Limitations of the High-Low C-V Technique for MOS Interfaces With Large Time Constant Dispersion

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Limitations of the High–Low $C–V$ Technique for MOS Interfaces With Large Time Constant Dispersion

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Abstract—We discuss the limitations of the high–low $CV$ technique in evaluating the interface trap density ($D_{IT}$) in MOS samples with a large time constant dispersion, as occurs in silicon carbide (SiC). We show that the high–low technique can seriously underestimate $D_{IT}$ for samples with large time constant dispersion, even if elevated temperatures are used to extend the range of validity.

Index Terms—AC conductance technique, interface states, interface traps, silicon carbide (SiC), wide-bandgap semiconductor.

I. INTRODUCTION

Silicon carbide (SiC) MOS technology has made great progress since the first MOS devices were reported in the 1990s [1], but despite the progress, high interface trap density ($D_{IT}$) at the SiC/SiO$_2$ interface continues to be a concern for SiC MOSFETs. Even though silicon and SiC share the same native oxide, the properties of the interfaces are different. Yet, most techniques used to characterize the SiC MOS interface are borrowed directly from the silicon repertoire. The high–low $CV$ method is one such technique.

Several reports in the literature stress the importance of interpreting SiC high–low data with care [2],[3]. $D_{IT}$ data are valid only over the energy range where the high-frequency capacitance does not include any trap contribution and the low-frequency capacitance includes the contribution of all traps within a few $kT$ of the Fermi level. The long response times of deep states make it hard to achieve true low-frequency conditions at room temperature when the Fermi level is deep in the bandgap. When the Fermi level is close to the majority carrier band, traps at the Fermi level respond quickly, making it hard to achieve true high-frequency conditions at those biases. Both high- and low-frequency conditions are simultaneously satisfied only over a narrow energy window, and the $D_{IT}$ extracted from the high–low technique is only valid within this window.

In addition to the long response times of the interface states, the SiC/SiO$_2$ interface typically has a large time constant dispersion [2], [4], [5]. In silicon, the time constant dispersion is attributed to a variation in the density of fixed charges across the interface [6]. The time constant dispersion in SiC is typically two to three times larger than that in silicon, and its origin will be discussed in a separate paper [7]. In this paper, we focus on the effect of this time constant dispersion on data interpretation using the high–low $CV$ technique.

II. THEORETICAL BACKGROUND

Fig. 1 shows the small-signal ac equivalent circuit of a MOS interface with interface states. In the high–low $CV$ technique, the terminal capacitances $C_{LF}$ and $C_{HF}$ are measured as a function of gate voltage at two frequencies $\omega_{LF}$ and $\omega_{HF}$. The interface state density $D_{IT}$ at a given bias is calculated using [8], [9]

$$D_{IT} = \frac{1}{q} \left( \frac{C_{ox}C_{LF}}{C_{ox} - C_{LF}} - \frac{C_{ox}C_{HF}}{C_{ox} - C_{HF}} \right).$$  \hspace{1cm} (1)$$

The surface-potential–gate-voltage ($\phi_s–V_G$) relation of the sample is then used to correlate $D_{IT}$ at a given bias with the position of the Fermi level in the bandgap at that bias. To calculate the $\phi_s–V_G$ relation of the sample, the high-frequency curve is treated as the ideal curve with no interface trap response, i.e., $C_{IT}(\omega_{HF}) = 0$. This makes the equivalent circuit at high frequency a series combination of $C_{OX}$ and $C_D$. The surface potential is then calculated from the extracted $C_D$ at each gate voltage. This method provides erroneous surface potential...
data close to the majority band, since $C_D$ has an interface trap capacitance contribution. This extra capacitance shifts the energy positioning of the $D_{IT}$ points closer to the majority band. Another common technique uses the low-frequency $CV$ curve to calculate the $\phi_s-V_G$ relation [10].

One of the most rigorous methods for evaluating the silicon MOS interface is the ac conductance technique [6]. This technique measures the equivalent parallel conductance $G_p$ and capacitance $C_p = C_D + C_{IT}$ at several fixed gate voltages as a function of frequency. To model the frequency dependence of $G_p$ and $C_p$, the theory assumes a random variation of fixed oxide charge across the interface, which produces a random variation in surface potential characterized by a Gaussian probability distribution.

For a $p$-type sample biased in depletion, the oxide charge across the interface, which produces a random underestimate, following this broadening causes the high–low technique to be typically around two. The same broadening is also evident in samples having a large time constant dispersion. Fig. 2(a) shows the $G_p/\omega$ curves on a 4H-SiC MOS capacitor biased in depletion. The sample was formed on a $p$-type SiC epilayer doped $\sim 2 \times 10^{16}$ cm$^{-3}$ on a $p^+$ (0001) SiC substrate. The oxide was grown by pyrogenic oxidation at 1150 °C and annealed in nitric oxide at 1175 °C for 2 h. The sample was measured at room temperature in the dark using an HP 4284A LCR meter. The extracted parameters are $D_{IT} = 3.4 \times 10^{12}$ cm$^{-2}$ · eV$^{-1}$, $\sigma_{us} = 4.0$, and $\tau_p = 4.9$ μs. The extracted parameters are used to generate a theoretical $C_{IT}(\omega)$ curve. (b) (Points) Measured $C_p(\omega)$, (solid line) calculated $C_{IT}(\omega)$, and (dashed line) extracted $C_D(\omega)$. The $C_{IT}(\omega)$ curve is generated using (2b) with the fitting parameters from the $G_p/\omega$ data. $C_D$ is obtained by subtracting the calculated $C_{IT}$ from the measured $C_p$. As expected, $C_{IT}$ is independent of frequency$^4$, confirming that all the interface trap capacitance has been removed.

III. RESULTS AND DISCUSSION

To examine the accuracy of the high–low $C-V$ technique in samples with large time constant dispersion, we assume a $\sigma_{us}$ distribution. For a $p$-type sample biased in depletion, the interface state admittance can be written [6]

$$\frac{\langle G_p(\omega) \rangle}{\omega} = \frac{q}{2} \int_{-\infty}^{\infty} \frac{D_{IT}}{\omega \tau_p} \ln \left[ 1 + \omega^2 \tau_p^2 \right] P(u_s) du_s$$  \hspace{1cm} (2a)

$$\langle C_{IT}(\omega) \rangle = \int_{-\infty}^{\infty} \frac{D_{IT}}{\omega \tau_p} \tan^{-1}(\omega \tau_p) P(u_s) du_s$$  \hspace{1cm} (2b)

where

$$\tau_p = \frac{1}{c_pN_A} \exp(u_s)$$  \hspace{1cm} (2c)

$$P(u_s) = \frac{1}{\pi \sigma_{us}^2} \exp \left[ -\frac{\left( u_s - \langle u_s \rangle \right)^2}{2\sigma_{us}^2} \right].$$  \hspace{1cm} (2d)

Here, $u_s$ is the surface potential normalized to $kT/q$, $\tau_p$ is the time constant for hole capture by states at the Fermi level, $c_p$ is the capture coefficient for holes, $N_A$ is the ionized acceptor concentration, and $\sigma_{us}$ is the standard deviation of surface potential or, in general, a measure of the time constant dispersion.

The $G_p/\omega$ data obtained by measurement are fitted to (2) to extract the interface state parameters $D_{IT}$ and $c_p$ at the energy of the Fermi level at each gate voltage [4].

In 4H-SiC, the $G_p/\omega$-versus-ω curves at each bias are broader than that in silicon [2], [4], [5], indicating a larger time constant dispersion. Fig. 2(a) shows the $G_p/\omega$ data (points) and the best fit using (2a) with a $\sigma_{us} = 4$. In contrast, for silicon $\sigma_{us}$ is typically around two. The same broadening is also evident in the capacitance data [Fig. 2(b)]. As will be shown in the following, this broadening causes the high–low technique to underestimate $D_{IT}$ in samples having a large time constant dispersion.

To examine the accuracy of the high–low $C-V$ technique in samples with large time constant dispersion, we assume a $\sigma_{us}$ distribution typical of 4H-SiC, and we simulate $C-V$ curves at two frequencies: $\omega_{HF} = 2 \pi \times 10^6$ rad/s (1 MHz) and $\omega_{LF} = \pi$ rad/s (0.5 Hz). The simulations are performed using the equivalent circuit in Fig. 1, with $G_p$ and $C_{IT}$ calculated using (2a)–(2d) and the $C_D$ obtained using the “exact” MOS $C-V$ theory [11]. We then regard these simulated curves as experimental data and perform the standard high–low analysis to extract an “experimental” $D_{IT}$ versus $E$. Finally, we compare the “experimental” $D_{IT}$ to the $D_{IT}$ originally assumed in the simulation. The simulations are performed for different $\sigma_{us}$ values and at different temperatures to evaluate the range of accuracy of the high–low technique. For illustration purposes, these simulations are performed for p-type MOS capacitors, and the temperature dependence of the ionized acceptor concentration in 4H-SiC is explicitly included [11].

The dashed line in Fig. 3(a) shows the assumed $D_{IT}$ distribution. This distribution is representative of the $D_{IT}$ typically reported on p-type 4H-SiC [2], [4], [5]. The capture coefficient $c_p$ is assumed constant throughout the bandgap. The results of simulated high–low measurements at room temperature are shown in Fig. 3(a). For a $\sigma_{us}$ of four, typical of 4H-SiC [2], [3], [4], the extracted $D_{IT}$ is underestimated at all energies. The reason for the error can be seen from Fig. 3(b), which shows the calculated $C_{IT}(\omega)$ at three bias points, labeled A, B, and C in Fig. 3(a). The extracted $D_{IT}$ will only be accurate if $C_{IT}/qD_{IT}$
Fig. 3. (a) $D_{IT}$ as a function of energy for the assumed $D_{IT}$ distribution (dashed line) and deduced from the high–low technique for $\sigma_{us} = 4$. The relative accuracy of the deduced $D_{IT}$ is shown in the inset. (b) $C_{IT}$ versus frequency for $\sigma_{us} = 4$ at bias points A, B, and C in (a). The measured $D_{IT}$ at points A, B, and C is proportional to the difference between the $C_{IT}$ values at the low and high frequencies [the dashed lines in (b)] and is accurate only when the low-frequency normalized $C_{IT}$ is equal to unity and the high-frequency normalized $C_{IT}$ is zero. For samples with large time constant dispersion, these conditions may not be met for any bias point, as illustrated in part (b). All curves were simulated using $a_{CP} = 1 \times 10^{-11}$ cm$^3$/s, which is representative of many 4H-SiC samples [2], [4], [5].

Fig. 4. Extracted $D_{IT}$ versus energy for a $\sigma_{us} = 4$ sample at different temperatures. The points illustrate how data are often reported in the literature. To obtain a reliable estimate of $D_{IT}$, it is useful to make overlapping measurements at several temperatures and accept only the highest $D_{IT}$ obtained at any given energy.

Fig. 5. Normalized $C_{IT}$ as a function of frequency for different time constant dispersions. As $\sigma_{US}$ increases, the curves become broader; more traps begin to respond at the high frequency ($2\pi \times 10^6$ rad/s), and fewer states respond at the low frequency ($\pi$ rad/s). This leads to an underestimation of $D_{IT}$.

is simultaneously unity at the low measurement frequency $\omega_{LF}$ and zero at the high measurement frequency $\omega_{HF}$. This condition is clearly not met for bias points A or C. This is due to the large time constant dispersion for interface state response, which causes a very gradual transition from low- to high-frequency behavior. Using fixed measurement frequencies of $\omega_{HF} = 2\pi \times 10^6$ rad/s and $\omega_{LF} = \pi$ rad/s, there is only a narrow range of bias for which the high- and low-frequency requirements can be satisfied simultaneously.

High–low measurements on SiC are often performed at multiple temperatures to extend the energy range over which $D_{IT}$ can be measured. Fig. 4 shows the simulated high–low measurements on a sample with $\sigma_{us} = 4$ and the $D_{IT}$ distribution in Fig. 3(a), performed at several temperatures. The plotted points illustrate the way results are often reported in the literature, and an abrupt jump in $D_{IT}$ is observed at the point where measurement temperature is changed. Measurements at a single temperature provide valid data over only a limited energy range, but a fairly accurate picture can be obtained by combining data taken at several temperatures and accepting only the highest $D_{IT}$ at any given energy. This is shown in Fig. 4 by extending the 300 °C data to lower energies (solid line) and adding additional data at 150 °C (dashed line).

Fig. 5 shows the $C_{IT}$ versus $\omega$ for several values of $\sigma_{US}$. As $\sigma_{US}$ increases, more states respond at the “high” frequency, and fewer states respond at the “low” frequency, increasing the error in $D_{IT}$. Note that the bias point in Fig. 5 is close to optimum, in that the capacitance transition occurs midway between $\omega_{HF}$ and $\omega_{LF}$. As bias is varied, the capacitance transition moves to higher or lower frequencies, increasing the error, as illustrated in Fig. 3(b). When the high–low technique is performed at elevated temperatures, the range over which the high–low data are valid shifts toward midgap, as can be seen in Fig. 4. This is because the higher thermal energy reduces interface state response times.

In the high–low technique, there is no direct method to determine the range over which the data are valid, and this
is a major weakness of the technique. The ac conductance technique avoids this problem, since data are obtained over a range of frequencies, rather than at two discrete frequencies, and the entire $G_P(\omega)$ and $C_{TT}(\omega)$ curves are revealed. $D_{TT}$ data are extracted only at those biases for which a clear conductance peak is observed versus frequency. This restricts the range of energies over which conductance data can be obtained. The conductance technique is not inaccurate outside this range, it simply provides no data for biases that are too close to the majority carrier band or too deep in the bandgap. (The energy range can be shifted closer to the band edge by measuring at low temperatures or deeper into the bandgap by measuring at high temperatures.) Unlike the high–low technique, the conductance technique makes no a priori assumptions regarding frequency response of the states. A comparison of the high–low and conductance techniques for the analysis of SiC MOS devices is given in [2].

Determination of surface potential is an important part of the data reduction process, since surface potential is used to calculate the bandgap energy of the interface states opposite the Fermi level. In the conductance technique, one can subtract the interface state capacitance $C_{TT}(\omega)$ from the measured parallel capacitance $C_P(\omega)$, leaving a frequency-independent depletion capacitance $C_D$, as illustrated in Fig. 2(b). This depletion capacitance is then used to determine the surface potential. In the high–low technique, $C_D$ is inferred from the high-frequency capacitance under the assumption that none of the interface states can follow the signal. This assumption fails when the Fermi level is near the band edge. Consequently, $C_D$ is overestimated, and the surface potential is thought to be closer to accumulation than it actually is. Like the $D_{TT}$ errors discussed previously, there is no direct way to determine whether the energy position of states near the band edges is being determined correctly.

In this paper, we have considered a p-type sample, but our results apply equally to n-type samples. Our conclusions are relevant for any MOS interface with large time constant dispersion, such as MIS capacitors on III–V semiconductors.

**IV. CONCLUSION**

The high–low technique must be used with caution on samples with large time constant dispersion. This is particularly true of MOS interfaces on SiC and quite likely true for interfaces on III–V semiconductors as well. It is not possible to determine directly from the high–low data over what energy range the data are valid, compounding the problem. The ac conductance technique measures admittance over a range of frequencies and makes no a priori assumptions about the frequency response of the states. For this reason, it is the preferred technique in cases where large time constant dispersion is present.

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**REFERENCES**


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