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MOSCAP Oxide Charges

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MOSCAP Oxide Charges

by

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An undergraduate honors thesis submitted in partial fulfillment of the requirements for the degree of Bachelor of Science in University Honors and Physics

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Abstract

Transistors form the backbone of digital devices, allowing for amplification and digital “switching” of electrical signals. One of the simplest devices is the Metal-Oxide-Semiconductor Capacitor (MOSCAP). The MOSCAP is analogous to a parallel plate capacitor with the Si wafer and metal layer corresponding to the positive and negative plates, and the oxide layer corresponding to the dielectric material. During the fabrication process it is likely that trapped charge impurities will become incorporated into the oxide layer of the MOSCAP. This thesis project will examine the effects of these charges on the capacitance and voltage characteristics of a MOSCAP.

Introduction

The world drastically changed in 1947 with the development of the first transistor by Shockley, Bardeen, and Brattain at Bell Labs. Large vacuum tube chambers could now be replaced with smaller solid-state devices. As scientists and engineers worked, these devices shrunk not only in size but in price, largely due to improvements in industrial fabrication techniques. These improvements led the way to the large scale production of transistors, causing an exponential increase in computing power, commonly known today as “Moore’s Law”. In effect, transistors have become ubiquitous in today’s society, finding their way into cars, appliances, and medical devices.

By far the most common type of transistor made today is the MOSFET (Metal Oxide Semiconductor Field Effect Transistor) which are the basis for the more advanced logic devices (CMOS) that make up ICs (Integrated Circuits). Going even further, MOSFETs can be built from MOSCAPs (Metal Oxide Semiconductor Capacitor). The key step in mass producing these devices is the fabrication process, where great care must be taken to eliminate any contaminants. This thesis will provide a look into the fabrication of MOSCAPs, and how varying material
properties will affect their performance. More specifically, this paper will investigate the effect of oxide fixed charges on device capacitance.

In order to study MOSCAPs it is essential to have sufficient background in solid state and device physics. This paper will strive to provide a sufficient introduction into the properties of metals, oxides and semiconductors. It will also explore the characteristics of MOSCAP devices: junction types, fabrication, charges, and models. Afterwards, the experimental methods used to verify the theoretical model of MOSCAP performance will be discussed. The data collected from this will then be analyzed and compared to the theoretical predictions. From this, there will be sufficient evidence to thoroughly examine the performance of MOSCAPs.

1.1 Introduction to Solid State Physics

The first foundation of MOSCAP device physics is the atomic organization of each material. The current physical model describes electrons around an atom as occupying orbitals, mathematical expressions which dictate the most probable regions that an electron will occupy. These orbitals can be described using four quantum numbers: ‘n’, the principal quantum number which relates to quantum energy, ‘l’, the azimuthal quantum number which relates to magnitude of angular momentum, ‘m’, the magnetic quantum number which relates to a component of the angular momentum, and ‘s’ the spin number which relates to either the spin up or spin down position. Each ‘n’ value can be thought of as a series of shells around the atom. For each principal quantum number there is a subset of azimuthal, magnetic, and spin numbers, which describe an electrical orbital for a given quantum energy level. While these shells are mathematically defined over all space, it is more useful to only consider the cases where the probability of an electron in each orbital is larger than zero. Each shell is categorized into different types: if ‘l’ is zero, then it is a ‘s’ orbital which is spherical, if ‘l’ is one, then it is a ‘p’ orbital which is a “dumbbell” shape, if ‘l’ is two then the orbital is a four lobed shape. There are more orbital shapes as ‘l’ increases, but these are less important for practical applications.¹
Only two electrons can occupy a particular set of principal, azimuthal, and magnetic quantum numbers. These correspond to the two spin numbers for spin up and spin down. This is called the “Pauli Exclusion Principle” and is very important as it limits the number of electrons in each shell. Following the Exclusion Principle is Hund’s Rule, which states all the possible energy states of a particular shell must be filled before doubling up the electrons via the Exclusion principle. Using the Pauli Exclusion Principle and Hund’s Rule, Pauli devised the idea of “aufbau”, in which orbitals are built up piece by piece like a house.

These principles can be used to explain the process of chemical bonding. Two atoms of the same energy level will each have a corresponding orbital. As they move closer together, their orbitals will overlap. At this point, the two orbitals will combine to make two new orbitals, one at a higher (bonding) energy and one at a lower (anti-bonding) energy state. The difference in energy states between the new orbitals is referred to as the bond energy. Depending on how many electrons are in the orbitals, either the lowest energy orbital will be occupied, or both the highest and the lowest orbital will be occupied. If only the low energy state is occupied, then the system has formed a chemical band because the combination provides a more preferable (lower) energy state. If instead the higher energy state is occupied, there will not be a stable bond.

Crystal structures can be thought of as large collections of atoms, and as such the formation of crystals can be thought of as analogous to the formation of a bond. Like chemical bonds, the orbitals of the electrons are combined. However, unlike chemical bonds, the combination gives a range of energies (energy band) and not just two bonding and anti-bonding states like with two atoms. Electrons in this band of orbitals are considered delocalized or corresponding to the whole crystal system and not just a particular atom. Generally only two bands are typically needed to explain the properties of solid state devices. The lowest band, called the valence band, represents bonding orbitals, while the highest band, called the conduction band, corresponds to delocalized anti-bonding orbitals.

Since the conduction band is more delocalized than the valence band, electrons in the anti-bonding orbitals are freer to move about the crystal structure.
These free moving electrons are called mobile carriers. To be promoted from the valence band to the conduction band, an electron must be excited in some way to overcome the energy gap between the two bands. In insulating materials, this energy gap is very large (several eVs) so it is rare for electrons to jump into the conduction band. With metals, the energy gap is very small or non-existent, so electrons frequently move to conduction band. As a result, insulators have poor electrical characteristics while metals have excellent electrical characteristics.

1.2 Semiconductors

As the name implies, semiconductors have properties between that of an insulator and a metal. It has a small energy band gap of a few eVs which allows it to have some electrons in the valence band and conduction band. In this way, semiconductors have a non-trivial amount of mobile carriers. Semiconductors also have two types of band gaps: direct and indirect. A direct band gap semiconductor such as gallium arsenide will allow an electron to move from valence to conduction bands without losing momentum. In a direct band gap, the maxima of the valence band are directly underneath the minima of the conduction band. When an electron moves from the valence band and combines with a hole in the conduction band, the resulting annihilation gives off energy as a photon. As a result, direct band gap materials have optical properties well suited for laser diodes and other optical devices.

In an indirect band gap such as in silicon, the maxima of the valence band is not directly underneath the minima of the conduction band. This implies that an electron traveling from valence to conduction bands will have a change in momentum. It follows that in semiconductor, the valence band is filled, while the conduction band is empty at lower temperatures. When an electron is excited into the conduction band, it leaves behind a hole, which is similar to positively charged electron. These holes can move around the crystal since they are also delocalized.
This results in a quasi-particle which can move freely and carry a charge, making it another mobile carrier.

An important aspect of solid state devices is the nature of these holes and electrons within the semiconductor. It is important to view mobile electrons as not a single electron but rather an increase in electron density over a small area. Likewise, a hole is viewed not as a single absent electron but a decrease in the electron density over a small area. These changes in density move around the crystal lattice and actually have a different mass than that of a single electron.  

To establish the electrical characteristics of a semiconductor it is helpful to use Fermi-Dirac statistics. The Fermi-Dirac distribution is used to determine the probability that an electron state is occupied at a certain energy level. The Fermi energy is thus defined as the energy at which the state has a probability of 0.5 and can also be thought of as the free energy of mobile carriers. Since a semiconductor typically has a full valence band and an empty conductor band, the Fermi energy level will be half the distance of the band gap, between the valence and conduction bands. This is a good approximation for semiconductors, but in reality, due to the complex structures, the Fermi level is a little below the middle of the band gap in silicon. Since the temperatures are not high, the Fermi-Dirac distribution can be modeled using the Maxwell-Boltzmann distributions. This distribution can lead to an expression for the concentration of electrons and holes in a semiconductor, the intrinsic mobile carrier concentration, \( n_i \):

\[
\begin{align*}
n_i &= N_c N_v e^{-\frac{E_a}{kT}}
\end{align*}
\]

where \( N_c \) is the effective electron concentration, and \( N_v \) is the effective hole concentration, \( E_a \) is the activation energy needed to transition from reactant to product, \( k \) is Boltzmann’s constant, and \( T \) is the absolute temperature. In a semiconductor at normal temperatures, a few electrons will transition to the conduction band, leaving behind holes in their place. This concentration is known as the mobile carrier concentration.

### 1.3 Doping
The next critical aspect of semiconductor physics is doping. Doping occurs with the insertion of shallow level impurities into a pure semiconductor. Supposed an element such as phosphorous is added to a silicon crystal. Since phosphorus has five free electrons, the first four will bond to the lattice with silicon, leaving one extra electron unbonded. This extra electron occupies a donor state just below the conduction band, but is excited up into the conduction band by the vibrational energy of the lattice. The electron then becomes delocalized, free to move about the lattice. At this point, the phosphorus atom is considered a positively ionized impurity. The result of this, is that by adding phosphorus to the silicon crystal, the amount of conduction electrons has increased, thereby improving the overall conductivity of the material.

The other method of doping introduces an element with three electrons instead of five, such as boron. As a result, boron leaves an extra hole in just above the valence band, called an acceptor state. The hole gets moved to the valence band due to the vibrational energy of the crystal, and as a result creates a delocalized negative mobile carrier. This makes boron a negatively ionized impurity. Similarly to the phosphorus, by adding boron, the overall conductivity of the semiconductor has increased.

Typically, Group VB elements such as phosphorus are considered donor impurities, while Group IIIB elements act as acceptor impurities. These impurities are called dopants. If the doping is done with a donor impurity, it is called n-type doping because more electrons are being sent to the conduction band, making the majority of the mobile carriers negatively charged. Conversely, if the doping is done with donor impurities, it is called p-type doping because more holes are being sent to the valence band, making the majority of the mobile carriers positively charged.

When doing p and n-type doping it is important to clarify the different types of mobile carriers. First, there are intrinsic mobile carriers, the electrons and holes that are free to move in a pure semiconductor crystal. Second, there are majority carriers, which correspond to electrons in n-type and holes in p-type
semiconductors. Finally, there are minority carriers, which are holes in n-type and electrons in p-type semiconductors. The majority carrier is usually orders of magnitude larger than the minority and intrinsic carriers, although minority carriers are a vital part of the inversion process. Having a large proportion of majority carriers will cause a shift in the Fermi energy level. It is self-evident, that if there are more electrons in the conduction band that the energy level (Fermi level) where there is a 50% chance of being occupied must shift towards the conduction band. Similarly, the Fermi level will shift closer towards the valence band when holes are the majority carrier. Typically, doping is done using ion implantation.

1.4 Wafer Fabrication

In order to make an integrated circuit, the first step is to fabricate a wafer. These wafers are composed of single crystal silicon. The silicon comes from sand which contains SiO₂. In order to get high grade silicon, the silicon dioxide must first be reduced:

\[ \text{SiO}_2 + 2C \rightarrow \text{Si} + 2\text{CO} \]

The silicon must then be purified again using hydrogen chloride:

\[ \text{Si} + 3\text{HCl} \rightarrow \text{SiHCl}_3 + \text{H}_2 \]

The SiHCl₃ is then distilled and reduced until only high grade polycrystalline material is left over.

Once electronic grade silicon has been obtained, it is possible to grow a wafer. The main technique for wafer growth is the Czochralski (CZ) process. In CZ growth, the pure silicon is placed into a crucible and then heated until it melts. A crystal seed is then placed into the molten silicon. Note that the diamond cubic structure of silicon allows for multiple orientations which have different electrical properties. Depending on the device being fabricated with the wafer, a different crystal with a different orientation will be used as the seed. The seed is slowly pulled up until the desired wafer size is solidified on the crystal. From there the crystal structure is
gradually pulled up until a larger crystal is formed. Once the large crystal is cooled it is then cut into thin slices, making the silicon wafers.\textsuperscript{7}

Another method used to make wafers is Float Zone (FZ) refining. In this method, a silicon rod is melted and then recrystallized. The recrystallized rod then goes through a furnace, which only heats one section (zone) at a time. The impurities are thus moved to the bottom of the rod. The rod continues cycle through the furnace until the desired purity has been reached. As a result, the FZ method gives the purest silicon.\textsuperscript{8} However, the impurities found in the CZ process actually improve the durability and reliability of the wafers by increasing the mechanical strength and allowing for internal gettering which will be discussed shortly. Therefore CZ growth is used primarily for industrial applications which don’t require extremely pure silicon.

### 2.2 Impurities and Gettering

In any silicon wafer there will be some impurities that affect the properties of the semiconductor. For instance, doping atoms can be thought of as point defects within the crystal. Foreign atoms can contaminate the crystal by either forming in the lattice or inside the lattice structure (interstitial).\textsuperscript{9} These atoms typically form electronic states inside the band gap which cause poor electronic performance. In addition, these atoms can group together to form precipitates within the crystal, which will cause device failure.

While most foreign atoms have negative effects, not all do. One example is of a beneficial foreign atom is oxygen. In the CZ process, oxygen is dissolved into the crystal from the crucible, forming interstitial impurities. As long as the oxygen amount is small enough to not form precipitates, it can provide two beneficial characteristics. The first is to improve the mechanical durability of the crystal. The second is to add donor states into the crystal, adding extra doping to the crystal. During the CZ process, great care must be taken to not introduce carbon, which will accelerate the formation of oxygen precipitates.\textsuperscript{9}
Following the discussion of impurities, it is important to mention gettering. Gettering is a process used to move impurities from active areas in the device. This is crucial to ensure device reliability and durability. The underlying principle behind gettering is that a damaged section of a crystal lattice will “get” or capture defects and foreign impurities. Since there is already a defect, it takes less energy for impurities to fall into the defect than to form an interstitial defect in the lattice. In external gettering, the back is typically damaged so the front (where the ICs are placed) will not have defects. The “damage” is usually caused by adding argon ions or polysilicon to the back of the wafer which draw out the defects. Alternatively, internal damages can be used to trap and collect defects which typically done with oxygen atoms. The first step is to remove the oxygen from the front of the wafer by heating it, causing the oxygen to move out of the surface. The wafer is then annealed, allowing for the remaining oxygen atoms to cluster together, nucleating. Once the nucleation reaches as critical mass it will then cause defects inside the crystal which can be used for gettering. Internal gettering has become more popular in the recent years due to its simplicity and use of existing elements in the fabrication process.

2.3 Oxide Growth

Once a wafer has been fabricated, the next step is oxidation. An oxide layer provides insulation for the semiconductor which is crucial for capacitors. The primary advantage of silicon versus other semiconductors (germanium) is that it easy combines to make silicon dioxide. By heating up the wafer in a container filled with oxygen, a glass film (thermal oxide), is formed (thermal oxidation). Note that quartz glass is an amorphous and not a solid crystal. Thermal oxidation is preferred over chemical vapor deposition (CVD) because it provides a more uniformed structure that is less subject to change over time. Another critical property of amorphous silicon dioxide is that it provides a mask for doping, due to a low diffusion rates.
There are two main methods of oxidation: dry oxidation and wet oxidation. Dry oxidation occurs by heating the silicon wafer in a chamber filled with oxygen gas. The result is silicon dioxide. Wet oxidation occurs by heating the silicon wafer in a chamber filled with water vapor. The result is silicon dioxide and hydrogen gas. It is crucial to closely monitor the temperature and amounts of each substance in these reactions. In order to get the desired device specifications, exact thicknesses and timings are required. The growth of the thin film and the time required can be modeled using the Deal-Grove model.\textsuperscript{11}

Under the Deal-Grove model, thermal oxidation is broken into transport and reaction processes. The oxygen gas in the chamber moves first (transports) to the silicon surface. Next the oxygen diffuses and dissolves (reaction) into the interface of gas and solid. The Deal-Grove model then gives two useful expressions for the oxide thickness and growth time.\textsuperscript{12} The first, called the parabolic growth regime, is diffusion limited, meaning that the limiting factor is the diffusion of oxygen through the oxide film. The second expression, the linear growth regime, is considered mass transport limited, meaning that the limiting factor is not enough oxygen close to the substrate. The parabolic regime expression is given by:

$$x = Bt$$

where $x$ is the oxide layer thickness, $B$ is the parabolic rate constant that can be found in a lookup table, and $t$ is the time. Similarly, the linear limiting expression is:

$$x = \frac{B}{A}(t + t_0)$$

where $B/A$ is the linear rate constant that can be found in a lookup table, $t_0$ is the initial time, $x$ is the oxide thickness, and $t$ is the time.

The growth of thin films is also highly correlated to temperature.\textsuperscript{12} The linear and parabolic growth rate constants can be related to temperature via the Arrhenius equation:

$$R = R_0 e^{-\frac{E_a}{kT}}$$

Where $R$ is the rate constant, $R_0$ is the initial rate constant, $E_a$ is the activation energy needed to transition from reactant to product, $k$ is Boltzmann’s constant, and
$T$ is the absolute temperature. The activation is usually around 2 eVs whereas the initial rate constants are $\sim 10^4$.

A final consideration in growth of thermal oxides is the pressure. Increasing the pressure will cause more oxygen atoms to move to surface, decreasing the overall time needed to reach a desired thickness. This is important in industrial settings where it is necessary to limit the amount of high temperature the substrate is exposed to, in order to mitigate negative effects on durability. Decreasing the temperature will simultaneously increasing the pressure will allow the wafers to be fabricated in the same amount of time. Another industrial application of pressure dependence is the ability to make very thin films by using a lower pressure. In summary, the Deal-Grove model is used in industrial settings to model the required device parameters.

### 3.1 Material Interfaces

Before getting into the specifics of metal-oxide-semiconductor capacitors it is important to analyze the properties at the interface of different materials. Recall that Fermi energy is the free energy of mobile carriers which implies that at equilibrium, the Fermi level of a solid will be uniform. In addition to the Fermi level, the work function is used to describe the electronic properties at material interfaces. The work function is defined as the energy required to remove an electron from a material in a vacuum. It can be thought of as the energy difference between the Fermi level and the vacuum, and as a result each material will have a different work function. For a given work function, there is also an associated threshold voltage at which current will begin flowing, activating the device. $^{13}$ This threshold voltage is typically a few volts.

Now, with this knowledge, it is possible to analyze the interface of a metal and a semiconductor. Since intrinsic (undoped) semiconductors have a larger work function than metals, when the two materials are placed together, the electrons will flow from the metal to the semiconductor until equilibrium is reached. The electrons
flow from the metal to the semiconductor because the semiconductor has a lower Fermi level (lower energy states). Because electrons are being added to the conduction band of the semiconductor, it is essentially being n doped close to the interface, causing a shift in the Fermi level. Since the Fermi level must remain constant, the bands must “bend” close to the interface. The electrostatic field that is caused by the potential differences between the two materials keeps the “bend” in the bands near the interface.  

Introducing doping to a semiconductor has a similar effect. For example, in a p-type semiconductor, there will be a larger work function, and as a result there will be a steeper “bend” in the band since more electrons will need to travel from the metal for equilibrium. Since p-type semiconductors have holes in the valence band, these extra electrons will recombine with them first. This will cause a reduction in the amount of mobile carriers available near the interface. If the semiconductor is heavily doped (>10^{18} \text{ cm}^{-3}), holes will still remain the majority carrier, but will be reduced. This is called depletion. If the doping level is low enough (<10^{13} \text{ cm}^{-3}), all the holes will be recombined with electrons in the valence band, and the electrons will then move to the valence band. This changes the majority carrier from holes to electrons, making the p-type semiconductor n-type at the interface. This is called inversion.

Similarly, if one has combines a metal to an n-type semiconductor, there will also be a "bend" in the bands of the semiconductor near the interface. Electrons travel from the metal to the semiconductor, entering the conduction band first as there are no holes in the valence band. This will cause an increase in the amount n-type dopants in the semiconductor. This process is called accumulation because the amount of mobile carriers increases. Also, it is possible to have the Fermi level and the work function difference of the materials cancel out with the right doping concentration. Since the effective work function difference is zero, there is no "bend" in the band. This is called flat-band.
3.2 MOSCAP Interface Characteristics

These principles can be applied to a MOSCAP, which is simply a metal-semiconductor interface with an insulator stuck between them. Like the previous cases, the Fermi level must remain constant at equilibrium, creating an electric field between the materials due to the difference in work functions. However, due to the insulator in-between, charges of opposite sign build up on either side, putting some of the electric field in the insulator. This decreases the field penetration in the semiconductor. While the MOSCAP will still exhibit accumulation, depletion, flat band, and inversion, the overall effect of adding an insulator is a reduction of these regimes.

By applying a bias voltage to the MOSCAP it is possible to change the Fermi level. This is due to the energy that is gained by the electrons getting pushed through the field. In this way it is possible to adjust the voltage such that the MOSCAP can undergo accumulation, flat band, depletion, and inversion. Since the amount and polarity of the charge will change depending on whether it is in the accumulation, depletion, or inversion mode, the capacitance will also change. In effect, the applied voltage will control the capacitance.

3.3 MOSCAP CV Curve Characteristics

As an example, consider a MOSCAP with a p-type semiconductor. When a negative voltage is applied, the holes will move towards the interface, causing accumulation. These holes will form an accumulation layer where there is an increase of mobile carriers. As a result, a sheet of positive charge will build up at the silicon-silicon oxide interface, and a sheet of opposite charge will form on the silicon-metal interface. These sheets of charge are similar to a parallel plate capacitor. The equation for capacitance per area $C_{ox}$ is

$$C_{ox} = \frac{\varepsilon_{ox}}{\chi_{ox}}$$

(5)

Where $\varepsilon_{ox}$ is the dielectric constant of the oxide layer (0.34 pF/cm for SiO$_2$) and $\chi_{ox}$ is the oxide thickness.
As the voltage increases (positively), the amount of mobile carriers (holes) will decrease until the flat band is reached. When the voltage increases, the capacitor will enter into depletion. In this regime, the positive voltages will repel the holes from the oxide interface. The region without the holes is called the depletion region. This region has an overall negative charge. As a result the capacitance can again be modeled as a parallel plate capacitor with the depletion capacitance, $C_d$, per area given by:

$$C_d = \frac{\varepsilon_d}{\chi_d}$$  \hspace{1cm} (6)

where $\varepsilon_s$ is the semiconductor dielectric constant and $\chi_d$ is the depletion layer width.

As the voltage gets very large, the MOSCAP will undergo inversion. In this regime, the positive bias is such that the majority carriers (holes) are pushed away from the interface and the minority carriers (electrons) are attracted to the interface. The sheet of electrons form an inversion layer within the depletion layer, halting the growth of the depletion layer. The result is that the semiconductor becomes n-type at the interface. The capacitance per area, $C_s$, is given by:

$$C_s = \frac{\varepsilon_d}{\chi_s^{\text{max}}}$$  \hspace{1cm} (7)

where $\chi_s^{\text{max}}$ is the maximum depletion layer width.

To measure the CV curve of a MOSCAP, the silicon-silicon oxide wafers are coated with a metal (aluminum) or polysilicon (used for transistor fabrication). To improve conductivity, the back of the silicon wafer may also be coated with a metal. The two metal contacts are then connected to an impedance analyzer which applies a voltage sweep, measures the current, and calculates the capacitance by integrating the current over time. In a perfect MOSCAP there would be no current inside the oxide insulator, however in real world systems there is typically a small leakage current which is accounted for in most modern measurement systems.

The MOSCAP CV characteristics are illustrated by the following chart:

---

$\varepsilon_s$  \hspace{1cm} $\chi_d$  \hspace{1cm} $\varepsilon_d$  \hspace{1cm} $\chi_s^{\text{max}}$
CV measurements are typically done at low (quasistatic) and high frequencies. At low frequencies (Hz range), the mobile carriers will be at the oxide interface during accumulation and inversion, making the MOSCAP capacitance equal to the total capacitance of the oxide layer. In the depletion mode, since there are no mobile carriers at the oxide interface, the total capacitance will be the series combination of the depleted region and the non-depleted region, decreasing the overall capacitance. In high frequency measurements (MHz range), the total capacitance for the accumulation and depletion modes are the same as in the low frequency range. However, in the inversion mode, the high frequency is too fast for an inversion layer to form. As a result, the total capacitance for the inversion mode is the same as for the depleted mode, the series combination of depletion and oxide capacitance.
4.1 Oxide Charges

Now, the CV characteristics of MOSCAPs are further complicated by the introduction of trapped charges in the oxide layer during fabrication. These charges are effectively trapped inside the oxide layer, near the oxide interface, and therefore affect the depletion region by shifting the potential. These charges are called oxide charges and are divided into four types: interface trapped charges (also known as surface states), fixed oxide charges, oxide trapped charges, and mobile oxide charges. Surface states are caused by structural defects and are typically removed using a low temperature annealing process. Oxide trapped charges are holes or electrons that are trapped from ionizing radiation. These can also be removed using low temperature annealing. Mobile oxide charges are contaminants from Na\(^+\), Li\(^+\), or K\(^+\) ions that can occur during the fabrication process. These charges are often identified using a bias stress test. Fixed oxide charges are typically positive charges located near the semiconductor-oxide interface and are a result of the thermal oxidation process. These charges are the primary interest of this paper.

It is helpful to analyze the effect of the fixed oxide charges using the flat band condition. The flat band capacitance, \(C_{FB}\), can be shown to be:

\[
C_{FB} = \frac{C_{\text{max}}}{1 + \frac{C_{\text{max}}/C_{\text{min}} - 1}{2\sqrt{n_i(N_A - N_D)/n_i}}}
\]

where \(C_{\text{max}}\) and \(C_{\text{min}}\) are the maximum and minimum capacitance, \(N_A\) and \(N_D\) are the acceptor and donor concentrations, and \(n_i\) is the intrinsic carrier concentration.

Once the flat band capacitance has been calculated, the flat band voltage can be visually determined from the CV curve. If the charges are assumed to be stuck near the interface, an expression for the flat band voltage, \(V_{FB}\), can be given as:

\[
V_{FB} = \phi_M - \phi_S - \frac{Q_f}{C_{\text{ox}}}
\]

where \(\phi_M\) and \(\phi_S\) are the work functions for metal and semiconductor, and \(Q_f\) is the fixed charge density near the surface of the oxide. For a given MOSCAP, the change in flat band voltages, called the flat band shift, can be found using:

\[
|V_{FB}| = \frac{Q_f}{C_{\text{ox}}}
\]
Finally, in the inversion regime, it is possible to calculate the threshold voltage, $V_T$, of the MOSCAP. This equation is given by:

$$V_T = V_{FB} + 2\phi_F + \frac{\sqrt{4qN_B\epsilon_F\phi_F}}{C_{ox}}$$  \hspace{1cm} (11)

where $\phi_F$ is the work function of the Fermi energy level, and $N_B$ is the substrate charge concentration. The important feature of this expression is that the change in the threshold voltage will be the flat band shift for the same MOSCAP with different trapped charge amounts.

So,

$$|V_T| = |V_{FB}|$$  \hspace{1cm} (12)

Since the threshold voltage is a critical parameter for MOS devices\textsuperscript{14}, (especially CMOS transistors), it will be possible to relate a change in fixed oxide charges, $Q_f$, with the overall performance of these devices using the above equations and two MOSCAP CV curves (with and without oxide charges).

**Methods**

In order to test how the oxide charges affect device performance, capacitance and voltage curves were taken. CV curves, as explained previously, show the effective capacitance as a function of different voltage biases applied to the metal contacts. By measuring the CV curve for a MOSCAP with trapped charges, and comparing it to one without charges, it would be possible to see a change in characteristic behavior of the capacitor, and therefore infer the change in performance.

Originally, an experiment was planned using MOSCAPs manufactured at SHARP Labs in Camas, Washington. These were made as P type silicon with boron doping and oxide thickness of 110 nm, (see appendix). These wafers were then broken into smaller pieces. The backsides of these pieces were scratched and light layers of a conductive silver paste were added, in order to improve the metal contacts. Once the silver had dried after about 3 days, the wafers were placed into a measuring stage, where the wire leads were placed on the metal contacts using a
microscope. After the connection was made, a HP 4275A impedance analyzer was used to measure the capacitance from -5 to +5 Volts using an alternating current of 1 MHz. These readings were then connected to ICS software on a computer.

Unfortunately, after repeated trials, there was no viable data measured just static noise. After switching to a HP 4192A and then a Solartron SI 1260, no results were measured. During the measurements with the Solartron, there was a failure, likely a blown capacitor, which caused the Solartron to be under repair for a few days. After trying again with the Solartron without results, a new strategy was devised.

The measurements could be simulated using software tools developed for MOSCAPs. The particular software used was based on PADRE. Two measurements were taken using simulated MOSCAPs: one with trapped charges, and another without trapped charges. In this way, the only variable changed was the amount of charges introduced into the oxide layer. A trapped charge density of $10^{16}$ cm$^{-3}$ was added in order to provide an extreme difference between the two simulated MOSCAPs. In the simulated MOSCAPs, the wafers were P doped, with an insulator thickness of 0.1 micrometers. The measurements were simulated with a bias of -5 to +5 Volts, with an alternating current of 5 MHz and 1 Hz.
Results and Discussion

The results of the simulations were then calculated using the PADRE\textsuperscript{18} and MOSCap\textsuperscript{19} software and are shown in the following graph:

\textit{Figure 2: CV response of contaminated (blue/yellow) and uncontaminated (magenta/green) MOSCAPs.}

On the graph, the blue line corresponds to the high frequency response and the yellow line corresponds to the low frequency response of the MOSCAP with trapped charges. Also, the magenta line corresponds to the high frequency response and the green line corresponds to the low frequency response of the MOSCAP without trapped charges. Note the accumulation regime at negative voltages, depletion, flat band, and inversion at positive voltages.
As is visible from the graph, there is a noticeable shift in the CV curves of approximately 1 Volt. This shift corresponds to a change in the flat band voltage (equation 10). Since the threshold voltage is dependent on flat band voltage (equation 12), there will also be a shift in threshold voltage. As noted earlier, the threshold voltage is a critical parameter used in field-effect devices. Since the typical threshold voltages are close to 0.7 mV, a shift of 1 Volt will cause a dramatic change in the voltage needed to activate the device. This will have a negative effect on the reliability of the devices because they will be operating outside the designed tolerances. As a result, the trapped oxide charges will have a significant negative impact on device performance.

**Conclusion**

During the fabrication of MOSCAPs contaminants can be trapped inside the oxide layer. By introducing oxide charges into the MOSCAP the threshold voltage shifts causing device instability and general reliability problems. As a result, MOSCAP devices suffer worse performance with trapped charges than without trapped charges. For large scale industrial applications, this can be catastrophic. It is crucial for production that these trapped charges be kept to a density under $10^{15}$ cm$^{-3}$. With modern clean room procedures and annealing processes it is standard to have densities well below $10^{15}$ cm$^{-3}$. Nonetheless, trapped charges can still cause problems in device performance if they are not carefully tested for.

**Recommendations**

One recommendation is the need for physical measurements. Do to equipment malfunctions and general technical issues; I was unable to use the impedance analyzer with the fabricated wafers. It would be very beneficial to
compare those readings to those of the simulated results to determine the accuracy of the measurements.

Another recommendation would be to study the effects of mobile charges such as sodium or potassium ions. It would be possible to contaminate the MOSCAP with a sodium or potassium bath, and measure the shift in the CV curve. This shift could be used to calculate the shift in the flat band voltage, as well as determine the mobility and diffusivity of the charges.

Acknowledgements

I would first like to acknowledge my advisor Dr. Solanki for the use of his equipment and continued guidance with my project. I would also like to thank Dr. Evans and Sharp Labs for fabricating the MOSCAPs and starting me on this project. In addition, I would like to thank Prasanna Padigi for teaching me how to use the lab equipment and for advising me on any questions I had. Finally, I would like to thank Dr. Fallon, Dr. Wolf, Dr. Estes, and the Portland State University Urban Honors College for their guidance and support of my undergraduate thesis.
<table>
<thead>
<tr>
<th>Step</th>
<th>Process Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Start material: Prime 100 p-51 - 100 ohm cm</td>
</tr>
<tr>
<td>2.</td>
<td>2D CV dots lot</td>
</tr>
<tr>
<td>3.</td>
<td>Plasma clean</td>
</tr>
<tr>
<td>4.</td>
<td>SC1/SC2 clean</td>
</tr>
<tr>
<td>5.</td>
<td>HF dip</td>
</tr>
<tr>
<td>6.</td>
<td>Thermal oxidation 15min</td>
</tr>
<tr>
<td>7.</td>
<td>Frontside PR strip</td>
</tr>
<tr>
<td>8.</td>
<td>Frontside PR strip</td>
</tr>
<tr>
<td>9.</td>
<td>Plasma clean</td>
</tr>
<tr>
<td>10.</td>
<td>E-beam evaporation, Ti/AI: 3nm/150nm</td>
</tr>
<tr>
<td>11.</td>
<td>Photo lithography, 250um square array with 200um spacing</td>
</tr>
<tr>
<td>12.</td>
<td>RIE Ti/Pt etch, 45sec. In situ PR strip</td>
</tr>
<tr>
<td>13.</td>
<td>DI water rinse and SRD</td>
</tr>
<tr>
<td>14.</td>
<td>Forming gas anneal, 30min @450°C</td>
</tr>
</tbody>
</table>

**Notes:**
- SHARP
- Appendix A - Wafer Fabrication Data Sheet
Appendix A - Continued
References


18. Mark R. Pinto; kent smith; Muhammad Alam; Steven Clark; Xufeng Wang; Gerhard Klimeck; Dragica Vasileska (2014), "Padre," https://nanohub.org/resources/padre.