrödinger model [8].
Figure 3-5 Flow diagram of the fixed Schrödinger quantum correction DD model [8].
3.5 **Monte Carlo**

MC allows different parameterisations for electrons with different energy levels and thus it is more flexible in allowing integration of Poisson-Schrödinger. In this section, we will investigate on normal Monte Carlo first. The Monte Carlo method applied to semiconductor device simulations provides a direct solution to the BTE. It provides and accurate solution for the energy and momentum distribution in the presence of strong external electric field leading to nonequilibrium transport effects. The solutions consider all relevant scattering mechanisms, in a 2D/3D simulation domain which could be a very realistic representation of the simulated device. The Monte Carlo approach simulates the effects of random scattering events and the free flight carrier trajectories in electric field, bridging the classical Newtonian mechanics and quantum mechanical scattering rates to provide a numerical solution to the BTE.

Figure 3-6 illustrates a common simulation flow for an Ensemble Monte Carlo simulation engine. The initialisation of the MC simulations proceeds by introducing MC super particles into the solution domain according to the carrier density distribution obtained from DD simulations. The energy dependent total scattering rate is calculated based on all relevant scattering mechanisms. A random number in connection with the total scattering rate is then used to determine a ‘free flight’ time for each super particle. During the free flight period carriers move following classical lows of motion accelerated by the electric field. At the end of the free flight an additional random number is used to select the scattering mechanism, which is introduced once the free flight period ends. Here, the relevant scattering probability distributions along with the choice of the superparticle state following scattering event enable the device physics to be accurately reproduced by the MC simulations. Self-scattering is also introduced leading to a constant total scattering rate, in order to simplified the flight time selection process. Self-scattering is the most commonly-selected scattering mechanisms in the majority of cases reducing the efficiency of the MC simulations. Statistics are collected after scattering, with average values obtained, for particle velocity (3-43) and energy (3-44) taking into account the nonparabolic band dispersion, described by nonparabolicity factor. Using the expressions below.
\[ v = \frac{\hbar K}{m^* \left( 1 + 2\alpha E(K) \right)} \]  
(3-43)

\[ E(K)(1 + \alpha E(K)) = \frac{\hbar^2 K^2}{2m^*} \]  
(3-44)

Current and other ensemble averages are also obtained. Both self-consistent and frozen field simulation are possible using the Monte Carlo approach. In the former case, Poisson’s equation is solved at regular time steps (3-12), using the electron distribution achieved at the completion of each time step. This provides a new field distribution for the next time step. The process is repeated until convergence is achieved. In the latter case, the field distribution is fixed and based on the results of prior simulation DD-simulations. In terms of computational efficiency in the case of 3D MC, the main challenge is the frequent solution to Poisson’s equation which consumes most of the simulation time. Thus, it is crucial to adopt a highly efficient Poisson solver, such as the black/red successive over relaxation (SOR) method [108], [109] with Chebishev acceleration which is highly efficient in parallel processing environment. The biconjugate gradient-stable (BICGSTAB) METHOD [110], or the multigrid method [111] are also good candidates for accelerating the MC simulations.

Different approximations for the material band structure can be used in the MC simulations. The most straightforward approach is the use of single parabolic or nonparabolic bands [112], with ellipsoidal or spherical shape depending on the tensorial form of the effective mass. A more complex approach is to use full band structure based on k · p, pseudopotential [113]–[115] and tight-binding [116] methods. However, as the description of band structure becomes more sophisticated the computational efficiency is significantly eroded. The Monte Carlo method is superior to the DD method in simulating contemporary and future nano-CMOS transistors with fewer scattering events resulting in near-ballistic and non-equilibrium transport. Previous research confirms the increasing role of the non-equilibrium transport in the presence of carrier heating, as a result of the decrease in scattering event quantity [117]. Other studies also confirm the increasing significance of velocity overshoot and the non-local transport effects [118]. Overall, the Monte Carlo method offers adequate computational efficiency for 3D simulation, nano-scale semiconductor devices. It should be
noted that the method is a direct numerical solution to the BTE whole accuracy depends on the number of superparticles and the time step used in the simulations [119].

Figure 3-6 Flowchart showing the computational steps needed for Self-Consistent Quantum Monte Carlo [120].
3.6 Scattering in Monte Carlo

In order to accurately represent drive current variations and magnitude along with electrostatic effects, which primarily impact threshold voltage and sub-threshold current, it is crucial to introduce the relevant scattering processes into the Monte Carlo simulation. However, this requires consideration of the trade-off with physical accuracy and computational efficiency. The most common approach taken to achieve this is random number selection, which determines the scattering process employed in the simulation via a comparison table of the probabilities associated with each process [120]. Fermi’s golden rule (3-45), (obtained through the Schrödinger equation) provides the probability of perturbation Hamiltonian $H_{k',k}$ leading to a shift between state $k$ and $k'$ ($S(k, k')$):

$$S(k, k') = \frac{2\pi}{\hbar} |H_{k',k}|^2 \delta[E(k') - E(k)] \quad (3-45)$$

The scattering probabilities associated with each process, as derived from the above equation, are then tabulated, enabling the calculation of the total scattering rate and the selection of the particular scattering process at the end of the free flight. This approach is adopted for example for calculating acoustic and optical phonon scattering rates [112], as well as scattering rates for ionised impurities and interface roughness [121]. The simulation of the interface roughness scattering can also be achieved by using an alternative approach, wherein specular or diffusive reflection [122] is introduced through weighted selection [123]. However, the limitation of this approach is that it depends on the selection of the specular-diffusive scattering ratio which is device specific [124].

A new final state is randomly selected post-scattering. The randomly selected scattering mechanism determines the modification of the particle’s energy, direction and magnitude and direction of the momentum (k-vector). Numerous studies discuss the selection methods used for each of the individual scattering mechanisms.
In the case of random discrete dopant ‘*ab initio*’ ionised impurity scattering can be implemented in the Monte Carlo simulation. In this case, the impurity scattering is represented not via scattering rate and corresponding random number selection and respective scattering rate probabilities, but by the direct impact if the impurities Coulomb potential on real space trajectories. The *ab initio* method has been adopted in earlier research to study the impact of Coulomb scattering from random ionised impurities or a trapped charge [125]. Here, the particle-particle-particle-mesh (P³M) algorithm is used to calculate the short-range Coulomb force determining real space trajectory of each particle:

\[
F_{SR}(r) = -\frac{Q_r}{4\pi \varepsilon \left(r^2 + 0.5r_c^2\right)^{3/2}}
\]  

(3-46)

Where \(r_c\) is a cut-off radius selected to block unwanted carrier heating [126] and \(r\) is the distance to the point like charge. The long-range interactions are included via the solution of Poisson’s equation. This allows the random dopant scattering induced current variability to be simulated [127], [128]. This has proved to be an efficient method that could capture not only the localised carrier-impurity interactions but the carrier-carrier scattering in small devices where \(r_c\) is a threshold radius restricting carrier heating, and \(r\) is the distance to the charge. The solution to the Poisson equation provides the long-range interactions, enabling a high level of accuracy in simulating the current variability caused by random dopant scattering. Previous research confirms the ability of this approach to accurately represent both localised interactions between carriers and impurities as well as scattering between carriers in small-scale devices [127].
3.7 Quantum Corrections in Monte Carlo

The incorporation of quantum corrections into the Monte Carlo simulation method enables the accurate simulation of quantum confinement effects which play important role determining the performance of nano-CMOS transistors. The quantum-corrected Monte Carlo method provides accurate simulation results in highly-scaled devices with $t_{Si}$ thickness in the range of 10 - 7 nm [129]–[133]. Therefore, the incorporation of quantum corrections into the Monte Carlo model both upholds the approach’s computational efficiency and enables tunnelling and size quantisation effects to be captured. Different methods have been used to introduce quantum corrections in the MC simulations including the solution of the Schrödinger [134], [135] and Wigner [136] equations, as well as by using effective potentials derived from accurate quantum simulations or related to the DG approach [119], [137], [138]. The standard method introduces quantum-correction potential ($\psi_q$) as follows:

$$\psi_q = \psi_c + \psi_{qc}$$  \hspace{1cm} (3-47)

where $\psi_c$ is classical potential (derived from the solution to Poisson’s equation), and $\psi_{qc}$ is quantum-corrected potential (derived via the methods outlined below). rather than integrating the quantum-correction potential with the current continuity equation, as in the case of DD simulation, quantum-corrected potential is instead utilised to determine the force ($F$) impacting particles during the free flight period under the Monte Carlo method:

$$F = F_c + F_{qc}$$  \hspace{1cm} (3-48)

$$F_c = -\nabla \psi_c$$  \hspace{1cm} (3-49)

$$F_{qc} = -\nabla \psi_{qc}$$  \hspace{1cm} (3-50)
Compared to quantum transport simulations, which offer greater accuracy and detail at high computational expense, the introduction of quantum correction into semi-classical transport offers greater efficiency, particularly given that PS equations are typically solved in the cross sectional discretisation lanes once in only few times during the 3D Monte Carlo simulations [139]. This is the most accurate method used to introduce quantum corrections in GARAND MC engine. Alternatively, the DG method can be used for introducing the quantum correction force in GARAND MC. In this case the calibrated DG quantum correction method is used in the DD simulations used for initialisation of the MC simulations. The mobile charge distribution at the end of the DD simulations is used to obtained $\psi_{qc}$.

In the case of two-dimensional Monte Carlo simulation, a previous study presents a method for incorporating the Poisson- Schrödinger solution into the model with greater efficiency [139]. Here, periodic solutions to the Schrödinger equation are calculated, with a perturbative method used to calculate eigen energies during the remaining time steps. In other research, a Wigner distribution function-based correction term has also been presented [136]. This is expressed as follows [140], beginning with the Wigner transport equation, as a modified BTE:

$$
\psi_q = \psi_c + \frac{\hbar^2}{4m^*r_k_B T} \left[ \nabla^2 \psi_q - \frac{1}{2k_B T} \left( \psi_q \right)^2 \right] 
$$  (3-51)

Calculating the solution to the above equation is challenging given its highly non-linear structure. This being said, compared to the earlier approach, Eqn. (3-51) provides the benefit of lower noise sensitivity. Previous research has confirmed the effectiveness of the correction term in two-dimensional Monte Carlo simulations. Additionally, other studies have adopted the method and applied it to incorporate numerous valleys’ electron distribution [140], with the equation also applied beyond the thermal equilibrium assumption. Both two-dimensional and three-dimensional Monte Carlo simulations have been carried out in the literature using the effective potential method to accurately capture confinement effects. Additionally, with multi gate MOSFETs, the method has also been
adopted to explore surface roughness scattering [141]. A scattering rate must be used to ensure accuracy in modelling whilst also overcoming the limitation of the effective potential method, in that it overestimates the degree to which carriers are turned away from the interface. Thus, the selection of a weighted specular/diffusion reflection is unsuitable in this case. The method has also been used to explore Coulomb interactions with unintended doping [142], with dopant channel placement being found to be effective in capturing degradation.

The adoption of the fitting parameter poses a noteworthy limitation to the method, however [142]. When the field is overestimated, the interface’s electron concentration is underestimated, resulting in the peak concentration moving away from the interface. In comparison with Poisson-Schrödinger solutions, if the parameter is reduced in order to correct this issue, this can result in a lower correction term than anticipated, along with a peak concentration that is too high. The fitting parameter values must be varied in the normal and parallel directions to the interface when the method is applied in 3D simulation.

The replacement of the Gaussian distribution with a Pearson IV distribution has been carried out in order to present an alternative effective potential approach. Depending on the distance between the carrier and semiconductor-oxide interface, the Pearson distribution can be tuned at different points in the vertical direction. This is achieved using four parameters, rather than the single parameter used in the Gaussian approach. The benefit of this method is greater consistency with the Poisson-Schrödinger solution-based electron distribution [142]. However, validation and implementation of this method has only been achieved in 2D simulation to date, with no extensive simulation research adopting the approach. The following equation can be used to obtain a more computationally efficient Schrödinger-based correction term, compared to the incorporation of a PS-based solution into Monte Carlo simulation:
\[ \psi_q(z) = -k_B T \log(n_q(z)) - \psi_c(z) + V_0 \]  

(3-52)

where \( n_q(z) \) is quantum density (derived from the solution to the Schrödinger equation) [134]. Here, \( n_q(z) \) does not reflect values, but distribution shape. Quantum density is calculated by periodically applying the eigenvalue solver to slices along the quantisation direction, which upholds self-consistency. The correction term expressed in Eqn. (3-52) is modified based on the result, although the solution to the Poisson equation is reached separately based on the Monte Carlo electron distribution. This allows the simulation to achieve the same computational efficiency as traditional models. This approach does not require consideration of the Fermi level, since it only requires the quantum carrier density shape. The method can be applied beyond the assumption of thermal equilibrium by means of the varying carrier temperatures between the slices. Additionally, consistency with the solution to the Poisson- Schrödinger is achieved due to the method’s adoption of the Schrödinger solution. Both two-dimensional [143], [144] and three-dimensional [143], [145], [146] applications of the method have been carried out, with correction and transport being two- and three- dimensional, respectively. In both cases, a good level of accuracy has been found in terms of size quantisation. However, the method has been found to be unsuitable for capturing tunnelling effects [147].

### 3.8 Mobility and I-V calibration

One of the key parameters impacting semiconductor device performance in DD simulations is the carrier mobility (\( \mu \)). The dependence of mobility with the field (mobility models) has traditionally been the way that DD simulations have been parameterised and calibrated to correctly match theoretical results with experiment. There is a strong correlation between the drive current and the mobility, with the drive current increasing as mobility increases. Since scattering mechanisms such as surface roughness, lattice/phonon and ionised impurity scattering have an impact on mobility, the accurate representation of carrier mobility
requires the key scattering mechanisms to be taken into account. Matthiessen’s rule (3-20) is used when multiple scattering mechanisms are introduced in the mobility models.

The carrier mobility is reduced by ionised donor/acceptor-induced impurity scattering in doped semiconductors. A typical mobility model in DD simulations has three components including low-field mobility model, perpendicular-field mobility dependence and lateral-field mobility dependence. There is a large variety of mobility models used in DD simulations with different level of complexity and sometimes the low field mobility and the field dependency cannot be clearly separate.

The low-field mobility model is further reduced by the perpendicular-field model. The combination of the two models determine the $I_D-V_G$ characteristics of the CMOS transistor at low drain bias. At high drain bias the mobility if further reduced by the lateral-field mobility model which accounts for the velocity saturation effects at high electric field. The three mobility models used in this study are explained in further detail below.

### 3.8.1 Low field mobility dependence

The low-field mobility value determines the base mobility mainly determined by the phonon and the impurity scattering. Near the interface the low filed mobility is modified to take into account the impact of the interface on the mobility in the MOSFET channel. It can also be modified by the strain used to enhance the performance of contemporary CMOS transistors. The Masetti (concentration-dependent) [148], Arora [149] (concentration-dependent) or Philips [150] (concentration-dependent, including screening effects) mobility models are available for selection in the atomistic simulator GARAND.
3.8.2 **Vertical field mobility dependence**

In order to capture the effects of surface roughness scattering at the semiconductor/insulator interfaces, the low-field mobility value is adjusted through the use of the perpendicular-field mobility model to deliver the perpendicular field-dependent mobility ($\mu_\perp$). This value is then further adjusted using the chosen lateral field-dependent mobility model. The available perpendicular field-dependent models in GARAND include the Yamaguchi [151], Lombardi [152] and Thin-Layer [153] models.

The Yamaguchi model [151], (Eqn. (3-53)) is an empirical model that delivers a reduction in the low field mobility of the inversion layer as a response to the confinement field perpendicular to the interface and therefore to the direction of the carrier flow in the channel:

$$\mu(E_\perp) = \frac{\mu_0}{\sqrt{1 + D(r_\perp) \left(\frac{E_\perp}{E_c}\right)^\alpha}}$$

(Eqn. (3-53))

where $\mu_0$ is the low-field mobility, $E_c$ is the critical electric field (used as a fitting parameter for the model), and $\alpha$ is an exponential factor used as a second fitting parameter. The above equation (3-53) presents the numerous parameters expressed in the model, and these are also outlined in previous research [151].

The low field mobility $\mu_0$ impacts low drain bias current in the linear region of operation, $E_\perp$ impacts the start of the bending in the $I_D$-$V_G$ characteristics, and $\alpha$ impacts the curvature of the bending. This can be illustrated by using the simple current model in the linear region, work function can be used as a fitting parameter $V_T$ taking into account the impact of the short channel effects:
Chapter 3. Simulation Methodology

\[ I_D(V_G) = W \mu(E_{\perp}) C_{Ox}(V_G - V_T) \frac{V_G}{L} \]  
(3-54)

3.8.3 \textbf{Lateral field mobility dependence}

The use of a lateral field-dependent model causes high-field velocity saturation effects to be reflected in mobility. The mobility \( \mu \) value is taken as the final value for isotopic application across the simulation region.

The following equation presents the Caughey-Thomas field-dependent mobility model [154], which is in empirical accordance with velocity-field characteristics:

\[ \mu(E_{\parallel}) = \frac{\mu_o}{\left(1 + \left(\frac{\mu_o E_{\parallel}}{v_s}\right)^\beta\right)^{1/\beta}} \]  
(3-55)

The fitting parameter \( v_{sat} \) impacts the slope if the \( I_D-V_G \) curve is at high \( V_D \), whilst a \( \beta \) impacts bending. The following equation expresses the temperature-dependent saturation velocity \( v_{sat} \) [155]:

\[ v_{sat} = v_{sat} \left(\frac{300}{T}\right)^{v_{exp}} \]  
(3-56)

Temperature-dependent fitting parameter \( \beta \) can also be expressed as:
\[ \beta = \beta_o \left( \frac{300}{T} \right)^{\beta_{\text{exp}}} \]  

(3-57)

The high drain bias the simple current model in the saturation region is expressed as:

\[ I_D(V_G) = \frac{W}{2L} \mu(E_{\parallel}) \cdot C_{\text{OX}} \cdot (V_G - V_T)^2 \]  

(3-58)

In the simple current model in the saturation region the lateral field can be approximated as \( E_{\parallel} = V_D/L \), with work function again used as a \( V_T \) as the fitting parameter taking into account the impact of the short channel effects.

3.9 TCAD Tools and Data Analyses

This research has adopted the GSS 3D statistical atomistic TCAD simulator GARAND and the automation and optimisation engine Enigma, which offer significant accuracy and efficiency in device calibration, optimisation and modelling.

3.9.1 GARAND

GARAND used comprehensively in this research is a TCAD simulator specifically developed to simulate statistical variability and reliability in contemporary and future CMOS transistor. The GSS (Synopsys) website details a number of applications of the GARAND tool in this context. GARAND has offers different simulation modules, including Monte Carlo, drift-diffusion with PS and DG quantum corrections. Additionally, GARAND also provides the latest physical models, supporting accuracy at atomic scale. The tool provides the ability to use density gradient quantum corrections for both electrons and holes simultaneously. Additionally, GARAND also offers extremely efficient computation with
guaranteed convergence in complex 3D simulations. Furthermore, GARAND takes into account random discrete dopants, trapped discrete charges, gate stack granularity, line edge roughness, and other key sources of statistical variability individually and in combination. Figure 3-7 illustrates the statistical drift-diffusion simulation engine fits into the GARAND tool-chain.

3.9.2 **Enigma**

The statistical simulation of CMOS transistors requires simulation in 3D of large statistical samples which can be efficiently performed on computing clusters using farming. Data flow between tools within the GSS tool chain is maintained through the use of a specific database architecture. The automated cluster job submission and management system provided by the GSS tool chain offers high productivity and efficiency, enabling high numbers of simulation tasks to take place at the same time in clusters with large number of processors. The results

*Figure 3-7 Flow diagram defining the relation between the GARAND drift-diffusion and Monte Carlo simulation modules.*
of the simulations are automatically gathered and summarised through the job submission and management interface with the GSS database.

3.10 Summary

This chapter has outlined in the basis of the 3D drift/diffusion simulation, presenting both the drift/diffusion and the density gradient equations as well as the algorithms employed solve them self consistently. Semi-classical approaches are used to simulate the charge transport based on the Monte Carlo method, where the charge trajectories between two scattering events are simulated using classical dynamics. Fermi’s golden rule [156] is uses, to calculate the scattering rates based on analytical non-parabolic band structure. The 3D EMC simulations are used to capture the non-equilibrium transport in aggressively-scaled NWT, leading to accurate estimation of the transistor performance. This chapter has also described the quantum correction based on the solution to 2D Poisson-Schrodinger equations for each discretisation cross-sectional plane perpendicular to the transport direction [157]. The quantum potential obtained from the PS solution determines the driving field, which controls the free flight of the particles. This chapter has also discussed the stability and efficiency of the DD simulation method, with a straightforward yet effective DG approach to the introduction of quantum effects into simulation. It was clarified that non-equilibrium transport effects cannot be represented by the DD model in nano-scale devices. Furthermore, transport variability due to ionised impurity scattering is also difficult to capture using DD- simulation. Monte Carlo simulation are used to calibrate the drift-diffusions of large multichannel NWTs and to perform statistical variability simulations [158].
Chapter 4. Si Nanowire Transistors Design

4.1 Introduction:

Electronics manufacturers, influenced by Moore’s law, have been drawn into innovating many key technical CMOS technology developments including high-k gate dielectrics technology and strain technology, which involve both new processes and new materials [26], [159]–[161]. Multi-gate MOS FET modelling is anticipated to be a primary focus for future developments. The superior electrostatic integrity provided by Nanowire (all gate around) Transistors (NWT) [7]–[9], [162] ensures that these innovative transistors remain popular candidates for achieving ultimate CMOS scaling limits [162]–[164]. The scaling limits are ultimately defined by the quantum mechanical effects which control the operation of NWT transistors [4]. Threshold voltage shift, resulting from confinement effects, together with the reduction of the gate-to-channel capacitance are reducing the available for transport charge within the channel [6]. Lower gate-to-channel capacitance will additionally negatively impact the electrostatic integrity. These effects increase with the reduction of the NWT dimensions and are therefore key to the assessment of NWT scaling limits [9].

Density gradient quantum corrections, in commercially available TCAD tools, represent the pragmatic approach for dealing with quantum confinement effects [7]. However, no investigations of the importance of quantum effects and the relevance of the DG technique for the modelling of ultimately scaled NWTs of varying cross-sectional shape at the scaling limits have yet been conducted. This means that the effect of NWT cross-sectional shape on ultimate scaling limits remains unaddressed, along with other critical design questions.

This chapter examines the ultimate scaling limits of NWTs with different cross-sectional shapes, using simulations employing Poisson Schrodinger (PS) quantum corrections implemented in the ‘atomistic’ drift-diffusion (DD) simulator GARAND [GSS]. Realistic
design parameters of the source-drain extensions and their impact on SS and DIBL are considered. In addition, the effect of the nanowire cross-sectional shape on the gate capacitance $C_G$, the charge available for transport $Q_G$ and the speed (evaluated by using the $Q_G/C_G$ ratio) are investigated and contrasted with $\tau$.

This chapter will commence with an introduction detailing the NWT design employed in this investigation. The next section, demonstrates the key elements of the NWT structure design. Section 4-3 describes the structure editor employed in this study. Section 4-4 provides an overview of the design details of 5 nm NWT. The simulation methodology and the density gradient calibration have been explained in Sections 4-5 and 4-6. The analysis of effect of quantum confinement on the gate capacitances and the available charge for transport in the channel are described in Section 4-7. Sections 4-8 examines the effects of quantum confinement on the nanowire electrostatic integrity, including SS and DIBL. Section 4-9 attempts to find the optimal cross-sectional aspect ratio. The penultimate Section 4-10 investigates the effect of varying source/drain doping designs on optimal channel length for different NWT cross sections. The conclusions are drawn in the final Section 4-11.

### 4.2 Nanowire Transistor Design Structure

A nanowire FET comprises of a silicon body within a nanowire shape acting as the conducting channel; whilst the gate dielectric layers and gate material wrap around the semiconductor to form the MOS structure, as depicted in Figure 4-1.

#### 4.2.1 Silicon Channel

Ever since the first silicon transistor was produced in the 1950s, silicon (Si) has been the basis of computer chips and the foundation of the semiconductor industry, with substantial investments in R&D over the following decades to ensure production of dependable scaled transistors. So, whilst other elemental or compound semiconductors may offer greater speed than Si at ultra-high-speed and ultra-low-power applications [165], a waste amount of
additional R&D expenditure in time and money would be required to deliver and optimise corresponding new chip manufacturing and design processes. A primary advantage of Si is its susceptibility to oxidation that produce silicon dioxide SiO$_2$, a gate dielectric with exceptional properties. The oxidation proceeds in dry or wet oxygen atmosphere at high temperature. Si also remains stable at high temperatures of operation. Additionally, Si is insoluble in water and this is beneficial in minimising various problems associated with manufacturing and functionality. Si exists naturally as a perfect crystal with only minute defects apparent in ultra-large ULSI integrated chips. Si is also inexpensive and commonly sourced across the Earth. These properties make Si the semiconductor of preference in the production of robust transistors and chips.

Based on a set of commercial criteria, for each technology node, a key research objective is to continually investigate a variety of transistor scenarios with the aim of ensuring that Si remains the basis for future chip manufacturing. Strained Si has demonstrated important key benefits for improving transistor performance, particularly with regard to p-type MOSFETs. Particular strain patterns result in tripling the hole mobility and doubling the on-state current of p-type transistors. Further to Si strain development, alternative channel materials are also being investigated for their potential of enhancing MOSFETs speed.

*Figure 4-1 schematic view and material details of the circular NWT (S/D contacts are not shown).*
4.2.2 Gate dielectrics layers

Excessive scaling of SiO$_2$ layer results in higher gate leakage as a consequence of quantum mechanical tunneling effects [166]. Minimising the gate leakage current in CMOS components, necessitates replacing the traditional SiO$_2$ gate dielectric layer with an alternative dielectric of higher dielectric constant ‘high-k’. HFO$_2$ has high permittivity (k > 20) and has therefore was introduced as an alternative gate dielectric.

Conversely, Hafnia and the majority of high-k MOS dielectrics have unstable interfaces with Si channels. Here the high-k layer reacts with Si under equilibrium conditions to produce an undesired interfacial layer (IL). To enhance the capacitance of the gate stack and match the high-k layer with the Si interface, an insulating interfacial layer is needed [167]. The insulating interface provided by SiO$_2$ circumvents undesired movements of threshold voltage and flat band voltage. It also offers improved thermodynamic stability [168]. Insertion of a thin IL of SiO$_2$ between the high-k gate stacks and Si channel promotes reliability by lowering ITC (interface trap density) defects at the Si-interface [29], [166], [167], [169]–[174]. The IL lowers gate stacked/Si roughness and improves the mobility [175]. The peak mobility in MOSFET devices with a high-k layer has been shown to be dependent not only on IL material, but also to be directly proportional to IL thickness for both poly-Si and metal gates [176], [177].

Conversely, the hafnia thickness tends to increase the low-frequency noise associated with the MOSFET current when compared with SiO$_2$ alternatives. The IL minimises noise, whilst reducing SiO$_2$ thickness results in greater, low-frequency noise as a consequence of interface trapping and mobility variations [178], [179].

The SiO$_2$ IL is found between the Si conducting channel and the high-κ dielectric layer can be deliberate or unintentional. The thin IL develops between the high-κ layer and the Si as a consequence of many high-κ dielectric deposition techniques resulting in thermally unstable interfaces with Si. The IL derives from the diffusion of oxygen from the high-κ layer towards the Si interface mainly during the process of activation annealing, rather than the process of high-κ deposition itself. The molecular bonding of Si atoms to oxygen
promotes the controlled diffusion of the oxygen by producing high-κ stacks with SiO₂ IL\cite{180}, \cite{181}.

Whilst the IL offers improved mobility, better stability and reduced noise for high-κ gate stacks, it also lowers the total gate capacitance \cite{174}, \cite{182}, \cite{183}. The total capacitance of MOS devices comprises the two dielectrics layers in series (4-1):

$$\frac{1}{C_{ox}} = \frac{1}{C_{SiO₂}} + \frac{1}{C_{high-κ}}$$  \hspace{1cm} (4-1)

The MOS capacitance $C_{ox}$ is determined by dielectric primitivity and thickness as described in equation (4-2):

$$C_{ox} = \frac{\varepsilon_{ox}\varepsilon_o}{t_{ox}}$$  \hspace{1cm} (4-2)

Where $\varepsilon_{ox}$ represents the relative dielectric permittivity, $\varepsilon_o$ represents the permittivity of vacuum, and $t_{ox}$ represents the oxide thickness. More specifically, the employing a dielectric with high $\varepsilon_{ox}$ more facilitates the scaling to the transistor dimensions. The expression “equivalent oxide thickness” (EOT) is employed to extrapolate the thickness of SiO₂ necessary to achieve capacitance of high-κ material of an equivalent area. The EOT equation (4-3) is:

$$EOT = \varepsilon_{SiO₂} \frac{t_{high-κ}}{\varepsilon_{high-κ}}$$  \hspace{1cm} (4-3)

Equations (4-1), (4-2), and (4-3) give rise to the EOT of two layers of gate stack (4-4):

$$EOT = \varepsilon_{SiO₂} \frac{t_{SiO₂} + t_{high-κ}}{\frac{\varepsilon_{SiO₂}}{\varepsilon_{high-κ}}}$$  \hspace{1cm} (4-4)
4.2.3 **Gate Electrodes**

Metal gates, in combination with Si and appropriate dielectric, can act as gate electrodes to form a MOS channel region. However, use of poly-Si as a gate electrode can result in problems with poly-depletion and resistivity effects. Therefore, metallic TiN gates are used in contemporary CMOS technologies [172].

4.3 **3-D Structure Modelling Tools**

Adaptable techniques are required for modelling the new innovative 3-D Nano-transistors. This study uses the Structure Editor Sigmund [GSS] to generate accurate simulation domains of nanowire structures stored in VTK format which can be loaded and used by the device simulator GARAND. The hierarchy of object structure flow employed by the Sigmund Structure Editor is depicted in Figure 4-2. The structure editor sets the 3-D simulation domain, which can then be populated with material regions, doping regions and contact. Sigmund also meshes the simulation domain prior storing it in the VTK file. This is a five-step process:

1- Identification of materials, doping type, dimensions and other device parameters.

2- Utilisation of the template geometry objects (triangular prism, cylinder, cuboid, tetrahedron, regular tetrahedron and sphere) to generate the 3-D structure. Amalgamations of the template objects permits creation of more complex structures.
Sigmund Structure Editor

- Material Map
- Structure Editor
- VTK file
  Input to Atomistic Simulator

Objects
- Triangular prism
- Cylinder
- Cuboid
- Tetrahedron
- Regular Tetrahedron
- Sphere

Doping Regions
- Block
- Gaussian 1-D
- Gaussian 2-D
- Gaussian 3-D
- Block-Gaussian

Mesh Region

Figure 4-2 Process flow of (Sigmund) structure editor.
3- Following the generation of the structure, doping can be introduced via inbuilt analytical doping regions employing a combination of position and materials options. The Sigmund Structure Editor offers five doping options: Block, Gaussian 1-D, Gaussian 2-D, Gaussian 3-D, Block-Gaussian.

4- Whilst meshing is defined at the same time as the objects, remeshing can be applied at any time to achieve target resolution.

5- Metal gate granularity and line edge roughness (LER) are additional options which can be used to supplement the device structure or selected objects.

4.4 5 mm Nanowire Transistor Design

Key design criteria:

1- Material composition of channel, interfacial layer, high-k gate stack.

2- Dimensions of device cross-sectional shape, gate length, effective oxide thickness and source/drain spacers.

3- Crystallographic orientations and strain of the semiconductor channel.

4- Doping concentrations, such as peak and roll-off doping.

Si is used for the simulated NWT channel, source, and drain regions. SiO₂ and Hafnia comprise the 0.4 nm interfacial layer and 0.8 nm high-k layer determine the gate stack (according to the ITRS scaling criteria). According to Equation 4-4 this results in gives equivalent effective oxide thickness EOT=0.6. Figure 4-3 depicts the 3D structure of a cylinder NWT. In this investigation, the electrostatic performance is examined for silicon channels with four different cross-sectional shapes: circle; ellipse; rectangular; square. To ensure an unbiased comparison, all cross-sectional shapes were designed with identical
surface areas: $A_{\text{cross-section}} = 9\pi \text{ nm}^2$, i.e. the diameter or (fin-pitch) of the circular NWT is 6nm. Table 4-1 gives the precise cross-sectional dimensions for all four investigated shapes. The two channel crystallographic orientations studied on (001) wafer are $<110>$ and $<100>$. Table 4-2 summarises design parameters for all devices: the effective gate length is 10-20 nm, the spacer thickness is 5nm, the source/drain peak doping concentration is $4 \times 10^{20} \text{ cm}^{-3}$.

The low channel doping boosts charge mobility with a significant improvement of the FET performance and it also reduce local (statistical) variability. It has been demonstrated that doped FinFETs retain a variability advantage over classical planar MOSFET if the channel doping is lower than a few $10^{18} \text{ cm}^{-3}$ [184]. The channel doping of Intel 14nm FinFET is about $10^{17} \text{ cm}^{-3}$. Better electrostatic integrity, offered by gate-all-around nanowire transistor (NWT) architectures allow reducing the doping of the channel [7], [8], [9], [162], [164], [185]–[189]. In this study, the channel doping is lowered up to $10^{14} \text{ cm}^{-3}$.

Sigmund Structure Editor was used to generate suitable NWT structures for this study. Figure 4-4 depicts the doping profile of the Si NWT studied with a gate length $L_g = 12$ nm and spacer length $L_s = 5$ nm.
Table 4-1 Physical dimensions of simulated devices

<table>
<thead>
<tr>
<th></th>
<th>Y(nm)</th>
<th>Z(nm)</th>
<th>Y/Z</th>
<th>Area (nm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Circular</td>
<td>6</td>
<td>6</td>
<td>1</td>
<td>9π</td>
</tr>
<tr>
<td>Elliptical</td>
<td>3.45</td>
<td>4.64</td>
<td>0.74</td>
<td>9π</td>
</tr>
<tr>
<td>Square</td>
<td>3.54</td>
<td>3.54</td>
<td>1</td>
<td>9π</td>
</tr>
<tr>
<td>Rectangular</td>
<td>3.06</td>
<td>4.13</td>
<td>0.74</td>
<td>9π</td>
</tr>
</tbody>
</table>

Figure 4-3 A 3D representation of cylindrical NWTs with S/D contacts.
Figure 4-4 The doping profile of the Si NWT with gate length Lg = 12 nm and spacer length Ls = 5 nm.

Table 4-2 Parameters of the simulated devices

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T_{II}$ (nm)</td>
<td>0.4</td>
</tr>
<tr>
<td>$T_{high-k}$ (nm)</td>
<td>0.80</td>
</tr>
<tr>
<td>Gate length (nm)</td>
<td>14</td>
</tr>
<tr>
<td>Spacer thickness (nm)</td>
<td>5.0</td>
</tr>
<tr>
<td>S/D peak doping (cm$^{-3}$)</td>
<td>$4 \times 10^{20}$</td>
</tr>
<tr>
<td>Channel doping (cm$^{-3}$)</td>
<td>$10^{14}$</td>
</tr>
<tr>
<td>Substrate orientation</td>
<td>001</td>
</tr>
<tr>
<td>Nanowire orientation</td>
<td>$&lt;110&gt; &amp; &lt;100&gt;$</td>
</tr>
</tbody>
</table>
Figure 4-5 Wave functions in the $\Delta_1$, $\Delta_2$, $\Delta_3$, and $\Delta_4$ degenerate valleys in the perpendicular (slice) cross-section of Si NWT (PS simulation).
Figure 4-6 Four NWT’s cross-sections simulated in this subsection. Comparison of the charge distribution in the NWT cross-section obtained from three simulations methods; Poisson-Schrödinger, density gradient the classical DD. The simulations at low drain voltage and at gate voltage \( V_G = 0.60V \).
4.5 Simulation Methodologies

Subsequent to the generation of the VTK files with required NWT structures, the simulation was conducted to examine three simulation options: Poisson-Schrödinger (PS) quantum correction directly implemented in the drift-diffusion (DD) simulator; a Density Gradient (DG) quantum corrections and classical simulation. The PS and DG simulations have been compared to assess the accuracy of the DG simulations of quantum confinement effects in the NWTs appropriate for sub-5-nm CMOS applications with varying cross-section.

Non-parabolic band structure effects were not captured in the 2D cross-sectional slices along the gate length of the simulated 3D transistors forming the 2D solution of the Schrödinger equation. A LAPACK solver was utilised to solve the 2D Schrödinger equation in the effective mass approximation. The wave function penetration in gate oxide is taken into account. Figure 4-5 depicts the wave functions in the $\Delta_1$, $\Delta_2$, $\Delta_3$, and $\Delta_4$ degenerate valleys in the cross-section of an cylindrical Si NWT at gate bias of 0.60V. The probability distribution obtained from the wavefunctions combined with the occupation density resulting from the Fermi level and eigenvalues (subband energies) are used to calculate the 2D charge distribution. The PS charge distribution is coupled to the DD transport. A DD simulation is initially performed until convergence is attained, and then the quasi-Fermi level resulting from the converged DD solution is employed as a fixed reference for the PS solution [9]. The effective quantum- potential is used as the driving potential in the solution of the current-continuity equation, where the charge distribution in the NWT cross-section mirrors the charge distribution resulting from the solution of the Schrödinger equation [7], [185].

The charge distribution in the NWT channel from the Poisson–Schrödinger (PS), density gradient (DG) and classical simulation is comported and analysed. Figure 4-6 depicts the 2D charge distribution at the centre of the NWT channel. Figure 4-6 illustrates how the 2D charge distribution resulting from uncalibrated (standard) DG simulations may not effectively capture the quantum mechanical effects in respect of the mobile charge distribution in the channel. Results of Poisson–Schrödinger (PS) and density gradient (DG)-
based quantum corrections are subject to a detailed comparison to evaluate the applicability of the DG approach for simulation and design of scaled NWTs in the following section.

4.6 Density Gradient calibration

Whilst the PS quantum corrections deliver highly accurate quantum charge distribution in the simulated NWT channel, the large number of cross-sectional solutions for the Schrödinger equation causes substantial slowing of the simulations with concomitant reduction in productivity and efficiency [162], [164]. Therefore, to address these issues a compromise solution involves calibrating the DG quantum corrections to the PS charge distribution and then utilising the DG simulations. The DG formulation includes a quantum correction term, \( \psi_{\text{qm}} \), proportional to the second derivative of the square root of the carrier density as explained in chapter three (equation (3-37)).

The DG equation is calibrated against quantum-mechanical simulations using Poisson-Schrödinger solution using the DG effective mass, as a fitting parameter. Furthermore, the DG effective mass is considered to be anisotropic, resulting in different effective mass components associated with each of the coordinate system directions \( m_x \), \( m_y \) and \( m_z \). As explained in chapter three, equation (3-38) is the form of the solved DG equation.

The DG effective masses for both the conduction band in Si and the SiO\(_2\) are calibrated to match the charge distributions resulting from PS solutions associated with the major and minor axes of the NWT cross-section and the integrated mobile charge in the channel at a 0.60V gate voltage. It is important that the orientation dependence is taken into account by the PS solution. The DG effective masses simply represent fitting parameters. Additionally, the quantum confinement is only relevant normal to the direction of the transport, and the DG effective mass along the transport direction is set so as to be sufficiently large to prevent artificial S/D tunnelling.

Figure 4-6 illustrate the charge distribution obtained from the standard DG and calibrated DG of Si NWT. Figure 4-7 illustrate the charge distribution differences between different NWT cross-sections resulting from PS, DG and classical DD simulations. It can be seen
that the calibration of the drift-diffusion simulations can significantly impact charge distributions. A fuller illustration of the reliability of the calibration process is provided in Figure 4-9 in comparison with the 1D charge distributions resulting from the PS and DG solutions along the main ‘diameters’ of the NWT cross-section.

Optimum channel gate control with excellent electrostatic integrity is offered by GAA design. However, quantum mechanics exerts significant effects in such ultra-scaled GAA NWT, which must be addressed in order to produce accurate performance results for the device using simulations. Four different NWTs cross-sections are studied here using simulations: cylindrical, elliptical, square and rectangular. Although the elliptical and rectangular NWTs have different characteristic dimensions, all devices have the same cross-sectional area of $9\pi \text{ nm}^2$. $<110>$ and $<100>$ and crystallographic orientations on (001) wafer are examined. Table 4-2 summarises the cross-sectional dimensions for all six nanowires.

![Figure 4-7 Charge distributions obtained from the standard DG and calibrated DG of Si NWT (y-z) and (x-z) cross-sections.](image)
Figure 4.8 Capacitance-voltage (C-V) characteristics of the simulated NWT with six different cross-sections.

Figure 4.9 1D charge distributions obtained from the PS simulation and DG simulation. The cutline is along of perpendicular ‘diameters’ (y and z) of the NWT’s slice at the mid of the gate.
4.7 NWT Mobile Charge and Gate Capacitance

The influence of the quantum mechanical (QM) effects on the gate voltage dependence of the gate capacitance for the different cross-sections depicted in Figure 4-6 and Table 4-1 are examined. An infinite gate length has been assumed to emphasise the effect that cross-section has on the gate capacitance. Figure 4-8 depicts the capacitance-voltage (C-V) characteristics of the simulated NWT with six cross-sections. The impact of the QM effects on the gate capacitance are clearly demonstrated. In the quantum simulations the gate capacitance is reduced by a mean value of approximately 32% compared to the results of the classical simulations. Most significantly, the magnitude of the QM effects is for different cross-sectional shapes, despite the identical cross-sectional area. Figure 4-8 shows that NWT with an extended elliptical cross-section (elliptical 2) exhibits the largest gate capacitance. This is probably due to the compensating effect that the elliptical shape induces in respect to the anisotropic spatial confinement / electrostatic confinement resulting from the dissimilar quantum masses along the cross-sectional diameters which provide a greater extent uniform charge distribution that the wrapped-around gate can control more effectively.

The mobile charge in the channel per unit length $Q_M$ at a particular gate voltage $V_G$ is directly proportional to the NWT gate capacitance per unit length $C_G$ and is given by the following equation:

$$Q_M = C_G(V_G - V_T)$$  \hspace{1cm} (4-5)

Where $V_T$ is the threshold voltage and $V_G - V_T$ represents the gate overdrive.

This indicates that reduction of the NWT gate capacitance will result in lower mobile charge in the channel and also in lower transistor performance. It is interesting that the precisely calibrated DG model closely resembles the PS results and therefore can be considered as an efficient approach to capture the quantum mechanical effects in TCAD device simulations, even for complex 3D transistors. Figure 4-10 depicts the gate voltage dependence of the
mobile charge in the channel for the NWTs in Table 4-1 (NB this refers to the integral mobile charge in the channel per unit area at low drain bias).

Eqn. (4-5) indicates that the reduction in gate capacitance results in a reduction of mobile charge in the channel, which consequently is an indicator of poorer NWT performance. Similar to gate capacitance, the mobile charge for NWTs with identical cross-sectional areas depends on the cross-sectional shape. The effects of NWT shape on potential NWT performance can be evaluated from Table 4-3 which compares $Q_M(V_G=0.60V)$ for identical $Q_M(V_G=0.0V)$. To perform this comparison the $Q_M(V_G)$ curves were aligned by changing the gate work function. Like the gate capacitance, the elliptical 2 NWT exhibits the largest mobile charge in the channel. Table 4-4 includes an additional case compared with the NWTs outlined in Table 4-1: the structures in this case have been rotated by 90 degrees whilst retaining the original crystal orientation. This perpendicular rotation of the optimal performing NWT leads to an improved $Q_M/C_G$ which represents around 11% and 10% improvement of $Q_M/C_G$ ratio. Here, the asymmetric spatial confinement induced by the elliptical shape aggravates the similar electrostatic confinement induced by the various quantum masses along the cross-section diameters, which gives rise to a charged distribution that is predominantly focused on only one of the two diameters and consequently less tightly controlled by the wrapped-around gate. Table 4-3 illustrates how the $Q_M/C_G$ ratio can be utilised as an indicator for the ‘intrinsic’ NWT speed. $Q_M$ and $Q_M/C_G$ are directly correlated which highlights not only the approximate nature of Eqn (4-5), but also the requirement to assess both $Q_M$ and $C_G$. Extended elliptical NWT exhibits the optimum $Q_M/C_G$ when compared to the other cross-sections. Table 4-5 provides the values for $Q_M$, $C_G$ and $Q_M/C_G$ ratio for the wires with identical matching cross-sectional areas as presented in Table 4-3, but also including the <100> channel orientation. Analyzing the results summarized in Table 4-3, Table 4-4, Table 4-5, gives rise to the following main conclusions.

First, all wires with the <100> channel direction have greater charge in the channel compared to the <110> wires. This confirms that varying the crystal orientation of the silicon channel results in change of mobile charge within the channel and therefore is an important design parameter. Consequently, the gate capacitance also increases leading to enhanced
electrostatic control. Additionally, the extended elliptical and rectangular NWTs offer about ~11% and 8% performance improvement compared to the circular and square shapes.

*Figure 4-10* Dependence of the mobile on charge gate voltage for the NWT’s (C-V) characteristics illustrated in 4-10 Capacitance-voltage (C-V) characteristics of the simulated NWT with six different cross-sections.
Table 4-3 $Q_M(V_G=0.600V)$, $C_G (V_G=0.600V)$ and $Q_M/C_G$ ratio at identical $Q_M (V_G=0.00V)$ for NWTs at $L_G=12nm$. With crystallographic channel orientation $<110>$.

<table>
<thead>
<tr>
<th>Shape</th>
<th>$Q_M(\times 10^6/cm)$</th>
<th>$C_G(fF/cm)$</th>
<th>$Q_M/C_G(10^{17}/F)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Circular</td>
<td>2.126</td>
<td>0.929</td>
<td>2.2876</td>
</tr>
<tr>
<td>Elliptical$_1$</td>
<td>2.374</td>
<td>0.996</td>
<td>2.3839</td>
</tr>
<tr>
<td>Elliptical$_2$</td>
<td>2.581</td>
<td>1.015</td>
<td>2.5429</td>
</tr>
<tr>
<td>Square</td>
<td>1.972</td>
<td>0.905</td>
<td>2.1784</td>
</tr>
<tr>
<td>Rectangular$_1$</td>
<td>2.266</td>
<td>0.985</td>
<td>2.3005</td>
</tr>
<tr>
<td>Rectangular$_2$</td>
<td>2.480</td>
<td>0.996</td>
<td>2.4888</td>
</tr>
</tbody>
</table>

Table 4-4 $Q_M(V_G=0.600V)$, $C_G (V_G=0.600V)$ and $Q_M/C_G$ ratio at identical $Q_M (V_G=0.00V)$ for NWTs at $L_G=12nm$. With crystallographic channel orientation $<110>$ where the structure is rotated by $90^\circ$.

<table>
<thead>
<tr>
<th>Shape</th>
<th>$Q_M(\times 10^6/cm)$</th>
<th>$C_G(fF/cm)$</th>
<th>$Q_M/C_G(10^{17}/F)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Circular</td>
<td>2.1476</td>
<td>1.0094</td>
<td>2.1275</td>
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<tr>
<td>Elliptical$_1$</td>
<td>2.6362</td>
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</tr>
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<td>Elliptical$_2$</td>
<td>2.964</td>
<td>1.1907</td>
<td>2.4892</td>
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<tr>
<td>Square</td>
<td>1.992</td>
<td>0.9834</td>
<td>2.0260</td>
</tr>
<tr>
<td>Rectangular$_1$</td>
<td>2.527</td>
<td>1.1734</td>
<td>2.15386</td>
</tr>
<tr>
<td>Rectangular$_2$</td>
<td>2.765</td>
<td>1.1632</td>
<td>2.37758</td>
</tr>
</tbody>
</table>
Table 4-5 $Q_M(V_G=0.60V)$, $C_G(V_G=0.60V)$ and $Q_M/C_G$ ratio at identical $Q_M(V_G=0.0V)$ for NWTs at $L_G=12nm$. With crystallographic channel orientation $<100>$.

<table>
<thead>
<tr>
<th>shape</th>
<th>$Q_M(\times10^6/cm)$</th>
<th>$C_G$ (fF/cm)</th>
<th>$Q_M/C_G(10^{17}/F)$</th>
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<td>1.0711</td>
<td>2.5564</td>
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</table>

4.8 Effect of Quantum Mechanics on SS and DIBL

Figure 4-11 depicts the effects of gate length on DILB (defined as $\Delta V_I/\Delta V_D$) and on the subthreshold slope (SS). Noticeable difference can be observed in the electrostatic integrity between the NWTs with different cross-sectional shapes, with the circular nanowire and the prolonged elliptical nanowire performing marginally better than the others. The quantum effects have less influence on DILB and SS compared to their impact on of $Q_M$ and $C_G$, where the difference is approximately 11%. Figure 4-12 and Figure 4-13 depict the gate length dependence of the error between the classical and QM estimate of DILB and SS respectively. The error is gate length dependent and is inversely proportional to gate length.
For DIBL, the error increases from few percent to approximately 7% with a decrease in gate length from 20 nm to ~10 nm. The equivalent error for SS increases from ~0.5% to ~3.4%.

Figure 4-11 Impact of the gate length on SS and DIBL for NWT with six individual cross-sections.
Chapter 4. Si Nanowire Transistors Design

Figure 4-12 Percentage error (%) when using DD simulation without quantum corrections for estimating DIBL.

Figure 4-13 Percentage (%) error when using DD simulations without quantum corrections for estimating SS.
4.9 **Optimum Cross-Sectional Aspect Ratio**

It has been clearly demonstrated that the cross-sectional shape of NWT’s gate has strong impact on gate capacitance $C_G$ and mobile charge $Q_M$ in the NWTs. It has been demonstrated that the NWT with $<110>$ channel orientation and elongated elliptical cross-section where the long diameter is parallel to the Si wafer surface exhibits the highest mobile charge for any given gate voltage. The major to minor axis ratios (cross-section aspect ratio [AR]) have been highlighted as a important design parameter determining the device performance. Nonetheless, experimentally NWTs are still typically produced in two versions: circular (or elliptical) NWT and nanosheet (or nanoslab) FET. Each version comes with its own advantages and disadvantages. The fundamental difference between these two versions remains the cross-sectional AR. However, critical design questions addressing the optimal NWT cross-sectional aspect ratio, remain unanswered.

In an attempt to remedy this situation, the effect of varying the AR of GAA NWT while keeping a constant cross-sectional area has been investigated. An interesting finding emerged from this investigation was that the observed signatures of isotropic/anisotropic charge distributions exhibit the same attributes as the golden ratio (Phi).

The structure, and dimensional characteristics associated with NWTs suitable for 5 nm Si CMOS technology will be investigated, along with the ratio of quantum mobile charge to gate capacitance, using the intrinsic delay ($\tau$) as a marker for the intrinsic speed and NWT performance. Quantum confinement effects are also considered to ensure the accuracy of the simulations.

The NWT structure comprises a single lateral NWT. To evaluate optimal AR, the NWT structure illustrated in Figure 4-3 is utilised. Here the Si channel is wrapped by a 0.4 nm SiO$_2$ IL and a 0.8 nm HFO$_2$ (High-k) layer in the gate region. The channel doping level is $1 \times 10^{14}/\text{cm}^3$, rising to $1 \times 10^{20}/\text{cm}^3$ in the extensions, and to $4 \times 10^{20}/\text{cm}^3$ in the source/drain regions. The x-axis is used for charge transport direction alignment. Table 4-6 outlines the cross-sectional dimensions for 9 elongated elliptical shape nanowires with matching cross-sectional areas of $10\pi \text{ nm}^2$. The NWT y-axis diameter ranges between 4.440 nm to 9.0 nm,
whilst the corresponding diameter along the z-axis ranges between 9.0 nm to 4.44 nm. Consequently, the associated aspect ratios vary between 2.020 to 0.490. Cross-sectional dimensions include wires representing Phi and 1/Phi (numeric) [186].

PS quantum corrections deliver a precise charge distribution within simulated NWT channels. Figure 4-14 depicts Phi ovals graphically represented on a 2D cross-section at the centre of the gate. The analysis of isotropic charge distributions resulting from the PS simulations exhibit identical characteristics to the golden ratio. Whilst charge distributions are voltage-dependent, phi oval signatures are observed at all gate voltages.

In order to make an unbiased assessment of the influence of NWT cross-sectional dimensions on device performances, Figure 4-15 illustrates the gate voltage dependence of the quantum mobile charge in the channel for the nine NWTs summarised in Table 4-6. Alignment of Q_M-V_G curves is achieved via amendment of the gate work function. The most interesting finding from the Q_M-V_G features illustrated in Figure 4-15 is that the NWT with Phi and 1/Phi aspect ratios deliver the highest quantum charge when compared with all other NWT scenarios. In addition, the lowest charge within the channel is associated with the NWT with a perfectly circular cross-section when compared with the charge in the channel of elliptical devices.

Table 4-7 and Table 4-8 compare the simulated gate capacitance (C_G) and the mobile quantum charge in the channel per-unit-length (Q_M) at specific gate voltages V_G for two crystal orientations, <110> and <100>. Eqn (4-5) has shown that Q_M is directly proportional to the NWT gate capacitance. Consequently, lowering NWT gate capacitance also results in a lower mobile charge within the channel and therefore poorer transistor performance. Both tables indicate that for either crystal orientation, the capacitance is greatest for the wires that conform to the golden ratio. In addition, for wires with the <100> channel orientation, the charge in the Phi and 1/Phi examples is practically the same due to identical effective masses in Y and Z directions [185].

Therefore, it could be suggested that the two recent research attempts to construct NWTs with cross-sectional AR≈1 (circular NWT [11] and AR≈3.5 (sheet NWT) [12], have not
taken into consideration the fact that AR is an important design factor determining the NWT performance (NB in this investigation, the term “sheet NWT” refers to NWT where 0.5 ≥ AR ≥ 2).

Figure 4-16 illustrates the $I_D-V_G$ characteristics of the simulated NWTs. Consistent with the data obtained thus far, the NWT with AR equal to Phi or 1/Phi exhibits the largest ON-current and this is valid at both low and high drain voltages.

Figure 4-17 highlights how the intrinsic delay ($\tau$) varies with the gate length. The intrinsic delay $\tau$ is directly inversely proportional to the speed of the device, where:

$$\tau = C_G \left( \frac{V_{DD}}{I_{eff}} \right)$$  \hspace{1cm} (4-6)

where $C_G$ represents the total gate capacitance, and the effective current is given by:

$$I_{eff} = \frac{I_H + I_L}{2}$$ \hspace{1cm} (4-7)

where

$I_H$= $I_D(V_G=V_{DD}, V_D=V_{DD}/2)$

$I_L$= $I_D(V_G=V_{DD}/2, V_D=V_{DD})$

$V_{DD}$=0.60V

And the leakage current is given by

$I_{off}$= 0.60 $\mu$A/$\mu$m
Figure 4-17 shows that the NTWs with dimensions 8.1 nm x 5 nm and 5 nm x 8.1 nm have the shortest intrinsic delay. The assessment of the intrinsic delay corroborates the results of $Q_M/C_G$ summarised in Table 4-7.

<table>
<thead>
<tr>
<th>$Z(\text{nm}) \times Y(\text{nm})$</th>
<th>$Y/Z$</th>
<th>$\text{Area (nm}^2\text{)}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.44 $\times$ 9.0</td>
<td>2.0200</td>
<td>10.0$\pi$</td>
</tr>
<tr>
<td>5.0 $\times$ 8.1 ($\Phi$)</td>
<td>1.6200</td>
<td>10.0$\pi$</td>
</tr>
<tr>
<td>5.7 $\times$ 7.0</td>
<td>0.8100</td>
<td>10.0$\pi$</td>
</tr>
<tr>
<td>6.0 $\times$ 6.66</td>
<td>0.900</td>
<td>10.0$\pi$</td>
</tr>
<tr>
<td>6.32 $\times$ 6.32</td>
<td>1.00</td>
<td>10.0$\pi$</td>
</tr>
<tr>
<td>6.66 $\times$ 6.0</td>
<td>1.110</td>
<td>10.0$\pi$</td>
</tr>
<tr>
<td>7.0 $\times$ 5.7</td>
<td>1.220</td>
<td>10.0$\pi$</td>
</tr>
<tr>
<td>8.1 $\times$ 5.0 ($1/\Phi$)</td>
<td>0.620</td>
<td>10.0$\pi$</td>
</tr>
<tr>
<td>9.0 $\times$ 4.44</td>
<td>0.490</td>
<td>10.0$\pi$</td>
</tr>
</tbody>
</table>

*Table 4-6 Physical dimensions of the cross-section of simulated NWTs*
Figure 4-14 The charge distribution in the 5.0 nm× 8.10 nm NWT cross-section (obtained from the Poisson-Schrödinger simulations) and Phi ovals (white ovals).
Table 4-7 $Q_M/C_G$ ratio at identical $Q_M$ ($V_G=0.0V$) for NWTs with crystallographic channel orientation <110>.

<table>
<thead>
<tr>
<th>Z(nm)x</th>
<th>$Q_M$ ($\times 10^7$/cm)</th>
<th>$C_G$ ($10^{-10}$/F/cm)</th>
<th>$Q_M/C_G$ ($10^{17}$/F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;110&gt;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.44x9.0</td>
<td>2.733</td>
<td>1.140</td>
<td>2.3970</td>
</tr>
<tr>
<td>5.0x8.1</td>
<td>2.842</td>
<td>1.147</td>
<td><strong>2.4760</strong></td>
</tr>
<tr>
<td>5.7x7.0</td>
<td>2.750</td>
<td>1.129</td>
<td>2.4350</td>
</tr>
<tr>
<td>6.0x6.66</td>
<td>2.580</td>
<td>1.109</td>
<td>2.3260</td>
</tr>
<tr>
<td>6.32x6.32</td>
<td>2.548</td>
<td>1.109</td>
<td>2.2970</td>
</tr>
<tr>
<td>6.66x6.0</td>
<td>2.631</td>
<td>1.116</td>
<td>2.3560</td>
</tr>
<tr>
<td>7.0x5.7</td>
<td>2.773</td>
<td>1.135</td>
<td>2.4400</td>
</tr>
<tr>
<td>8.1x5.0</td>
<td>2.892</td>
<td>1.160</td>
<td><strong>2.4910</strong></td>
</tr>
<tr>
<td>9x4.44</td>
<td>2.727</td>
<td>1.137</td>
<td>2.3980</td>
</tr>
</tbody>
</table>

**Figure 4-15** Gate voltage versus the mobile charge in the mid of the channel for the NWTs with different cross-section aspect ratio and same cross-sectional area. The crystallographic channel orientation is <110> and $L_G=12$ nm.
Table 4-8 $Q_M/C_G$ ratio at identical $Q_M$ ($V_G=0.0V$) for NWTs with crystallographic channel orientation $<100>$.

<table>
<thead>
<tr>
<th>$Z(nm) \times Y(nm)$</th>
<th>$Q_M$ ($\times 10^7/cm$)</th>
<th>$C_G$ ($10^{-10}F/cm$)</th>
<th>$Q_M/C_G$ ($10^{17}/F$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;100</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.44x9</td>
<td>2.732</td>
<td>1.135</td>
<td>2.407</td>
</tr>
<tr>
<td>5.0x8.1 Phi</td>
<td>2.890</td>
<td>1.159</td>
<td>2.492</td>
</tr>
<tr>
<td>5.7x7.0</td>
<td>2.779</td>
<td>1.138</td>
<td>2.444</td>
</tr>
<tr>
<td>6.0x6.66</td>
<td>2.640</td>
<td>1.118</td>
<td>2.360</td>
</tr>
<tr>
<td>6.32x6.32</td>
<td>2.610</td>
<td>1.116</td>
<td>2.337</td>
</tr>
<tr>
<td>6.66x6.0</td>
<td>2.640</td>
<td>1.118</td>
<td>2.361</td>
</tr>
<tr>
<td>7.0x5.7</td>
<td>2.779</td>
<td>1.138</td>
<td>2.442</td>
</tr>
<tr>
<td>8.1x5.0</td>
<td>2.891</td>
<td>1.159</td>
<td>2.491</td>
</tr>
<tr>
<td>9x4.44</td>
<td>2.732</td>
<td>1.134</td>
<td>2.409</td>
</tr>
</tbody>
</table>

Oblate NWT

Circular

Prolate NWT

Figure 4-16 The impact of all 9 cross-section AR on the $I_D-V_G$ characteristics. Dashed curves are at $V_D=0.050V$, and solid curves are at $V_D=0.70V$. $L_G=12.0 \text{ nm}$ with $<110>$ crystallographic channel orientation.
A more systematic design of experiment would assess the way in which NWTs FOM interact with other parameters associated with AR. Figure 4-18 depicts the design of experiment, including the impacts of gate length, effective oxide thickness and NWT cross-sectional dimensions on $I_{off}$, $I_{on}$ and SS. The focus is on 5 nm NWT, three-dimensions ($4 \text{nm} \times 4.6 \text{nm}$, $5 \text{nm} \times 8.1 \text{nm}$ and $6 \text{nm} \times 6.9 \text{nm}$) with cross-sectional areas ($4.6\pi \text{ nm}^2$, $10.125\pi \text{ nm}^2$, and $10.35\pi \text{ nm}^2$) involving the golden ratio in the design of experiment. In this study only the $<110>$ channel orientation was investigated.

The results indicate that the highest drive current is observed for the NWT with the smallest gate length and oxide thickness. The SS mirrors the electrostatic integrity of the transistors. The NWTs possessing a long channel with narrow cross-sectional dimensions offer greater
gate control, resulting in a steeper SS. Reducing the nanowire cross-sectional area from 41.5 nm² to 18.56 nm² (while keeping the same cross-sectional aspect ratio) offers in the region of 1.4mV/decade and 8mV/decade SS for respective gate lengths of 18nm and 12 nm.

Notwithstanding these marginal electrostatic improvements, drive current of the device is diminished by 38%. Reducing the gate length while increasing the NWT cross-sectional area can raise the drive current, but then the influence of parasitic effects becomes a significant issue leading to reduced performance when compared with larger NWT based

Figure 4-18 Experimental design (4D) for NWT and the effects of the NWT’s dimensions, Oxide thickness, and Lg on Ioff, DIBL, VT, and Ion.
devices. The NWT with gate dimensions 12nm-16nm exhibits an acceptable trade-off between the $I_{on}$, $I_{off}$ and SS.

4.11 Conclusions

The quantum mechanical impact on the electrostatic performance of NWTs suitable for CMOS technologies at and beyond 5-nm mark have been investigated. It has been highlighted how the shape of NWT influences the gate capacitance and the mobile charge in the channel. Furthermore, circular and elliptical NWT, with channel orientations <110> and <100>, have been investigated showing enhanced performance (with respect to electrostatic driven performance and ‘intrinsic’ ($Q_M/C_G$) ratio) when compared with square and rectangular ultra-scaled GAA NWTs. For instance, the circular and elliptical nanowires exhibit a greater quantum mobile charge ($Q_M$) in the channel and also a higher ($Q_M/C_G$) ratio compared to the square and rectangular nanowires. In addition, the results show that the 2D charge profile and the quantity of charge in the channel depend on the channel direction. For instance, all nanowires with the <100> channel have approximately ~11% greater mobile charge compared to the <110> nanowires. Such findings are important for determining the optimal designs of NWTs. Signatures of isotropic charge distributions within Si NWTs were observed exhibiting the same attributes as the golden ratio (Phi), the significance of which is well known in the fields of art and architecture.

The research demonstrates that the quantity of mobile charge within the channel, along with the intrinsic speed of the device are determined by device geometry and may be affiliated with the golden AR (Phi) of the nanowire transistors. This investigation has demonstrated NWT with aspect ratios equal or close to the golden ratio (Phi) can enhance gate capacitance and mobile charge in the channel and therefore could optimise the intrinsic speed of the device. We have also investigated the influences of the gate length on the time delay and the main FoM, such as $V_T$, $I_{OFF}$ and $I_{ON}$ and DIBL.
Chapter 5. Performance of Si NWTs

5.1 Introduction:

The semiconductor industry must continually cope with performance eroding factors including short-channel effects, high leakage current, and other issues associated with highly-scaled planar metal-oxide-semiconductor field-effect transistors (MOSFETs) [74], [190]. The Fin field effect transistor (FinFET) has been developed in an attempt to overcome these issues, with 7 nm FinFETs technology presently receiving much attention in research and development [191]. Whilst it is necessary to reduce FinFET size in order to retain scaling [4], statistical and process variability both increase as a result of the reduced FinFET dimensions [5]. The difficulty involved in FinFETs scaling increase with the Fin shape and geometry becoming difficult to manage when the fin is made slimmer and elongated in an effort to maximise drive current. Nanowire transistors (NWTs) have therefore been proposed as a potential FinFET, replacement [6]. The next-gen technology node could indeed be served by the gate-all-around NWTs as an alternative to FinFETs due to their superior electrostatic integrity [88].

The semiconductor industry is now focusing on the possibility of a 15% increase in the saturation current ($I_{sat}$) needed beyond 7 nm technology nodes in to deliver the expected performance increase. For the 5 nm technology transistor, there is an approximate 1.6 mA/µm $I_{sat}$ target, with Table 5-1, Figure 5-1, and Figure 5-2 presenting three technology nodes alongside the 14 nm FinFET.
Previous research [10], [11], [192] addresses the potential to increase the single NWT’s performance issues through the fabrication of a single transistor with two lateral nanowire channels. In order to increase the drive current, taller fins must be fitted to the 7 nm FinFET chip. However, the vertically-stacked lateral Si NWT would have a smaller chip footprint and would be shorter, compared to the FinFET counterparts.

![Figure 5-1 Comparison of Intel’s 14 nm FinFET and 5 nm NWT with one, two, and three lateral channels.](image)

For each new generation, a 15% increase in saturation current could be achieved using Si NWTS, as a result of their electron transport characteristics. Performance improvements can also be realised through the use of strain in the channel; through the inclusion of multi-lateral channels within devices; and by engineering the channel orientations, cross-sectional geometries and other design parameters of the transistors. Furthermore, device structure engineering could also provide better balance between leakage currents and performance.

Much of the existing literature has explored only single-channel NWT in simulation, with very few studies simulating multi-lateral 5 nm node NWTS at the time of conducting this research. For this reason, this chapter attempts to address this shortcoming and discuss the simulation of NWTS with multiple channels and various channel lengths. Contact resistance, non-equilibrium transport and quantum confinement effects are all taken into account during
the simulations in order to provide reliable and accurate results appropriate for highly-scaled NWTs. Importantly, this chapter also aims to address two main questions: firstly, whether a single-channel silicon (NWT) at 5 nm technology node is able to achieve the semiconductor industry target; and, secondly, if this is not possible, what number of nanowire channels are required in a single NWT is in order to reach this target.

The following section of this chapter describes the device, whilst the simulation methodology is presented in Section 5.3. Section 5.4 discusses the calibration methodology. The results are presented in Section 5.5 and Section 5.6. Finally, conclusions from the simulation are presented in Section 5.7.

![Figure 5-2 Saturation current scaling target from 14 nm to 5 nm technology.](image)

<table>
<thead>
<tr>
<th>Node</th>
<th>14 nm</th>
<th>10 nm</th>
<th>7 nm</th>
<th>5 nm</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Layout pitch</strong></td>
<td>42nm</td>
<td>29nm</td>
<td>21nm</td>
<td>14nm</td>
</tr>
<tr>
<td>$I_{sat} \times 15%$ (mA/um)</td>
<td>1.040</td>
<td>1.1960</td>
<td>1.3750</td>
<td>1.580</td>
</tr>
</tbody>
</table>

*Table 5-1 Layout pitch and corresponding saturation current for (14-5nm) technology.*
5.2 Device Description:

The previous chapter elucidated the relationship between performance, electrostatic integrity, quantum effects and cross-sectional geometry in highly-scaled NWTs. The results demonstrate that the strongest performance is found in NWTs with elliptical cross-sections compared to the performance NWTs with circular and square cross-sections.

The current section analyses the performance of Si n-channel gate-all-around NWTs with elliptical 7 nm x 5 nm and 8.1 nm x 5 nm cross-sections. Each of the simulated devices has a 0.4nm SiO$_2$ interfacial and 0.8nm HfO$_2$ (High-k) gate dielectric layers. As illustrated in Figure 4-5 and 4-6, doping concentrations of $10^{14}$/cm$^3$, $10^{20}$/cm$^3$ and $4\times10^{20}$/cm$^3$ were used in the channel, extensions and source/drain, respectively. The study includes simulation of 10 nm, 12 nm, 16 nm and 20 nm gate length NWTs, with single-, double- and triple-channels. The source/drain contacts were set at the tops of the devices in each case (see Figure 4-5) for overall device structure.

5.3 Ensemble Monte Carlo (EMC) Simulation:

Charge transport and performance of a single nanowire NWT was analysed using Ensemble Monte Carlo (EMC) simulations including band structure, relevant scattering mechanisms [193], [194], volume inversion [195], and other physical relevant for Si NWT scaled below 14 nm. As a result of the impact of confined acoustic phonon and body thickness variation-induced scattering [101], a rapid reduction in mobility been observed below 4 nm. Extensive downward scaling of NWT thickness has also been achieved, with a near-5 nm nanowire diameter. Significant quantization effects can observed in the nanowire cross-section, as a result of a high electric field and small confinement dimensions [196]. Consequently, discrete sub-bands are formed from the conduction band, with the maximum charge density trends shifting towards the channel centre and away from the metal-oxide interface. These quantum effects impact device properties and should therefore be taken into account when analysing device performance. Thus, in this study, the impact of quantum confinement...
effects on saturation current are explored using a self-consistent three-dimensional quantum mechanical Monte Carlo (3D-MC) simulator. The primary benefit of quantum-corrected 3D-MC simulation approach is the ability to simulate the quantum effects across the entire NWT structure. Furthermore, this approach also takes into account the transition of electrons between quantized states, resulting in greater accuracy when self-consistently calculating nonequilibrium electron distribution and current.

The MC method provides a direct solution to the Boltzmann transport equation (BTE), and therefore can capture non-equilibrium, quasi ballistic transport in nanometre scale semiconductor devices. It is suitable to apply to the simulation of Fin-FET, nanowire transistors and other novel 3D transistors. Furthermore, it is more accurate and predictive compared to drift-diffusion (DD) simulations. The smaller the transistor the more non-equilibrium the transport becomes with carrier heating [197] and with an increasingly significant velocity overshoot as a non-local effect.

The Ensemble Monte Carlo (EMC) [198] approach with Poisson-Schrödinger (PS) quantum corrections was adopted for the purpose of these simulations. As mentioned before the Monte Carlo approach provides a direct solution to the BTE. The method takes into account key electron scattering mechanisms such as intra- and inter-valley electron-phonon scatterings, surface roughness scattering, ionised impurity scattering, and the stochastic characteristics of electron transport. The k·p method was also utilised to calculate a full-band structure, which was approximated with and analytical band structure model comprised of ellipsoidal non-parabolic valleys. The movement of hundreds of thousands of particles within a field distribution was tracked in order to achieve self-consistent simulation, with Poisson’s equation solved in 3D every 0.50 fs. The 3D-EMC simulation began with initial conditions set as the DD solution and the corresponding potential profiles and charge distribution. During simulation, self-consistent time-varying electrostatic potential and field distributions were upheld through the solution of the linear Poisson equation and the application of QM correction. In highly-scaled channel transistors, the 3D-EMC simulation approach has been shown to offer high levels of accuracy in forecasting ON-state transistor performance and in the handling of non-equilibrium transport [199].
The $I_D$-$V_G$ characteristics obtained from the DD and MC-based NWT simulations (see Section 3.2) are illustrated in Figure 5-3. In the case of highly-scaled NWTs, DD-based simulation is unable to capture the non-equilibrium effects, although many recognise the efficiency and stability offered by the approach in alternative scenarios, with DD simulation being for large complex structures including external contacts.

The results of the MC simulation deliver drain current of $I_D = 1.16$ mA/µm at high drain bias, with a marginally lower drain current than the DD simulation results at low drain bias. The results demonstrate that the DD simulation does not provide the desired accuracy in forecasting ON-state transistor performance despite the incorporation of density gradient quantum correction for charges. The DD and quantum-corrected MC simulation subthreshold slopes are similar. DD calibration is then performed in respect of the quantum-corrected MC simulation results. This step allows the DD approach to fully capture the MC/PS simulation results. The calibrated DD model then can be used to simulate complex multichannel NWTs and to carry out statistical variability and reliability simulation, as well
as a simulation of the interplay between these two elements (see Figure 5-4 for a high-level simulation flowchart).

![Drift-Diffusion Simulation](image)

**Figure 5-4 The simulation tool calibration flow chart.**

### 5.4 Drift Diffusion Calibration

Whilst the MC approach is widely recognised as being amongst the top transistor simulation approaches in terms of accuracy, it places immense strain on the CPU usage [200]. Furthermore, the computational speed, efficiency and flexibility of the DD method has been found to be supplementary to the accuracy of the MC simulation approach. For instance, using a 3.00GHz Intel Xeon 16 Core CPU, simulation of each bias point of Figure 5-3 was achieved in just 58 minutes using the DD approach, and 96 hours using the MC approach (1% error, mesh = 0.2).
Drift diffusion calibration is conducted in order to identify the DD simulation parameters suitable for reducing the difference between the results provided by EMC simulations. Here, the density gradient confinement effective mass, gate work function, and mobility models with their corresponding parameters are amongst the parameters selected for calibration.

\[ \text{ID-VG calibration is conducted based on a three-stages strategy, with the first stage targeting the sub-threshold region, the second stage targeting the low field current voltage characteristics, and the third stage targeting the high field current voltage characteristic. Here, the impact of each parameter calibration stage is virtually independent on the following stages. The mobility models and additional parameters along with the relevant I-V characteristic areas that they most significantly influence are illustrated in Table 5-2.} \]

<table>
<thead>
<tr>
<th>Parameter / Mobility models</th>
<th>Major impact on I-V region</th>
</tr>
</thead>
<tbody>
<tr>
<td>Density gradient effective mass</td>
<td>Subthreshold; Low drain (LD) and high drain(HD)</td>
</tr>
<tr>
<td>Work-function</td>
<td>All; shift in VG</td>
</tr>
<tr>
<td>Low-Field mobility models</td>
<td>Low drain (LD); above threshold</td>
</tr>
<tr>
<td>Perpendicular-Field mobility models</td>
<td>Low drain (LD)</td>
</tr>
<tr>
<td>Lateral-Field mobility models</td>
<td>High drain(HD)</td>
</tr>
</tbody>
</table>

An oxide interface is found in the NWT structure during the calibration first stage, with a density gradient quantum correction carrier confinement effective masses normal z-direction (m^{dz}) and y-direction (m^{dy}). These affect the subthreshold slope (SS) by way quantum confinements’ decrease in effective oxide capacitance. This being said, since the PS-
calibration of the density gradient effective masses has already been carried out, it does not need to be changed at this stage. Modification of the work function at any of the calibration stages realigns the $V_T$.

In the (Poisson-Schrödinger (PS) / Drift-Diffusion (DD)) simulation, up to three mobility models can be selected to use in a given simulation: 1) Low-field mobility (that considers ionized impurity scattering); Arora mobility model [149], Masetti mobility model [148], and Philips mobility model [150]. 2) Perpendicular field-dependent mobility (that considers surface roughness scattering); Yamaguchi [151], Lombardi [152], and thin-layer mobility model [153]. 3) Lateral field-dependent mobility (for velocity saturation); Caughey-Thomas velocity saturation model [154]. In this work, the following three carrier mobility models have been used, Masetti, Lombardi, and Caughey-Thomas. At low $V_D$, the NWT $I_D-V_G$ characteristics are determined by the low-field mobility for a gate voltage around and just above $V_T$, then the device calibration process is the calibration of the complete $I_D-V_G$ characteristics at low drain bias. During the 1st calibration iteration a perpendicular electric field model has been activated and the lateral electric field model disabled. In the final stage of the calibration process, the complete $I_D-V_G$ behaviour at high applied $V_D$ is calibrated. During the first calibration iteration a lateral electric field model has been enabled and the perpendicular electric field model disabled. At all stages of the calibration the $V_T$ can be realigned by modifying the work function.

5.5 5nm NWT: MC simulation

Simulations of the $I_D-V_G$ characteristics of the NWTs under consideration are carried out following the aforementioned methodology. The $I_D-V_G$ characteristics of a 14 nm FinFET transistor is compared to that of a single Si-NWT simulation in Figure 5-5, with the results benchmarked against the 5 nm CMOS’ target drive current of 1.58 mA/µm. This was justified during the introduction. Alignment of the two characteristics provides the same leakage current. The results demonstrate that the 14 nm FinFET has a lower $I_{sat}$ than the single nanowire transistor at both high and low drain biases. Here, $SS$ is approximately 62.55
[mV/dec], while DIBL is approximately 9.04 [mV/V]. Although the drive current of the single NWT is higher than the drive current of the FinFET, the $I_{\text{sat}}$ value falls significantly short of the 1.58 mA/µm target at high (0.7V) drain bias (Figure 5-5). It should be highlighted that all currents shown in Figure 5-5 are normalised by the transistor pitch (see Table 5-1).

![Figure 5-5 I_D-V_G characteristics of Intel’s 14 nm FinFET (experimental) comparable to the performance of 5 nm Si NWT (3D-MC simulations). Lg=12nm for the NWT. Dashed, solid curves represent I_D-V_G characteristics at drain voltage 0.070V and 0.050V respectively.](image)

5.6 Strain and channel orientation effects

Chip manufacturing is based on silicon, despite the potential use of Ge, GaAs, InGaAs, InAs [201] and other high-mobility materials for the enhancing of the NWT performance. The performance of silicon-based transistors can be improved by suitable surface orientation and channel direction choices. The Si NWT performance and mobility can also be further enhanced through the introduction of strain [202]. Therefore, identifying solutions to
achieve the highest strain levels possible at the present time is a high priority for the semiconductor industry. Recent research reports a successful attempt to move significantly beyond the limits of SiGe-based virtual substrates by using a 4.5% elastic strain of 7.6 GPa uniaxial tensile stress in Si nanowires with a width of 30 nm [203]. In traditional FETs, high STI-based compressive stress result in a saturation current of scaled <110> nFETs being close to <100> nFETs. The velocity saturation of <110> pFETs is lower than that of nFETs, with mobility also being higher in <110> pFETs compared to <100> pFETs, rendering the scaled <110> pFETs more preferable [204]. Significant research is focused on <110> surface orientation, which has been heavily adopted in gate-all-around FETs, FinFETs and other FET structures [205].

Extreme downward scaling of NWT diameter to 5 nm has been achieved. Significant quantization effects can be seen in the corresponding NWTs., [196]. Consequently, discrete sub-bands are formed from the conduction band, with the maximum charge density shifting towards the channel centre and away from the metal-oxide interface. It is also possible for the charge density to have an isotropic distribution. These quantum effects have an impact on device performance and must therefore be taken into account during device simulation analysis. Therefore, this study investigates the impact of the quantum mechanical effects on the NWTs saturation current using a self-consistent three-dimensional quantum mechanical Monte Carlo (3D-MC) simulation approach. The ability of the quantum-corrected 3D-MC approach to capture accurately the quantum effects is one of the main advantage of this approach.

We begin by analysing 7 nm x 5 nm NWTs with channel orientations of <110> and <100> with regards to strain impact. Here, axis rotation necessitates that each valley’s effective mass tensor must be converted, and thus the impact of switching between the <110> to <100> channel orientation without strain must be assessed first.
Figure 5-6 $I_D$-$V_G$ characteristics compares the simulation results (MC) for two stacked channels (7nm x 5nm) each NWT has a different channel orientation $<110>$ and $<100>$. $L_g=12$nm. Dashed, solid curves represent $I_D$-$V_G$ characteristics at drain voltage 0.070V and 0.050V respectively. The gate length is 12 nm.

Figure 5-6 presents the $I_D$–$V_G$ characteristics of (7nm × 5nm) Si NWT with channel orientations of $<110>$ and $<100>$ without strain. The mobile charge is higher in the $<100>$ channel orientation than in the $<110>$, in line with the chapter 4 results. Therefore, the channel’s mobile charge increases when the crystal orientation of the silicon channel is changed. Furthermore, a 4% increase in saturation current is shown in the $<100>$ NWTs compared to the $<110>$ NWTs. The cross-sectional area and shape are the same in both NWTs.

The $I_D$-$V_G$ characteristics for a 90° NWT rotation are presented in Figure 5-7. The $I_D$-$V_G$ results of the $<110>$ NWT are largely the same as for the $<100>$ NWT and unrotated NWT, due to symmetrical charge distribution (as result of identical electron effective masses in the principal confinement plane). Anisotropic charge distribution occurs in the case of the $<110>$ NWT due to the differences between the electron effective masses. The results indicate that NWT performance can be enhanced by changing the $<110>$ channel orientation, with NWT performance significantly impacted by its geometry.

The enhanced transport and superior electrostatic control of strained Si nanowires make them viable candidate for further CMOS scaling. Consequently, there is significant interest
in increasing the NWT performance by strain. The introduction of strain can result in significant $I_{sat}$ current enhancement. However, further research is necessary to explore this possibility at NWT cross sections below 7 nm, taking into account also the quantum confinement effects[206].

The key benefits of strain include reduction in the effective masses and increase of the band separation all increasing mobility [207], [208]. The impact of strain on Si is discussed in the existing literature [209]. However, the valley splitting due to quantum confinement can minimise strain effect in transistors scaled transistors. Figure 5-8 illustrates conduction band constant energy ellipsoids ($\Delta$) with six valleys in silicon across the confinement plane and uniaxial tensile strain, with the x-axis denoting transport direction.

The Figure 5-9 and Figure 5-10 illustrate the $I_D-V_G$ characteristics of four single NWTs with varying strain values and a <110> and <100> channel orientation respectively. The results indicate that a 5-27% increase in $I_{sat}$ magnitude and 3-21% increase in saturation current can be achieved through the introduction of channel strain in the <110> and <100> channel orientations, respectively. This being said, strain of 2.0 GPa still did not allow us to achieve the target drive current in either case.
Within the crystal unit cell, atom location is impacted by tensile strain, resulting in a change in effective mass conductivity and band structure, with strained semiconductors demonstrating changes in band edge curvature, band warping, band splitting and band edge energies.

![Diagram showing conduction band constant energy ellipsoids (Δ) six valleys in silicon along the confinement plane and uniaxial tinsel strain. The transport direction is the x-axis.](image)

*Figure 5-8 conduction band constant energy ellipsoids (Δ) six valleys in silicon along the confinement plane and uniaxial tinsel strain. The transport direction is the x-axis.*
Figure 5-9 $I_D-V_G$ characteristics compares the simulation results (MC) for 4 (7nm×5nm) NWT with different level of tensile strain channel orientations $<110>$. $L_g=12$nm. Dashed, solid curves represent $I_D-V_G$ characteristics at drain voltage 0.070V and 0.050V respectively.

Figure 5-10 $I_D-V_G$ characteristics compares the simulation results (MC) for 4 (5nm×7nm) NWT with different level of tensile strain channel orientations $<100>$. $L_g=12$nm. Dashed, solid curves represent $I_D-V_G$ characteristics at drain voltage 0.070V and 0.050V respectively.
5.7 Vertically stacked lateral NWT

Whilst more difficult to simulate and manufacture, introducing multiple channels in a single NWT is one potential solution to enhance $I_{sat}$. Per-footprint drive current deficiency could be addressed through the development of vertically-stacked lateral NWTs at the 5 nm node, with reduced contacted gate pitch [11], [12]. The ITRS roadmap also incorporates vertically-stacked NWTs [210]. The difficulty in simulating vertically-stacked NWTs stems from its structure size, which remains an issue even with the adjustment of the discretization mesh. Direct simulation of multiple channel NWTs is computationally prohibitive using MC simulation. The MC simulations cannot also take into account the contact resistance. Therefore single channel NWT MC simulations with PS-based quantum correction were performed for the purpose of calibrating a single NWT’s DD simulations needed to assess the vertically-stacked NWT performance [211], [212]. Once satisfactory calibration has been achieved, the vertically-stacked lateral NWTs were simulated using the calibrated DD. The vertically-stacked NWT with 7 nm x 5 nm elliptical channel is illustrated in 3D in Figure 5-12.

Single-, double- and triple-channel devices of four gate lengths were compared and analysed as illustrated in Figure 5-11. It was not possible to reach the target drive current in the single channel NWT devices, meaning that the 5 nm CMOS technology’s scaling requirements could not be met. For instance, double-channel devices – discounting the source/drain (S/D) contact resistance – can exceed the target drive current in the case of an $L_G=12$ nm device, but this cannot be achieved at $L_G=20$ nm.

Based on these results, the target performance is achievable through the use of a double-channel lateral NWT. However, this is only possible if the S/D contact resistance is not taken into account and the gate length is of a certain size.
Compared to the 5 nm CMOS node target value of (15%) $I_{\text{sat}}$ current values are significantly greater for the triple-channel lateral nanowires, as shown in Figure 5-11. Therefore, the results indicate that the 5 nm CMOS scaling performance target could be safely achieved through the introduction of three lateral channels stacked in a single NWT device. Still these results do not take into account the S/D contact resistance.

![Figure 5-11](image)

**Figure 5-11** Characteristics of the $I_D$ versus the $V_G$ obtained from calibrated DD for the NWTs with one, two, and three channels. The DD is calibrated for each gate lengths (10nm, 12nm, 16nm, and 20nm). Dashed, solid curves represent $I_D$-$V_G$ characteristics at drain voltage 0.070V and 0.050V respectively.

![Figure 5-12](image)

**Figure 5-12** 3D schematic view of the vertically stacked NTW of two lateral channels and material details.
5.8 Source/drain contact resistance

As noted, source/drain contact resistance (R) is not taken into account in the simulation illustrated in Figure 5-11. In the simulation analysis presented here four values of R considered as lump S/D resistance (100 Ω, 500 Ω, 750 Ω, and 1K Ω). The contact resistance is introduced at both the source and drain contacts (Figure 5-12).

The results illustrated in Figure 5-13, which does not include strain, illustrate the negative impact of the S/D resistance on $I_{\text{sat}}$. For example, the $I_{\text{sat}}$ value is approximately 0.82 mA/µm for single-channel NWTs, 1.49 mA/µm for double-channel NWTs and 1.92 mA/µm for triple-channel NWTs at a drain bias of 0.7V and contact resistance of 1KΩ. Interestingly, the double-channel NWTs also fall below the target $I_{\text{sat}}$ current. Therefore, the introduction of contact resistance into the simulations suggests that the 5 nm CMOS scaling requirements cannot be met using double-channel NWTs. Conversely, the target value is exceeded by triple- channel transistors, suggesting that performance at or above the scaling target can only be achieved using triple-channel Si NWTs.

Figure 5-14 illustrates an experimental design for single-, double- and triple-channel NWTs with 10 nm – 20 nm gate lengths at a contact resistance varying from 500 Ω to 2500 Ω. The devices presented in Figure 5-13 are shown again in Figure 5-15. However, in this case, channel strain of 2.0 GPa is introduced. Here, the double-channel NWT exceeds the target $I_{\text{sat}}$ current across all R values, indicating that the industrial target could be achieved using with two channels NWT.

As shown in Figure 5-9, a single-channel NWT’s drain current can increase by 27% as a result of 2.0 GPa strain. However, Figure 5-15 demonstrates that the double- and triple- channel lateral NWTs show a significant decline in the single channel drive current, which is the result of a current path voltage decrease across series resistance at the S/D contacts, and in S/D regions that are highly doped. In the double-channel NWT the bottom nanowire current decreases by 21.6% at 1K Ω, whilst the triple-channel NWT the current decreases by 23.5% at the same resistance, between the channel and the contact.
Chapter 5. Performance of Si NWTs

Current density: series resistance effect

The single-, double- and triple-channel NWTs’ current density is illustrated in 3D in Figure 5-16, which highlights the variance between current densities. Whilst the nanowires in the triple-channel NWTs have the same cross-sectional area, current density is higher in the top channel (nearest to the metal) compared to the lower and middle channels. This is represented by the large red region near to the drain, with the red becoming weaker at the second and third channels. Similar results are depicted for the double-channel device, with greater current density in the top channel compared to the bottom channel. Therefore, the results suggest that as the distance between the channel and the contact becomes greater, a reduction in current density is experienced in each of the lateral channels.

Figure 5-13 Characteristics of the $I_D$ versus the $V_G$ obtained by calibrated DD for the NWTs with one, two, and three channels at four different ohmic contact resistance. $L_g=12\,\text{nm}$. Dashed, solid curves represent $I_D-V_G$ characteristics at drain voltage 0.070V and 0.050V respectively.

5.9 Current density: series resistance effect

The single-, double- and triple-channel NWTs’ current density is illustrated in 3D in Figure 5-16, which highlights the variance between current densities. Whilst the nanowires in the triple-channel NWTs have the same cross-sectional area, current density is higher in the top channel (nearest to the metal) compared to the lower and middle channels. This is represented by the large red region near to the drain, with the red becoming weaker at the second and third channels. Similar results are depicted for the double-channel device, with greater current density in the top channel compared to the bottom channel. Therefore, the results suggest that as the distance between the channel and the contact becomes greater, a reduction in current density is experienced in each of the lateral channels.
The metal source/drain contacts are at the top of the highly doped source/drain regions, which are the same for every NWT configuration. However, it is only the top of the source/drain region that connects with the contact. Thus, the top channel has greater current density due to proximity to the metal contact (see Figure 5-16).

The current density of lateral NWT stack containing NWTs with varying cross section of 7 nm × 5 nm (top), 7.4 nm × 5 nm (middle), and 8 nm × 5 nm (bottom) is illustrated in Figure 5-17. Channels with larger cross-sections are able to conduct stronger currents than their smaller counterparts, thus resulting in asymmetrical current density through the channels. Therefore, the same current could be sent through the stack from the top and bottom channels, at least in theory.
Figure 5-15 Characteristics of the $I_D$ versus the $V_G$ obtained by calibrated DD for NWTs with one, two, and three channels. Each NWT strained by 2.0 GPa. Considering 4 different ohmic contact resistance. $L_g=12$nm. Dashed, solid curves represent $I_D$-$V_G$ characteristics at drain voltage 0.070V and 0.050V respectively.

Figure 5-16 Current density for NWTs with one, two, and three channels $L_g=12.0$nm.
Figure 5-17 (Top) the current density of NWT with three stacked channels. Cross-section. (Bottom) Slicing the S/D contact up to the middle of the stack.

Figure 5-18 (A) the current density of stack with three lateral NWTs: top 5nm×7nm, 5nm×8.6nm (middle), and 5nm×9.4nm bottom. (B, C, D) the current density of single (5nm×7nm) NWT in the top, middle, the bottom of the stack.
This theory, however, is refuted in Figure 5-17 where, it can be seen that current density cannot be completely equilibrated by upscaling the diameter of the lower nanowires, with the top and middle channels still having higher current density than the bottom channel. Additionally, the figure also demonstrates that current density varies even when a single-channel NWT is used in place of the stack (see B, C and D). This is due to the impact of the increasing access length, and corresponding increase in the resistance of the highly-doped S/D regions.

As shown in Figure 5-18, bringing the metal contacts to the side could be an option to reduce the S/D series resistance, but the differences between the vertical placement of lateral nanowires and the respective lateral NWT diameters can have an impact on transistor variability.

Whilst the use of stacked multi-lateral NWTs has been proposed as a solution, the nanowires’ diameters will impact the drive current, with the drive current increasing with the increase in the NWT diameter. Assuming that the cross-sectional area is increased by the same ratio, the following equation can be used to estimate the area:

\[
Area = x(a/2) \times (b/2) \times \pi \quad a = 7nm, \ b = 5nm, \ and \ x \quad (5-1) \\
> 1
\]

Here we investigate further the options the NWT based 5 nm CMOS node with integrating double or triple lateral NWTs whilst increasing the vertical diameter \(x \times 7nm, x > 1\). Here, the cross-sectional area will be larger at the same footprint, but the device’s electrostatic integrity will be affected. Figure 5-19 illustrates the increase in a single NWT’s cross-sectional area to double its size, which is equal to the stacked (double) NWT’s total cross-sectional area (Si).

The double NWT’s cross-sectional area can be expressed as:
\[ A_{\text{Double}} = 2 \times \frac{5}{2} \times \frac{7}{2} \times \pi = 17.5\pi \ (nm^2) \]

The tall single NWT’s cross-sectional area can be expressed as:

\[ A_{\text{Tall}} = \frac{5}{2} \times \frac{14}{2} \times \pi = 17.5\pi \ (nm^2) \]

Based on the results of the simulation, single increase area NWTs of the same gate length are found to have worst SS and DIBL compared to the double NWTs. Furthermore, double NWTs are found to be better in terms of the total quantum charge to gate capacitance ratio than single NWTs of the same (Si) cross-sectional area.

**Figure 5-19** Single NWT with the same area of the double NWT. The current density and charge distribution at \( V_g=0.7V, \ V_D=0.7V, \) and \( L_g=12nm \).

### 5.10 Single-, double- and triple-channel NWT capacitance

The simulation conducted in this study computes nanowire capacitances as \( C_{ij} = \frac{\partial Q_i}{\partial V_j} \), where \( i \) and \( j \) represent the four device contacts: gate (g), source (s), drain (d) and bulk (b). Here, bulk capacitance is excluded from the analysis. The gate capacitances related to the gate contact are expressed as \( C_{gg}, C_{sg}, C_{dg} \) and \( C_{bg} \); the source capacitances related to the
source contact are expressed as $C_{gs}$, $C_{ss}$, $C_{ds}$ and $C_{bs}$; and the drain capacitances related to the drain contact are expressed as $C_{gd}$, $C_{sd}$, $C_{dd}$ and $C_{bd}$

$$C_{ij} = \partial Q_i / \partial V_j = \begin{bmatrix} C_{gg} & C_{gs} & C_{dg} \\ C_{gs} & C_{ss} & C_{ds} \\ C_{dg} & C_{ds} & C_{dd} \end{bmatrix}$$

Single channel NWT  $C_{ij} = \begin{bmatrix} 9.83 & 5.28 & 4.47 \\ 8.07 & 5.19 & 2.72 \\ 1.56 & 0.13 & 1.84 \end{bmatrix} \times 10^{-18} F$

Double channel NWT  $C_{ij} = \begin{bmatrix} 19.4 & 10.3 & 8.86 \\ 15.2 & 9.35 & 5.12 \\ 4.20 & 1.09 & 4.19 \end{bmatrix} \times 10^{-18} F$

Trible channel NWT  $C_{ij} = \begin{bmatrix} 27.9 & 12.4 & 13.2 \\ 20.9 & 10.7 & 7.95 \\ 7.08 & 0.21 & 6.21 \end{bmatrix} \times 10^{-18} F$

With the stacking capacitance of NWT rapidly increasing, it is not only $I_{on}$ criteria that should be considered when using laterally-stacked NWs. This is particularly true based on the fact that modern CMOS technology is more often restrained in terms of power than speed. Nonetheless, NWTs, FinFETs and other multi-gate MOS-FET must still consider the issue of parasitic capacitances. A potential solution is the use of a diamond-shaped 3D multi-gate transistor [213]

### 5.11 Optimization of 5 nm NWT

The connection between the cross-sectional dimensions of NWTs and electrostatic integrity was discussed in details in Section 4.8 of this study, which also highlighted that the lowest intrinsic delays can be found in NWTs of $8.1\text{nm} \times 5\text{nm}$ and $5\text{nm} \times 8.1\text{nm}$. These results are based on the $Q_M/C_G$ criteria.

The failure to meet the expected scaling performance NWTs has been highlighted, in terms of $I_{on}$ per lay-out pitch. It has been suggested that the introduction of strain to the channel
can enhance Si NWTs’ electron transport, together with altering the channel orientation and geometry of the NWT [189], [176], [206].

The simulation of $I_D$-$V_G$ characteristics for a $<110>$ channel orientation 5 nm $\times$ 8.1 nm (Phi) NWT for four levels of strain is illustrated in Figure 5-20, which confirms that 5-27% increase in the saturation current magnitude occurs with the increase of the channel strain. In particular 27% increase in NWT performance (compared to the unstrained NWT) has been observed at a tensile strain of 2.0 GPa. Still the 5 nm CMOS industrial target is only just met by the $I_{on}$ value, with the S/D contact resistance excluded from this analysis.

The results clearly demonstrate greater $I_{on}$ in the 1/Phi NWT compared to the Phi NWT, indicating the potential for higher speed applications. This being said, the CMOS device density is minimised by the 1/Phi NWT which has a narrower footprint. Drive current could be maximised by high S/D doping, as shown in recent studies, which have gone beyond

![Figure 5-20](image)

*Figure 5-20 Characteristics of the $I_D$ versus the $V_G$ for 5.0nm$\times$8.10 nm NWTs at 4 different tensile strain. with one. $L_g=12.0nm$, channel orientations $<110>$. Dashed, solid curves represent $I_D$-$V_G$ characteristics at drain voltage 0.070V and 0.050V respectively.*
doping levels of $10^{21}$ cm$^{-3}$ [214]; however, the development of 5 nm technology still requires further work in the area of contact resistance. Phi NWT $I_{\text{on}}$ is compared with S/D doping of a maximum $1.2 \times 10^{21}$ cm$^{-3}$, with contact resistance (500 - 2500 Ω), and (10 – 22 nm) gate length (Figure 5-21). As illustrated in the Figure 5-21 $I_{\text{on}}$ value is 1.4 mA/µm at the 5 nm node with $L_g=12$ nm and $R=2314$ Ω [4], [215], [6].

The ITRS roadmap incorporates vertically-stacked lateral NWT nanowires in order to maximise per-footprint current and minimise contacted gate pitch. Vertically-stacked Phi and $1/\text{Phi}$ NWTs are illustrated in Figure 5-23, with their $I_DV_G$ characteristics presented in Figure 5-22.

![Figure 5-21 Correlation between NWT performance ($I_{\text{on}}$) and S/D peak doping, S/D contact resistance, and $L_g$. NWT cross-section is Phi (5.0nm×8.1nm) and channel orientation is <110>. $I_{\text{on}}$ is evaluated at $V_D=0.70$V.](image-url)
Figure 5-22 Characteristics of the $I_D$ versus the $V_G$ of vertically stacked of two lateral Phi (5.0nm×8.1nm) (red) and (8.1nm×5.0nm) 1/Phi (black) NWTs at $V_D=0.70$V. The gate length is 12.0 nm. The crystallographic channel orientation is <110>.

Figure 5-23 (left) Vertically stacked lateral NWT of both phi and 1/Phi. The gate pitch layout for Phi and 1/Phi NWTs.
5.12 Summary

In this chapter, the use of a quantum-corrected Monte Carlo simulator when exploring nonequilibrium transport in NWTs’ suitable for the 5nm CMOS technology generation. The simulations were performed in order to assess whether the semiconductor industry targets can be met through the use of a single-channel NWT. It was found that this is not possible with the saturation current falling short of the target value of 1.58 mA/µm. The simulations also reveal that NWT saturation current, still does not meet the target even if high tensile strain is introduced in the channel of single nanowire NWT. Whilst the quantum-corrected MC simulations are effective in predicting the transistor performance in the case of small NWTs, they cannot take into account the S/D contact resistance. Additionally, the MC simulation approach is highly time- and memory-intensive when simulating stacked NWTs. Following calibration (based on density gradient confinement effective mass, the gate work function, and mobility models and their relevant parameters), the DD simulation approach was adopted for the simulation of vertically-stacked NWTs.

The DD simulations have revealed that the industrial target value of the drive current can be met by NWTs with two vertically-stacked lateral channels and highly-doped S/D regions. This being said, the current is greater in the top nanowire compared to the bottom nanowire, since the top nanowire is closer to the S/D contact. The reason for this is the voltage decrease due to the S/D series resistance, and this can result in faster degradation of the top nanowire with time. A more even current density can be achieved by the slicing of the S/D contacts. Finally, this chapter also discussed the effects of the golden ratio Phi on device performance and electrostatic integrity in n-type silicon nanowire transistors for 5 nm CMOS technology, demonstrating that NWT performance can be maximised through the use of NWT shapes with near-golden ratio (Phi) aspect ratios.
Chapter 6. Study of statistical variability

6.1 Introduction

NWT statistical variability, associated with the discreteness of charge and granularity of matter, which cannot be controlled through tightening process control and which is a key contributor to variability in CMOS devices, is discussed in this chapter. It is possible to model deterministically the impact of stress and lithography on the systematic CMOS variability, and to capture this in the design and verification process. However, it is only possible to model and represent the impact of the statistical variability on the transistor characteristics probabilistically. When measuring a pair of identical adjacently-placed transistors, the two transistors can have characteristics from the extreme tails of the statistical distribution.

Today, the traditional approach of simulating, designing and describing CMOS transistors, in terms of smooth interfaces and boundaries and continuous ionised dopant charge, is becoming outdated. The contemporary transistors are increasingly atomistic in nature. Figure 6-1 illustrates the random discrete dopants (RDD) in a NWT with S/D peak doping $N_{S/D}=1\times10^{20}\text{cm}^{-1}$ and $N_{S/D}=8\times10^{20}\text{cm}^{-1}$, and with channel doping $N_{ch}=1\times10^{17}\text{cm}^{-1}$. The top of the image illustrates the random dopant distributions. The middle illustrates the impact of RDD on the 3D electric potential distribution. The bottom of the image illustrates the impact of RDD on the NWT’s potential distribution, depicted by the sliced y-x and z-x plane at the centre of the two NWTs.
Significant variations in the transistor characteristics can be introduced by the atomicity of matter and electric charge granularity. There is both variation in numbers and positions of activated dopants in the NWT’s active region. This results in a high level of variation to the transistor characteristics as a result of the microscopic differences between each transistor. Furthermore, gate oxide thickness is equal to approximately three atomic layers with an average of 1-2 atomic layers of interface roughness, resulting in considerable variation in oxide thickness. Consequently, correlations can be seen between the body thickness and the oxide thickness of the transistors. Both gate pattern line edge roughness (LER) and corresponding geometrical statistical variations are the inevitable result of photoresist granularity and illumination nonuniformities [216]. Furthermore, statistical variability can also arise as a result of metal gate or poly-silicon granularity, as well as high-κ dielectric granularity.

*Figure 6-1 The RDD source/drain peak doping $N_{S/D}=1 \times 10^{20} \text{cm}^{-1}$ and $N_{S/D}=8 \times 10^{20} \text{cm}^{-1}$, the channel doping $N_{ch}=1 \times 10^{17} \text{cm}^{-1}$ (top). The effects of RDD on the electric potential distribution on the conduction channel and also in the S/D regimes (mid). Two slices cut along y-x plane and along z-x plane at centre of the NWT to show the effect of RDD on potential distribution inside the NWT (down).*
In this chapter, the impact of the key sources of variability on NWTs are explored using variability aware MC- and PS-calibrated DD simulations.

6.2 Variability sources

In-depth numerical GAA NWT simulations has been performed in order to examine the magnitude of the statistical variability resulting from Metal gate granularity (MGG), line edge roughness (LER), random discrete dopants (RDD) and other sources [217] [218], [219]. The results will be discussed in the present chapter. Earlier research has illustrated the use of GARAND in the analysis of statistical variability through the simulation of various statistical variability sources [220]–[223]. The key sources of statistical variability will be discussed individually in greater details in the following subsections.

6.3 Random discrete dopants (RDD)

Notable changes in device characteristics due to the random and discrete nature of dopant charges (RDD) in the channel region can be observed in highly-scaled NWTs. The impact of RDD has been discussed in earlier research of both multi-gate MOSFET and traditional MOSFET [224], both experimentally [225], [226] and in simulation analysis [99], [227], [228]. The traditional approach to numerical device simulation is now changing as a result of the effect of the RDD on the output, the functionality and the reliability the relevant systems [229], macroscopic simulations now unable to represent accurately the transistor behaviour using a continuous charge distribution [98]. In the statistical ‘atomistic’ device simulations random discrete dopants are primarily created using classical TCAD process simulation-based doping profiles. A more modern and accurate approach is the determination of the discrete dopant atom locations by means Kinetic Monte Carlo ‘atomistic’ process simulations. In the RDD simulations ensembles of devices with microscopic different dopant distributions representing exactly the same average (macroscopic) dopant distributions must be created. Each location of the silicon lattice in GARAND DD simulation domain is scanned individually based on the methodology outlined in [230], [231], with dopants added at random to the crystal sites based on a probability value derived from the relevant levels of dopant to silicon using a rejection
method. The frequently-adopted in MC simulations cloud-in-cell (CIS) method [232] is then adopted to distribute each dopant’s charge to the eight neighbouring mesh nodes. Figure 6-1 presents a standard 3D potential distribution denoting the effect of randomly-introduced discrete topics in an average 7 nm n-channel NWT.

The existing literature addresses the issues arising from the use of fine mesh for individual charge resolution in atomistic DD simulation [233]. Here, the electrostatic potential, derived from the output of the Poisson equation, determines the electron concentration when using Fermi-Dirac or Boltzmann statistics in the traditional DD model. Consequently, mobile charge localisation in the deep Coulomb potential wells, resolved by the fine mesh occurs. In quantum mechanics terms, due to the high ground electron state in the Coulomb well, this localisation (trapping) is non-physical. The current voltage characteristics and the threshold voltage are artificially modified as a result of greater resistance in the S/D regions and changes in the depletion layer as a result of the artificial charge trapping. Additionally, charge trapping can also result in a greater influence of mesh size on the accuracy and the convergence of the traditional DD simulations. Here, an increase in trapped charge arises in line with the mesh refinement, which may be used to improve the resolution of the single Coulomb potential well.

Two of the approaches addressing the above issues in atomistic simulation scenarios include the application of screening [234] criteria to divide Coulomb potential into short- and long-range elements [233], as well as charge smearing [234]. The latter method, which is entirely empirical in nature can arbitrarily reduce the RDD impact particularly, in smaller devices. In the former method, limitations include the risk of double counting of mobile charge screening, as well as the random selection of cut-off points. Furthermore, long-range potential well depth at the charge point is significantly greater than the Coulomb well ground state, with significant charge trapping still a possible outcome.
Confinement effects associated with the Coulomb potential of discrete dopants are accurately represented through the application of DG quantum corrections, which also takes into account the confinement effects within the channel. DG corrections in DD simulations can be calibrated in respect of the more accurate results obtained from a coupled Poisson/Schrödinger (PS) solution.

**Figure 6-2** Saturation transfer characteristics of 5nm × 8nm NWT (1000) device ensembles under influence of RDD at low drain voltage (0.05V).

**Figure 6-3** Saturation transfer characteristics of 5nm × 8nm NWT (1000) device ensembles under influence of RDD at high drain voltage (0.7V).
The microscopic effects of RDD within device 7 nm NWT are illustrated in Figure 6-1, with RDD-based potential fluctuation also presented. Here, S/D peak doping is $N_{\text{S/D}} = 1 \times 10^{20} \text{cm}^{-1}$ and $N_{\text{S/D}} = 8 \times 10^{20} \text{cm}^{-1}$, whilst channel doping is $N_{\text{ch}} = 1 \times 10^{17} \text{cm}^{-1}$ (shown at the top of the image). The figure also demonstrates the impact of RDD on electric potential distribution in the conduction channel and S/D regimes. The impact of RDD on the potential distribution within the NWT is illustrated in the y-x and z-x slices through the NWT centre. The S/D regimes demonstrate significant fluctuation in electric potential, with the S/D extensions showing a lower level of fluctuation. The doping concentrate has an influence on the different levels of potential fluctuation seen in these simulations. Because there are a small number of ionised acceptors in the gate region of the Si channel, this results in smooth potential within the channel. It is interesting to note that because of the low doping in the channel, RDD $V_T$ fluctuations are minimised in NWTs compared to conventional MOSFETS.

This section discusses the effect of RDD on the on current $I_{\text{on}}$ and the threshold voltage $V_T$ in NWT with two cross sections of 5 nm and equal gate lengths. The RDD-based fluctuations are simulated using a statistical ensemble of 1,000 microscopically different NWTs. Significant fluctuations in S/D access resistance arise as a result of the potential fluctuation stemming from S/D extension donors, leading to greater on-current variance. RDD introduces also $V_T$ fluctuations as a result the potential fluctuations in the channel. Figure 6-3 depicts the set of 1,000 gate transfer characteristics at high drain voltage, while Figure 6-2 depicting the low drain voltage characteristics. There is a relatively low I-V dispersion in the sub-threshold region due to the relatively low-doped channel, with a rapid increase in on-current dispersion occurring as performed result of the access resistance variation which has greater impact at low drain bias.

Figure 6-4 and Figure 6-5 respectively illustrate the correlation between the on-current and the threshold voltage at high and low drain bias. The standard deviation of the threshold voltage ($\sigma V_T$) at high and low drain biases is 1.99 mV and 1.48 mV respectively. These low standard deviations suggest that $V_T$ fluctuations are effectively minimised due to the low channel doping, with no ionised impurities beneath the gate in most cases. At the high drain voltage, the on-current standard deviation ($\sigma I_{\text{on}}$) is 0.0412 mA/μm, with the low drain
voltage on-current standard deviation is 0.0125 mA/μm. The reason for this are the S/D access resistance fluctuations resulting from variation in numbers and positions of the S/D extension donors, indicating that scaling causes a marked rise in on-current variability. Additionally, it should be pointed out that RDD also impacts DIBL fluctuations as a result of a change in the effective potential as illustrated in Figure 6-6.

Figure 6-4 Scatter plots of Ion versus $V_T$ at saturation for 1000 device ensembles with RDD. At high drain voltage (0.7 V).
Figure 6-5 Scatter plots of $I_{on}$ versus $V_T$ at saturation for 1000 device ensembles with RDD, at low drain voltage (0.05 V).

Figure 6-6 Histograms of the impact of RDD- generate fluctuations on DIBL of 5nmx 8nm NWT.
6.4 Line edge roughness (LER)

Figure 6-7 and Figure 6-8 depict the impact of the line edge roughness (LER) on the simulated NWTs, which is one of the key sources of fluctuation introduced during lithography and consequent etching [87]. The reduction of LER to less than 1 nm is challenging, with the photoresist used in the 14 nm lithography systems [235], [236] having restricted molecular dimensions. Given this, LER will become a key source of intrinsic parameter fluctuation with rising importance over the coming years [87].

One-dimensional (1D) Fourier synthesis is conducted to create random junction patterns. Here, gate edges are created from a power spectrum relating to a Gaussian autocorrelation function, using as parameters the correlation length ($\Lambda$) and RMS amplitude ($\Delta$). RMS amplitude is essentially the standard deviation of the gate edge’s $x$-coordinate (based on the assumption of a gate edge parallel to the $y$-direction). The LER value is equal to three times the value of RMS amplitude ($3\Delta$).

![Figure 6-7 Si NWT After the final anneal process LER = 1.12 ± 0.02 nm LER = 0.85 ± 0.01 nm [87].](image)

The Gaussian autocorrelation function provides the power spectrum needed to determine the amplitudes of a sophisticated set of $N$ elements, generated using a random line algorithm. As illustrated in (6-1), the power spectrum for the Gaussian autocorrelation function is represented by $S_G$: 
Chapter 6. Study of statistical variability

\[ S_G(k) = \sqrt{\pi} \sigma^2 \Lambda e^{-k^2\Lambda^2/4} \]  

(6-1)

where \( k = i(\frac{2\pi}{N}, dx) \) is the discrete spacing utilised for the line, with \( 0 \leq i \leq N/2 \).

Whilst no two lines are identical, due to the random selection of element phases, all but the \((N/2)-2\) elements (which are independent elements) are chosen based on symmetry considerations, resulting in a real height function \( (H(x)) \) following inverse Fourier transform. Figure 6-9 and Figure 6-12 respectively illustrate the effect of LER on gate transfer characteristics in 1,000 5nm \( \times \) 8nm NWTs (with this number limiting the risk of statistical error) at low and high drain bias.

![Figure 6-8 General 3-D representation of the LER and WER of single NWT](image)

Figure 6-8 General 3-D representation of the LER and WER of single NWT

Figure 6-10 represent the impact of LER on the DIBL distribution, while Figure 6-11 and Figure 6-13 represent the correlations between \( I_{on} \), and \( V_T \) at high and low drain bias respectively. It is clear that DIBL fluctuations in this case are similar to the fluctuations caused by RDD as a result of changes in channel length across the transistor width. The LER results in a standard deviation of 5.58 [mV] for \( \sigma V_T \) (compared to 1.99 under RDD), and 0.0317 [mA/\( \mu \)m] for \( \sigma I_{on} \) (compared to 0.0312 under RDD), 0.2 [mV/dec] for SS (compared to 0.26 under RDD), and 0.96 [mv/V] for DIBL (compared to 1.01 under RDD). The DIBL
and $V_{T}$-$I_{on}$ distributions become wider, indicating greater variability, as the dimensions of the scaled devices are reduced.

**Figure 6-9** Saturation transfer characteristics of 5nm x 8nm NWT (1000) device ensembles under influence of LER at low drain voltage (0.005V)

**Figure 6-10** Histograms of the impact of LER-induced fluctuations on DIBL of 5nm x 8nm NWT
Figure 6-11 Scatter plots of Ion versus $V_T$ at saturation for 1000 device ensembles with LER. At high drain voltage (0.7 V).

Figure 6-12 Saturation transfer characteristics of 5nm $\times$ 8nm NWT (1000) device ensembles under influence of LER at high drain voltage (0.7V)
Chapter 6. Study of statistical variability

6.5 Metal gate granularity (MGG)

Intrinsic fluctuations in the transistor characteristics originate also from metal gate granularity (MGG). Following thermal annealing, crystallisation of the metal gate occurs. Randomly-sized grains emerge during the gate’s polycrystallisation, with various crystallographic orientations and different corresponding work functions. At the time of writing, this is one of the key sources of fluctuation in the distribution of transistor parameters.

In the gate region the local threshold voltage experiences random variation based on the work functions of different crystallographic metal grains at the metal/oxide interface. Each metal grain differs from the next in terms domain orientations, probability for occurrence, the work-function value, and the average grain size (which is determined greatly by the annealing temperature and metal utilised). As discussed in the literature, the grains in the TiN metal gate commonly used in the contemporary CMOS technology can have $<200>$ or $<111>$ orientations [237], thus leading to a 0.2 eV difference between the corresponding

*Figure 6-13 Scatter plots of $I_{on}$ versus $V_T$ at saturation for 1000 device ensembles with RDD. At low drain voltage (0.005 V)*
work-function. Additionally, research indicates that there is a 60% probability for the crystalline orientation with 4.6 eV work-function and 40% probability for the orientation with of a 4.4 eV work-function [238].

Figure 6-14 illustrates the electric surface potential in a NWT for MGG with average size of 3 nm, 4nm, 5nm and 6nm. Here, a statistical ensemble of 1,000 NWTs with microscopically different MGG configurations were simulated and the impact on V_T, I_on and DIBL distributions were evaluated. Figure 6-15 illustrates the complete I_D-V_G characteristics of the simulated NWTs with different average metal grain size. Clearly with the increase of the MGG size the variability increases. It is interesting to note that in the case of LER and RDD

![Figure 6-14 The electric surface potential in the body of the device in the presence of MGG for the four size of MGG 3nm, 4nm, 5nm, and 6nm.](image-url)
the average $I_D-V_G$ characteristic is above the median $I_D-V_G$ characteristic, whilst in the case of MGG, the average $I_D-V_G$ characteristic is below the median.

The effect of MGG on the standard deviations of $V_T$, $I_{on}$, SS and DIBL is illustrated in Table 6-1. Additionally, Figure 6-16 presents $I_{on}/V_T$ scatter plots at high drain bias (0.7 V) for 1,000 device samples with MGG average sizes of 3nm, 4nm, 5nm and 6nm.

In comparison to LER and RDD, as sources of variability, work function variability appears to have a significant impact on the spread of the dispersion of the transistor characteristics. Here, MGG diameter is positively correlated with the degree of dispersion. It is worth highlighting that a near-parallel change in current-voltage characteristics was observed.

6.6 SV combinations

This section we examine the combined effect of RDD, LER and MGG on $V_T$, $I_{on}$ and DIBL with an exploration of the combined impacts of the three approaches. Since MGG represents the metal gate granularity arising out of the annealing process at high temperatures, it is important to incorporate MGG into SV simulation due to the impact this has on increasing statistical variability. MGG occurs once S/D high-temperature annealing has been performed on the metal gate during the gate-first process [239]. On the other hand, the metal gate is created following S/D annealing in the gate-last process, which can significantly minimise the MGG effects. Simulation without the incorporation of MGG relates to the gate-last process, though the gate-first and gate-last processes exist side by side. In this study, If RDD, MGG and LER are independent the approximate standard deviation ($\sigma$) of the relevant transistor figure of merit corresponding to the combined effect of the three parameters is given in equation 6-2:

$$\sigma_{\text{total}} = \sqrt{(\sigma_{\text{RDD}})^2 + (\sigma_{\text{LER}})^2 + (\sigma_{\text{MGG}})^2}$$ (6-2)
Figure 6-15 Saturation transfer characteristics of 5nm x 8nm NWT (1000) device ensembles under influence of MGG for the four size of MGG 3nm, 4nm, 5nm, and 6nm at high drain voltage (0.7V)
Table 6-1 Impact of MGG size on standard deviations of $V_T$, $I_{on}$, SS, and DIBL.

<table>
<thead>
<tr>
<th>MGG (nm)</th>
<th>$\sigma V_T$ [mV]</th>
<th>$\sigma I_{on}$ [mA/um]</th>
<th>$\sigma SS$ [mV/dec]</th>
<th>$\sigma DIBL$ [mV/V]</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>17.46</td>
<td>0.04465</td>
<td>0.58</td>
<td>3.25</td>
</tr>
<tr>
<td>4</td>
<td>23.31</td>
<td>0.06050</td>
<td>0.77</td>
<td>3.99</td>
</tr>
<tr>
<td>5</td>
<td>27.80</td>
<td>0.073353</td>
<td>0.92</td>
<td>4.53</td>
</tr>
<tr>
<td>6</td>
<td>32.48</td>
<td>0.085672</td>
<td>0.97</td>
<td>5.00</td>
</tr>
</tbody>
</table>

Figure 6-16 Scatter plots of $I_{on}$ versus $V_T$ at saturation for 1000 device ensembles under influence of MGG for the four size of MGG 3nm, 4nm, 5nm, and 6nm at high drain voltage (0.7V)
Both the 5 nm × 8 nm (Phi) and 8 nm × 5 nm (1/Phi) NWTs simulated here have a gate length (Lg) of 12 nm, with the same cross-sectional area and channel orientation. A statistical sample of 1,000 microscopically different transistors was simulated. Figure 6-17 and Figure 6-18 illustrate histograms of the I_{on} and V_T distributions arising based on the individual and combined RDD, LER and MGG effects. LER is found to have a more significant impact on SV than RDD, while MGG having the strongest impact on I_{on} compared to RDD and LER. Furthermore, the results reveal that the standard deviation of I_{on} values for the 1/Phi NWTs falls below that of the I_{on} SD of the Phi NWTs at a channel orientation of <110>.

It can therefore be proposed that the ideal AR is provided by the Phi and 1/Phi ratios, which represents one of the most important findings of the simulation experiments. It is clear that the specific parameters of the sources of variability dictate the variability of V_T and I_{on}, therefore highlighting the need for further research into the impact of singular SVs on V_T and I_{on} for Phi and 1/Phi NWTs.

*Figure 6-17* I_{on} distributions subject to individual SV and combined SV (RDD, LER, and MGG) which are defined of "gate-first technology" of Phi NWT and 1/Phi NWT.
In this study, as noted, sets of 1,000 transistors with microscopic differences were simulated in order to test each variability source. The impacts of SV sources on $I_{on}$ of Phi NWTs echo those of 1/Phi NWTs, with a 0.1mA/µm change in Gaussian distribution noted. Since the mobile charge (QM) in the 1/Phi NWTs’ channel exceeds that of the Phi NWTs, this change is important. Additionally, the variability of MGG is significantly broader in the case of $I_{on}$ compared to other SV sources. The validity of these findings also holds true for variability in threshold voltage (see, Figure 6-17 and Table 6-2).

The $V_T$ distributions arising out of the impacts of RDD and LER (individual and combined) are depicted in Figure 6-18 and Table 6-3. The standard deviation values of $V_T$ are found to be positively related to $I_{on}$ in both the Phi and 1/Phi devices. Furthermore, the $V_T$ statistical variability of Phi NWTs appears to be better than that of the 1/Phi NWTs. The greater fluctuation in statistical parameters in the scaled devices has important implications for device performance with regards to transistor scaling.

<table>
<thead>
<tr>
<th>$I_{on}$ mA/µm</th>
<th>5nm x 8.1nm</th>
<th>8.1nm x 5nm</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SD</td>
<td>Mean</td>
</tr>
<tr>
<td><strong>WER</strong></td>
<td>0.0380</td>
<td>1.2460</td>
</tr>
<tr>
<td><strong>RDD</strong></td>
<td>0.0379</td>
<td>1.2030</td>
</tr>
<tr>
<td><strong>MGG</strong></td>
<td>0.0882</td>
<td>1.2230</td>
</tr>
<tr>
<td><strong>SV</strong></td>
<td>0.1031</td>
<td>1.1710</td>
</tr>
</tbody>
</table>

*Table 6-2: Mean and standard deviation values of $I_{on}$, subject to individual SV and combined SV (RDD, LER, and MGG) which are defined of “gate-first technology” of Phi and 1/Phi NWTs.*
The distribution and correlations between the figures of merit (FoM) of the Phi and 1/Phi NWT obtained from the TCAD simulations are illustrated in Figure 6-19. Here, RDD, LER and MGG as the key sources of variability (combined effects) are combined in the simulations. Based on the results presented in Figure 6-18 and Figure 6-19, $V_T$, $I_{off}$, and $I_{on}$ appear to be correlated; although DIBL and the other FoM are weakly correlated. One of the main findings from the simulations is the superior $I_{on}$ and lower variability in the 1/Phi NWTs, indicating a potential for greater speed and lower power design. This being said, with a wider footprint than the Phi NWTs, this will negatively impact the density of the CMOS device, thus representing a key limitation of the 1/Phi NWT.

Figure 6-18 Threshold voltage ($V_T$) distributions subject to individual SV and combined SV (RDD, LER, and MGG) which are defined of “gate-first technology” of Phi NWT and 1/Phi NWT.
Table 6-3 Mean and standard deviations values of $V_T$ subject to individual SV and combined SV (RDD, WER, and MGG) which are defined of “gate-first technology” of Phi NWT and 1/Phi NWT.

<table>
<thead>
<tr>
<th>$V_T$ mV</th>
<th>5nm x 8.1nm</th>
<th>8.1nm x 5nm</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SD</td>
<td>Mean</td>
</tr>
<tr>
<td>WER</td>
<td>5.807</td>
<td>107.85</td>
</tr>
<tr>
<td>RDD</td>
<td>2.001</td>
<td>110.66</td>
</tr>
<tr>
<td>MGG</td>
<td>27.82</td>
<td>110.41</td>
</tr>
<tr>
<td>SV</td>
<td>28.52</td>
<td>111.9</td>
</tr>
</tbody>
</table>

Figure 6-19 DIBL, $V_T$, $I_{OFF}$, and $I_{ON}$, distributions subject to combined SV (RDD, LER, and MGG) which are defined of “gate-first technology” of Phi NWT and 1/Phi NWT.
6.7 Cross-sectional aspect ratio improves SV

Due to quantum mechanical effects the shape and the cross-sectional aspect ratio of the NWT has significant impact on the electrostatic-driven performance in 5 nm NWTs. The \( V_T \), \( I_{on} \), DIBL and \( I_{off} \) distributions in Figure 6-20 have been subjected to combined to the combined effect of RDD, LER and MGG for the gate-first technology. The distributions are near-Gaussian, based on the results of the simulations of a statistical sample of 1,000 microscopically different transistors the nine NWTs from Table (4-6). The Phi and 1/Phi ratio NWTs have more favourable \( I_{on} \), DIBL, \( V_T \) and \( I_{off} \) distributions. This could be associated with the fact that the Phi and 1/Phi ratio NWTs having greater gate capacitance, larger mobile charge and better electrostatic integrity.

![Figure 6-20 DIBL, \( V_T \), \( I_{off} \), and \( I_{on} \), distributions subject to combined SV (RDD, LER, and MGG) with a trap sheet density of \( 1 \times 10^{12} \) cm\(^{-2}\) of all NWTs listed in table (4-6).]
6.8 Statistical variability of stacked NWT

In order to maximise the current-per-footprint whilst minimising the contacted gate patch, the ITRS roadmap incorporates vertically-stacked lateral NWT nanowires. Simulations indicate that there is no significant relationship between the number of stacked NWTs and its electrostatic integrity including DIBL and SS. For instance, a minor 1% shift in SS is noted when comparing single-, double- and triple-channel lateral NWTs [164]. Stacked NWTs are widely recognised as the superior alternative to Fin-FETs based on their high drive current ultimate reduction of gate length and high immunity to short channel effect. This being said, the SV remains of crucial significance to CMOS technology, although no earlier studies address the impact of stacked NWTs on variability. The SV of vertically-stacked lateral NWTs are analysed in this section. Figure 6-22 illustrates the current-voltage transfer characteristics of a statistical ensemble 1,000 two-channel atomistically different devices under the combined impact of RDD, LER and MGG. Figure 6-21 illustrates the same statistical characteristics, but for NWT with three-lateral channels instead of two. It is clear the triple-channel NWTs has lower variability, with the difference being strongly pronounced in the sub-threshold region. The threshold voltage ($V_T$) distributions of 1,000 single-, double- and triple-stacked NWTs under the combined impact of RDD, LER and MGG as SV are illustrated in Figure 6-23. Triple-stacked lateral NWTs were found to have a lower $V_T$ standard deviation ($\sigma$) compared to the double-stacked NWTs. Additionally, single NWTs demonstrated the greatest $\sigma V_T$ value of all three types. It is worth highlighting that the $\sigma V_T$ variation declines with the increase in the number of stacked lateral channels. As illustrated in the histogram presented in Figure 6-25, a 27.39% or 1.52 mV/V decrease in $\sigma$DIBL is seen with the introduction of two lateral channels. Additionally, a 13.73% decrease in $\sigma$SS is observed in the double-channel NWT, with a 16.67% decrease in the triple-channel NWT as shown in Figure 6-24. As illustrated in Figure 6-26, the results also demonstrate that the $\sigma I_{on}$ of the double-stacked NWTs exceeds that of the single NWTs by 25% (0.02). Furthermore, the $\sigma I_{on}$ of triple-stacked NWTs is 1% compared to that of the double-stacked NWTs. Additionally, as shown in Table 6-4, the introduction of stacked lateral channel NWTs reduces the variability or standard deviations of SS, DIBL and $V_T$. 
Figure 6-21 Linear transfer characteristics for the ensemble (1000) with main sources of SV (RDD, LER, and MGG) for Si NWT with three channels. \( L_g=12.0 \text{ nm}, V_D=0.70\text{V} \)

Figure 6-22 Linear transfer characteristics for the ensemble (1000) with main sources of SV (RDD, LER, and MGG) for Si NWT with two channels. \( L_g=12.0 \text{ nm}, V_D=0.70\text{V} \).
Figure 6-23 $V_T$ distributions subject to combined SV (RDD, LER, and MGG) of single, double and triple NWTs.
Chapter 6. Study of statistical variability

Figure 6-24 Comparison of the SS distributions subject to combined SV (RDD, LER, and MGG) of single, double and triple NWTs.

Figure 6-25 Comparison of DIBL distributions of single, double, and triple NWT subject to combined SV (RDD, LER, and MGG)
Table 6-4 Mean and standard deviations values of $V_T$, SS, and $I_{ON}$ subject to combined SV (RDD, LER, and MGG) of single, double, and triple NWTs

<table>
<thead>
<tr>
<th>NWT</th>
<th>$V_T$ [mV]</th>
<th>SS [mV/dec]</th>
<th>$I_{ON}$ [mA/µm]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>$\sigma_{V_T}$</td>
<td>Mean</td>
</tr>
<tr>
<td>Single</td>
<td>112.8</td>
<td>28.9</td>
<td>62.33</td>
</tr>
<tr>
<td>Double</td>
<td>89.86</td>
<td>22.08</td>
<td>62.45</td>
</tr>
<tr>
<td>Triple</td>
<td>74.6</td>
<td>18.93</td>
<td>62.55</td>
</tr>
</tbody>
</table>

Figure 6-26 Ion distributions subject to combined SV (RDD, LER, and MGG) of single, double, and triple NWTs
6.9 **Additional variability sources**

Process variability may be increased due to variation in the cross-sectional shape of the NWT, based on the fact that electrostatic driven performance and NWT geometry are correlated. Additionally, VSL NWT performance is determined by the position of lateral NWTs within the vertical stack. This is due to greater process variability as a result of series resistance between the S/D contacts and lateral channels.

6.10 **Summary**

The statistical variability of scaled NWT devices with characteristic cross section of 5 nm and with 12 nm gate length has been explored through extensive simulation and the corresponding results presented and analysed this chapter. The individual and combined effects of RDD, LER and MGG on $V_T$, SS, $I_{on}$ and DIBL have been simulated and discussed, with the findings suggesting that the key source of variability for all main electrical parameters considered in this research is MGG. For instance, $V_T$ standard deviation was 1.9 mV under RDD, 5.8 mV under LER, and 27.8 mV under MGG; with $I_{on}$ standard deviation being 0.0316 mA/um under RDD, 0.0312 mA/um under LER, and 0.0733 mA/um under MGG.

The results also indicate that NWT variability is impacted by cross-section shape and dimensions, with a significant reduction in SV found in NWTs with optimal aspect ratios. Furthermore, a reduction in the variability of threshold voltage drive current are observed in vertically-stacked NWTs. As noted, the reduction in $\sigma V_T$ found in vertically-stacked NWTs is due to the cancelling-out of variability in double lateral and triple-channel NWTs.
Chapter 7. Conclusions and Future work

7.1 Conclusions

This PhD study was carried out with the primary objective of simulating and optimising a realistically-scaled Si NWTs suitable for and beyond the 5nm CMOS technology generation. We have adopted a simulation approach that could effectively reflect quantum confinement effects with non-equilibrium transport. The aim of this thesis is to carry out comprehensive TCAD simulation research in order to study the physics of silicon nanowire transistor devices at the ultimate scaling limits. With the advanced modelling tools employed in this study, the performance limits of Si NWTs near the end of the CMOS technology scaling have been evaluated and important challenges in NWT device design have been addressed.

Chapter two describes the challenges that need to be addressed in scaling of conventional MOSFETs and presents the technology innovations that have been implemented and proposed for the future. The theory of the MOSFET scaling is also outlined.

The relentless requirement for high-performance and low power devices with ever increasing density of integration drives the CMOS technology to the ultimate nanoscale dimensions. Nanowire MOSFETs transistors (NWTs) are among the candidates to extend CMOS downscaling to ultimate limits, eventually replacing the triple gate FinFET architectures after the 7nm CMOS technology generation. The nanowire gate-all-around configuration has the best immunity against short channel effects (SCE) allowing ultimate transistor scaling. Since statistical variability is becoming a serious scaling limitation factor, the sources of statistical variability have been discussed in detail.

Semiconductor device design and optimisation depend highly on the accurate simulation of the device’s physical behaviour and electrical characteristics. By modelling semiconductor device behaviour as part of the CMOS Technology Computer Aided Design (TCAD)
Chapter 7. Conclusions and Future work

process, we can assess its key parameters and operation and understand how to improve its functionality and performance.

Chapter three has outlined the basis of the 3D drift/diffusion simulation, presenting both the drift/diffusion (DD) system of semiconductor equations and the density gradient approach to introduce quantum corrections in the DD system as well as the algorithms employed solve them self consistently.

In this study we also use the 3D ensemble MC simulation approach with accurate quantum corrections for the predictive simulation of nanowire transistor (NWT) charge transport and performance. For the simulation of large multichannel NWTs and for the simulation of statistical variability we use drift diffusion simulations thoroughly calibrated to the results of the Monte Carlo simulations. It was clarified that non-equilibrium transport effects in nano-scale devices cannot be represented by the DD model. Furthermore, transport variability due to ionised impurity scattering is also difficult to capture using DD-simulation. Monte Carlo simulation are used to calibrate the drift-diffusion simulations of large multichannel NWTs and to perform statistical variability simulations. Self-consistent Poisson- Schrödinger solver is adopted for studying the impact of the quantum confinement on the mobile charge and capacitance of NWTs with different cross section.

Chapter four examines the ultimate scaling limits of NWTs with different cross-sectional shapes, using simulations employing Poisson Schrodinger (PS) quantum corrections implemented in the ‘atomistic’ drift-diffusion (DD) simulator GARAND [GSS]. Realistic design parameters of the source-drain extensions and their impact on SS and DIBL are considered. In addition, the effect of the nanowire cross-sectional shape on the gate capacitance, the charge available for transport and the speed are investigated and discussed.

Signatures of isotropic charge distributions within Si NWTs were observed exhibiting the same attributes as the golden ratio (Phi), the significance of which is well known in the fields of art and architecture. The research demonstrates that the quantity of mobile charge within the channel, along with the intrinsic speed of the device are determined by device geometry and may be related to the golden AR (Phi) of the nanowire transistors. This investigation
has demonstrated that NWT with aspect ratios equal or close to the golden ratio (Phi) can enhance gate capacitance and mobile charge in the channel and therefore could optimise the intrinsic speed of the device. This study has also investigated the influences of the gate length on the time delay and the main FoM, such as $V_T$, $I_{OFF}$ and $I_{ON}$ and DIBL.

In chapter five, quantum-corrected Monte Carlo simulations are used when exploring nonequilibrium transport in NWTs’ suitable for the 5nm CMOS technology generation. The simulations were performed in order to assess whether the semiconductor industry targets can be met through the use of a single-channel NWT. It was found that this is not possible with the saturation current falling short of the target value of 15% increase in the saturation current. The simulations also reveal that NWT saturation current, still does not meet the target even if high tensile strain is introduced in the channel of single nanowire NWT. Whilst the quantum-corrected MC simulations are effective in predicting the transistor performance in the case of small single channel NWTs, they cannot take into account the S/D contact resistance. Additionally, the MC simulation approach is highly time- and memory-intensive when simulating stacked NWTs. Following calibration (based on the gate work function, density gradient confinement effective mass, and mobility models and their relevant parameters), the DD simulation approach was adopted for the simulation of vertically-stacked NWTs.

The calibrated DD simulations have revealed that the industrial target value of the drive current can be met by NWTs with two vertically-stacked lateral channels and highly-doped S/D regions. This being said, the current is greater in the top nanowire compared to the bottom nanowire, since the top nanowire is closer to the S/D contact. The reason for this is the voltage decrease due to the S/D access resistance, and this can result in faster degradation of the top nanowire with time. A more uniform current density can be achieved by the side placement of the S/D contacts. Additionally, NWT performance was found to be optimised when using NWT shapes with aspect ratios similar to the golden ratio (Phi), based on the exploration of the impacts of Phi on 5 nm CMOS n-type silicon nanowire transistor performance and electrostatic integrity.
The impact of variability sources in the DD-based simulator is then presented in Chapter six. The individual and combined effects of RDD, LER and MGG on $V_T$, SS, $I_{on}$ and DIBL have been simulated and discussed, with the findings suggesting that the key source of variability for all main electrical parameters considered in this research is MGG. For instance, the $V_T$ standard deviation was 1.9 mV under RDD, 5.8 mV under LER, and 27.8 mV under MGG; with $I_{on}$ standard deviation being 0.0316 mA/um under RDD, 0.0312 mA/um under LER, and 0.0733 mA/um under MGG.

The results also indicate that NWT variability is impacted by cross-section shape and dimensions, with a significant reduction in SV found in NWTs with optimal aspect ratios. Furthermore, a reduction in the variability of threshold voltage and drive current are observed in vertically-stacked NWTs. As noted, the reduction in $\sigma V_T$ found in vertically-stacked NWTs is due to the cancelling-out of variability in double lateral and triple-channel NWTs. Additionally, device performance was also found to be significantly driven by the vertical position of lateral NWTs in the vertical stack as a result of the heightened process variability caused by series resistance between the S/D contacts and lateral channels.

Overall, the results of this study indicate that NWTs may be a valuable option as next generation transistor for industry application.

### 7.2 Contributions

The main contributions of this study is the finding that multichannel NWT are suitable candidates for the 5nm technology node meeting the performance targets and the pitch requirements. Some of the other important contributions are summarized as follows:

1- Developed novel methodology for DG calibration against more physical PS solution. The study has demonstrated that after careful calibration against the PS results the DG approach can be used for practical 3-D simulations of NWTs [7], [9].
2- For the first time we have studied in details the impact of the quantum mechanical effect on the electrostatically driven performance of NWTs suitable for a 5 nm CMOS technology and beyond. We have demonstrated that the NWT shape has a strong impact on the gate capacitance and the mobile charge in the NWTs at high gate bias. The quantum mechanical confinement results in a reduction in the mobile charge, gate capacitance, and intrinsic speed [9], [162], [185], [240].

3- An interesting observation was made in this research regarding the fact that the signatures of isotropic charge distributions in silicon nanowire transistors (NWT) displayed identical characteristics to the golden ratio (Phi). In turn, a simulation was conducted regarding ultra-scaled n-type Si (NWT) suitable for 5-nmCMOS technology and beyond. The results reveal that the amount of mobile charge in the channel and the intrinsic speed of the device are determined by the device geometry and could also be correlated to the golden ratio (Phi) [185], [186].

4- Ensemble Monte Carlo (MC) simulations with quantum correction are used to predict accurately the drive current of Si (NWT). The complex the carrier transport in the vertically stacked lateral NWTs and their performance are evaluated using novel methodology involving the thorough calibration of the current in a single channel to accurate reference MC simulations.[162], [164], [188], [189]

5- We have investigated the correlation between channel strain and device performance in various n-type Si- NWTs suitable for 5-nmCMOS technology and beyond. We have establish a correlation between strain, gate length and a cross-section dimension of the transistors [189].

6- Our simulations have shown that a single nanowire transistor even with strained Si channel will not be able to provide the required drive current in comparison to the industrial target at 5nm CMOS technology generation [164], [188].
7- Based on predictive simulations have proposes a “vertically stacked lateral NWTs” structure for the 5-nm CMOS technology that meets the required industry scaling projection [164], [185].

8- The current density degrades from the top to the bottom of identically sized lateral NWTs in the vertical stack. The upper lateral nanowires, closer to the source/drain contacts carry more current than the ones that are farther away (close to the bottom) due to the series resistance related voltage drop and may become weak point suffering extensive ageing. Solutions including the idea of increasing the diameter of lower nanowires concerning the upper nanowires in the stack. However, although the drain current is increased with the increase of the NWT cross-sectional area, the voltage drop will increase across the highly doped S/D regions. Furthermore, the possibility of decreasing the doping concentrations near the top lateral NWT and increasing it gradually near the middle and the bottom lateral NWTs would decrease the total drain current (Ion). Therefore, we have developed strategy to optimise S/D contacting by the deformation of S/D contacts [16] [17].

9- For the first time we have carried a comprehensive computational study of the impact of the principal sources of statistical variability including RDD, LER, and MGG, on the threshold voltage, drain-induced barrier lowering, and drive current of vertically stacked lateral NWTs suitable for the 5-nm CMOS technology [164], [185].

10- We have investigated for the first time the position dependent performance and geometrical variation of the lateral nanowires in the stack as new sources of process variability (IWCN17).

11- We have found that $V_T$ standard deviation of triple lateral stacked NWTs has a lower standard deviation ($\sigma$) than the double stacked NWTs while the value of $\sigma V_T$ for a single NWT shows the highest $\sigma$ compared with double and triple NWTs [164].
7.3 **Recommendations**

It will be useful if future research focuses on the optimisation of power and/or performance of ultimately scaled NWT beyond the 5nm CMOS technology optimising carrier mobility, electrostatics and parasitic capacitance in order to determine the value of NWT devices.

Mobility tends to suffer as a result of quantum confinement effects in conventional 5 nm NWTs, caused by phonon and surface roughness scattering. Alternative materials may need to be adopted to improve carrier transport as the scaling continues, with SiGe (offering better hole transport) and Ge (offering better electron and hole mobility) recommended for this purpose. Furthermore, decreasing the effects of fringing field capacitance could enhance NWT performance. At high frequencies, NWTs with a diamond-shaped S/D structure might have good performance.

Greater transistor performance can be achieved through the employment of 3D integration and vertical channel orientation, resulting in more favourable circuit delay and greater performance due to better per-footprint effective width. This approach can be applied to NWTs, with nanosheet structures providing an interesting alternative for optimising the effective width per layout footprint. This should maintain power density whilst enhancing drive current. The effective width per layout footprint is smaller in nanosheets than in regular NWTs. Using a thin and wide nanosheet, the effective width can be increased significantly in comparison to conventional NWTs. A greater effective width along with improved overall performance can be achieved through the use of a wider nanosheet due to the larger footprint, while short channel effects remaining unchanged.

There are still many technological and simulation issues associated with the advancement of NWTs and nanosheet transistor technologies, including the optimisation of shape with a decrease in roughness, inner spacer, access optimisation and strain management. However, recent testing and research have demonstrated the value of this alternative configuration to the future scaling and development of the industry. Therefore, we recommend further study of the positive impact of device width on performance, highlighting the potential benefits of nanosheet transistors in the optimisation of future CMOS applications.
The scaling NWTs beyond 3nm CMOS technology is high challenge, expecting that in the end the horizontal configuration (such as lateral nanosheet) will punch physics limits. The vertical configuration of NWTs transforms the layout configuration from a 2D to a 3D, where the gate length can be extended without occupation a wide area on the wafer. This technology requires intensive research of design co-optimization.
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