Deep-Level Transient Spectroscopy

3rd year Physics Laboratories



Last compiled August 8, 2017

Contents

1	Introduction	3			
2	Some words about third year labs				
3	Safety Notes 3.1 Personal protective equipment (PPE)	3 4			
4	Theory 4.1Semiconductors4.2Schottky barrier diodes4.3Capacitance and the depletion width4.4Defects and deep-level traps4.5Deep level transient spectroscopy (DLTS)	4 4 5 6 7 8			
5	Experimental Work5.1The racks of electronics5.2The cryostat and sample holder5.3SRIM simulation5.4Performing the C-V measurement5.5Analysing the C-V data5.6DLTS measurement5.7Performing a C-T measurement	13 13 13 14 15 16 17 18			
6	Analysing the DLTS data 1				
7	Overview	19			
8	Important Numbers				
Re	References				

1 Introduction

Semiconductor devices have revolutionised the world in which we live and have become such a part of our everyday lives that we often take it for granted that they can and will perform the task that they were designed for. However, due to their high quality the semiconductors wafers used are extremely sensitive to defects arising from contamination, from impurities and ion bombardment.

In this experiment you will examine how the static capacitance -using Capacitance-Voltage (CV)- and transient capacitance -using Deep-Level Transient Spectroscopy (DLTS)- of a metalsemiconductor diode (Schottky diode) varies with applied bias, temperature and time. These measurement will allow you to gain a good understanding of the nature and effect of the defects present in the silicon semiconductor.

DLTS is a highly sensitive analysis technique that is used on a daily bases by the semiconductor industry. If you have a spare moment you might like to ask Prof. Google or Dr. Google Scholar where it has been used. If you get around to doing this you will see papers dating back to 1974 when D.V. Lang first pioneered the technique [1].

2 Some words about third year labs

These notes don't contain all the information you need - they're here to define the terms for you, and to give you an idea of the direction you should be taking. You'll find that you need to go foraging for more detailed information in other places - often in the references, and even more often in the rubble of your demonstrators' minds. You're going to learn a lot while you're doing that, so don't waste the effort - share all the work you're doing with us in your report! Remember that your report has to be self-contained. It doesn't have to be beautiful, but it should be clear and detailed. Perhaps the most important thing to remember is that when you write your report you're trying to teach your reader what you did, why you did it, and what you learned. That way, we get to learn something too! Enjoy!

We in part3 would love it if you could bring any mistakes to the attention of your demonstrators and email suggestions to part3@physics.unimelb.edu.au

3 Safety Notes

Liquid nitrogen has two main hazards associated with it: **frostbite** and **asphyxiation**. These are related to its low boiling point (-196°C, 77 K) and its large gas to liquid volume ratio (682, at 1 atm and 15° C).

Frostbite: The eyes are particularly at risk from frostbite (cold burns) due to their moisture content. Liquid nitrogen rapidly boils when it comes in contact with an object at room temperature and will splatter or spit. There is a risk of frostbite not only from direct contact but also from indirect contact with a surface that has been cooled by liquid nitrogen. Objects less than about -20°C will cause cold burns and if there is sufficient moisture on your skin you may become frozen to the object.

Asphyxiation: When liquid nitrogen evaporates it displaces air and reduces the oxygen concentration in the air. Symptoms of asphyxia can be sudden in the case of a deep inhalation of air with a low

oxygen content or gradual asphyxia where the symptoms are more subtle. For further information please consult the MSDS (Material Safety Data Sheets) for liquid nitrogen in the safety folder.

The dewar must be placed on level surface at all times except during transport, preferably the ground.

The room in which the dewar is stored and experiments are carried out must be kept well ventilated with all doors open while people are inside. This is to ensure that the oxygen content of the room does not drop below 18% due to ANY spill including the worst case scenario where the entire contents of the dewar evaporates.

3.1 Personal protective equipment (PPE)

Note: You do not need to buy/supply your own PPE. There is PPE available for the short time when you'll be refilling the dewar with liquid nitrogen.

The following safety equipment is only required when there is exposed liquid nitrogen present (e.g. when the cap or cryostat is not inserted into the dewar, during refilling):

Gloves: Appropriate gloves should be worn when handling the dewar.

Face and eye protection: A full face shield conforming to AS 1337 must be worn.

Clothing: Loose dry clothing that completely covers the arms and legs must be worn. Trouser legs preferably should not have cuffs that may trap liquid nitrogen.

Footwear: Shoes that completely cover the feet must be worn. (This goes for all experiments at all times in lab, too!)

4 Theory

Get ready, this theory section is going to be the most robust and detailed in part 3 labs.

Even knowing where to start when explaining DLTS can be tough. Remember to discuss how best to understand it with your demonstrators.

4.1 Semiconductors

You will find a good introduction to this topic in Streetman [2], its probably a good idea to have a bit of a read of this text at some point (hmmm, even wikipedia has some interesting stuff on it). So what is a semiconductor? Well you would probably be aware that they turn up in pretty much everything these days (shoes, cats... even the humble laptop has some inside), so what makes them so special? Why are they essential to the modern electrical device?

Question 1 *Ok, so what is a semiconductor? Make your description relatively comprehensive and cover at least the following topics:*

• Include a diagram of the band structure

- Compare this band structure to conductors and insulators
- The importance of the bandgap
- Energy levels within a semiconductor
- A description of the Fermi level, and what influences its location
- *The difference between doping and implantation into a semiconductor*
- A description of *defects*. What are they? What effect do they have?

This may seem like a lot, but having a good grasp of all of these before moving on is **essential** for this experiment.

Question 2 We are using silicon in our investigations in these couple of weeks. Why silicon? Germanium has become the subject of more intense research lately. Why? Hint: Germanium was initially suggested as the preferred material in semiconducting devices, but we're only just coming back to it now.

Right, so now we understand what a semiconductor is, time for the next question: how can we employ its fantastic qualities for our benefit? We therefore need to consider *devices*. We will now consider the Schottky barrier diode and how this device operates.

4.2 Schottky barrier diodes

The electronic definition of a diode is a device that allows current to flow in one direction (forward bias) and not in the other (reverse bias), The same is true for a Schottky barrier diode. To understand why this occurs one has to look at the band structure of a metal-semiconductor junction and what occurs when a bias is applied across the junction.

Question 3 In this experiment we will be using n-type (phosphorous doped) silicon as the semiconductor and the metal is gold. By keeping this in mind, describe the Schottky diode effect using these diagrams:

- a well labeled diagram of the band structure of both a metal and an n-type semiconductor separated in space.
- a diagram of the band structure when the metal is brought into contact with the n-type semiconductor.

In your diagram you should include these important terms: Fermi Level, Conduction Band, Valance Band, Metal Work Function (ϕ_m), Semiconductor Work Function (ϕ_s), Barrier Height (ϕ_b), Depletion Width (ω), and Built in Voltage (V_{bi}).

It should be noted that the work function of a metal describes the binding energy of the electrons and is hence related to their Fermi level. Also when the metal and semiconductor come into contact it becomes energetically favorable for electrons in the material to drift in such away that the Fermi levels line up across the interface. This drift creates a charge difference over the interface which creates a Built-in Voltage (V_{bi}). In n-type silicon, if the work function of the metal is larger than that of the

silicon, the electrons in silicon drift away from the metal-silicon interface region leaving behind a net positive charge in the silicon near this region. This region is known as the Depletion Region (depleted of majority charge carriers) and characterized by a width (ω).

In this experiment we will be looking at the electrical properties of a single Schottky barrier diode. This will be done by applying a voltage, or External Bias, or voltage pulses to the gold contact.

Question 4 Using a similar diagram to that used in the previous question, describe what occurs when a reverse External Bias (V) is applied across the metal-semiconductor junction.

Question 5 By considering what happens when forward and reverse bias is applied to our Schottky diode, what is the significance of the Built in Voltage V_{bi} and the Barrier Height ϕ_b in relation to current flowing through our devices? How does this lead to the diode effect?

Question 6 *What happens if we bring a metal with a smaller work function in contact with a n-type silicon?*

The Schottky diodes made for this experiment were fabricated by first cleaning the silicon wafer with solvents to remove any organic contaminates on the surface (degreasing). Next the thin native oxide (10-20 Å) that exists on the surface of bare silicon is etched off with hydrofluoric (HF) acid (very dangerous stuff!) and the sample is promptly transferred to a thermal evaporator for gold deposition through an aluminum mask.

4.3 Capacitance and the depletion width

Now, it should be understood from the previous section that, depending on the individual properties of the material and if an external bias is applied, a depleted region will form at a metalsemiconductor interface. The width of this region is related to these material properties and is given by [3]:

$$\omega = \sqrt{\frac{2\epsilon}{qn}(V_{bi} - (\frac{k_b T}{q}) - V)} \tag{1}$$

Where:

- ϵ is the dielectric constant of the semiconductor
- n the active dopant concentration in the semiconductor
- V_{bi} is the built-in voltage
- and V the externally applied bias

An important effect of the Schottky diode is that a capacitance will form across this depletion region.

Question 7 What exactly is capacitance? What is the formula for a parallel plate capacitor? How is our diode similar to a parallel plate capacitor?

Question 8 *By considering equation 1 find an expression for the capacitance over our device, note that the area* (A) *of the diode is known and can be simply measured.*

Right, so now we should have a direct relationship between the capacitance of our device, C, and externally applied bias, V. Armed with this expression we can measure many properties of the semiconductor by slapping a metal layer on top of the sample (thus forming a Schottky diode) and measuring how the capacitance varies with the applied bias. This is the key of the CV and DLTS measurements.

Lets take a closer look at this relationship. What are the thing we do not know in Equation 1? Well, here in this experiment we can precisely hold the diode at a known temperature, we can record the applied bias, and the importance of this equipment is that we can also accurately measure the diodes capacitance. So all this leaves is n, the doping concentration, and Vbi, the builtin voltage.

Question 9 Lets first consider the simple case where n is constant throughout the silicon bulk. From our expression, how can we calculate n and V_{bi} from measured capacitances (C) and applied biases (V)? Hint: try rearranging the expression in terms of $1/C^2$ and V.

Question 10 What happens now if n changes throughout the bulk of the silicon? How can n be measured now? Hint: We will be measuring the capacitance of lots of points at different applied voltage, how will a changing n effect the slope in our relationship? What does the slope mean between two points?

Question 11 *What might cause a change in n throughout the silicon? Where would you expect the largest change?*

4.4 Defects and deep-level traps

At the heart of semiconductor physics is the idea that one can alter the electrical properties of a semiconductor by the addition of impurities and defects in the semiconductor lattice. This idea is the basis behind the doping of a semiconductor as these impurities or defects can create donor or acceptor sites in the semiconductor lattice. Having said this, it is not always desirable to have additional donor or acceptor in the semiconductor lattice as they may change the electrical properties of a fabricated device in an undesired or unpredictable manner. It is for this important reason that the task of electrical defect analysis has been undertaken.

Question 12 *Here you should comprehensive explain the effects of different impurities and defects in semiconductors. You should make sure you cover the topics of: lattice vacancies and interstitial atoms, electrically active dopants and defects, shallow and deep charge traps, and the capture cross-section* (σ) *of a dopant.*

The silicon wafers used to create the diode in this experiment have undergone ion implantation, the species of ion is either phosphorous or hydrogen depending on which sample is used for your individual experiment. It is well understood that this process introduces many defects into the substrate [4, 5, 6]. This damage is initially of the form of interstitial atoms which have been knocked off lattice sites leaving behind a vacancy [6] this damage is known a Frenkel defects. At room temperature the

vacancy and interstitial can migrate and separate without recombination. This occurs for 4-10% of the Frenkel defects created during ion implantation [6]. These defects can go on to cluster together or form stable defect complexes with impurities. About 10-25% of the Frenkel defects that survive recombination form a di-vacancy cluster [7] which is a characteristic defect for ion implanted silicon [4, 8], or bond with impurities to form other stable clusters.

The defects of prime importance for electrical devices are those that are electrically active, meaning that they can trap charge carriers in the device. Electrically active traps are basically unoccupied states in the band-gap of a semiconductor that trap charge carriers. This occurs mostly due to defects having dangling bonds, or unpaired electrons, and the traps exist with in the band-gap of the silicon at some energy level depending on the structure of the defect. Defects can be characterised by their position in the band-gap of a semiconductor relative to the conduction band (trap energy, E_t), and their crosssection for trapping (capture cross-section, σ). The values reported for electron traps in ion-implanted silicon are found in table 1. This table shows the average value for the trap energies found in the literature. Absolute errors in the last digit are shown in parentheses, no error is quoted for occasions where there is only one reference for the defect, nor is the error quoted for capture cross-sections that have an error that is over an order of magnitude difference. V indicates vacancy, other letters indicate elements, subscripts 2 & 3 indicate the number of vacancies in a cluster and superscripts indicate the charge state of the trap. Other subscripts indicate either substitutional or interstitial atomic positions in the lattice. The silicon wafer implanted with phosphorus was implanted at an energy of 450 keV at a dose of 1×10^9 P/cm² while the samples implanted with hydrogen were implanted at an energy of 70 keV at a dose of 1×10^{10} H/cm².

It is also possible other samples will be used, depending on availability/interest/someone wanting to analyse their own data and involving you in the exciting world of research physics.

Ask your demonstrator which sample and implantation you will be using for this experiment.

The local doping concentration will change with defect concentration. However to find out exactly what type of defects are causing this altered doping concentration we will need to employ DLTS.

4.5 Deep level transient spectroscopy (DLTS)

In DLTS a pulsing bias is applied to the diode, this fills and drains the charge traps that exist within the semiconductor, figure 1 shows an example of such a pulse cycle. Figure 2 shows the effect of changing the bias on the trap population in the sample. Figure 2a) shows the effect of applying the V_{pulse} bias to the diode, here the traps begin to fill. We call this electron trapping. Figure 2 b) shows the effect of applying a lower offset bias V_{offset} (or reverse bias), here trapped electron in the 'increased ω ' region will be released, we call this electron emission. The region defined by ω during the V_{pulse} bias and V_{offset} bias is known as the measurement window, it will be important to select appropriate values of V_{pulse} and V_{offset} so that the measurement region includes all the deep level traps in your diode. The total concentration of traps in the measurement region is referred to as N_T while the concentration of full traps (or trapped electrons) is referred to as n_T .

Question 13 Write an expression for the concentration of filled traps within the measurement region straight after the V_{pulse} bias has been applied (n_T (t = 0)). This expression should be written in terms of the time V_{pulse} bias is applied t_p , the rate the electrons are trapped r_c , and a total trap concentration N_T . Assume all the traps are empty before the pulse, $n_T(t=0)=0$.

Defect type	Trap Energy (E_c-E_t)	Capture cross-section
	[ev]	$[\mathrm{cm}^2]$
H-related	0.10	-
H-related	0.13	-
НС	0.15(1)	-
VO	0.17(1)	$7(4)x10^{-15}$
C_sC_i	0.17(0)	$8 x 10^{-18}$
V-related	0.19(1)	3.5×10^{-17}
V_{2}^{2-}	0.22(1)	$5(4)x10^{-16}$
V_2O	0.27	-
V ₃ O	0.30	-
VO-H, other H–related	0.30(2)	$3x10^{-15}$
V-related	0.35(1)	5×10^{-16}
H-related	0.39(1)	1×10^{-17}
H-related	0.41(1)	-
VP _s	0.42(2)	$5(3)x10^{-15}$
$ V_2^- $	0.42(1)	$5(3)x10^{-15}$
H-related	0.45(1)	$1 x 10^{-17}$
V ₂ -related	0.47	_
H-related	0.50(1)	-
Unknown	0.59	_

Table 1: Defect characteristics as published in the literature for defects relevant to ion-implanted n-type silicon.

Hint: The trapping rate is assumed to be exponential and don't forget that there is a finite number of traps to be filled.

The concentration of trapped electrons in the measurement region (n_T) exponentially decreases and is effected by the emission rate (e_n) of an individual trap:

$$n_T(t) = n_T(t=0)e^{-e_n t}$$
(2)

Assuming $n_T \approx N_T$ this becomes:

$$n_T(t) = N_T(t=0)e^{-e_n t}$$
(3)

 e_n depends on temperature T, the trap energy level E_T and the capture cross section of the trap σ_c :

$$e_n = \gamma_n \,\sigma_c \,T^2 e^{-\left(\frac{E_c - E_T}{k_B T}\right)} \tag{4}$$

Where γ_n is a set of constants given by:

$$\gamma_n = 2\sqrt{3}M_c (2\pi)^{\frac{3}{2}} k_b^2 m^* h^{-3} \tag{5}$$

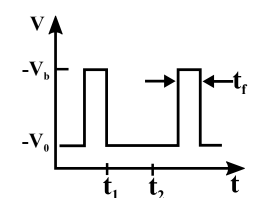


Figure 1: Required pulse cycle for DLTS [3].

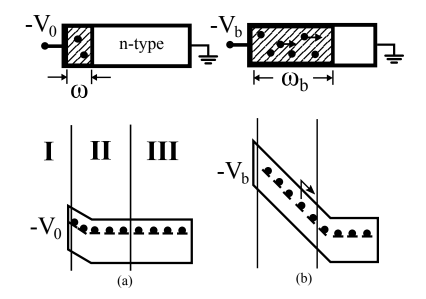


Figure 2: a) Deep level traps full, occurs when V_{pulse} bias is applied. b) Deep level taps emptying, occurs straight after the Voffset is reapplied.

Where M_c is the number of minima in the conduction band (6 for silicon) and m^{*} is the effective electron mass in the conduction band [9]. Since the trap energy level and the capture cross section are different for different defects, the emission rate en will be different for each defect. It is important to note that as the electrons are emitted from traps in the measurement region, during the time the V_{offset} bias is applied, the depletion width ω will be affected and hence the diode capacitance will also be affected.

Figure 3 shows how the capacitance of the Schottky diode changes over time as a result of the traps emptying. This capacitance transient can be expressed as:

$$C = C_0 \sqrt{1 - \frac{n_T(t)}{n}} \tag{6}$$

For the case where the doping concentration is n is much greater than the trap concentration $(nT_{ii} n)$ we can say:

$$C \approx C_0 \left(1 - \frac{n_T(t)}{2n}\right) \tag{7}$$

With DLTS we monitor the change in capacitance over some time interval (t1, t2). The change

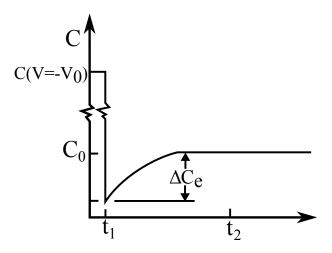


Figure 3: The time variation of capacitance as traps empty [3].

in capacitance over this interval (from Eq. 7) is hence:

$$\Delta C = C(t_1) - C(t_2) = C_0\left[\frac{N_T}{2n}\right] \left[e^{-e_n(T)t_2} - e^{-e_n(T)t_1}\right]$$
(8)

This change is capacitance is referred to loosely as the DLTS signal. Figure 4 shows how the DLTS signal changes as a function of temperature when only a single trap is present, this occurs due to the temperature dependence of the emission rate. This change in DLTS signal with temperature forms the DLTS spectrum.

Traps with different energy levels in the band gap will produce a peak at different temperatures. Note that in equation 8., the DLTS signal is proportional to the concentration of defects present N_T , and can be expressed as:

$$\frac{\Delta C}{C_0}\Big|_{max} = \frac{n_T}{2n} \frac{1-r}{r\frac{r}{r-1}}$$
(9)

Where $r = \frac{t_2}{t_1}$. We can then write for the trap concentration:

$$n_T = 2 \frac{\delta C}{C_0} \Big|_{max} n \frac{r^{\frac{r}{r-1}}}{1-r}$$
(10)

Question 14 Differentiate equation 8 with respect to T and show that the emission rate at the temperature that the DLTS signal is greatest is given by the following equation:

$$e_n(T_{peak}) = \frac{ln(\frac{t_2}{t_1})}{t_2 - t_1} \tag{11}$$

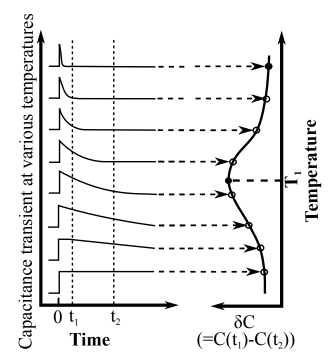


Figure 4: The temperature dependence of the DLTS signal [3]. Here the transient capacitance is in negative units (are they? Why? Make sure you understand).

Hint: you will NOT NEED to explicitly differentiate e_n *, just carry it through.*

Question 15 Derive equation 9 by subbing equation 11 into equation 8.

Equation 11 shows that the measurement times t_1 and t_2 sets what is known as a *rate window* for the emission rate. It can be shown that different rate windows cause the peak in the DLTS signal to shift in temperature. By taking DLTS spectra with various rate windows we may obtain a range of values for the peak temperature associated with each emission rate and hence each defect. Using these values, and noting that at the peak temperature the emission rate is given by your answer to equation 11, it is possible to calculate both the trap energy level and the capture cross-section.

Since the temperature at which the DLTS signal peaks changes with the rate window, we can generate a plot of the temperature dependant rate against $\frac{1}{T}$ using equation 4, this is known as an **Arhhenius** plot

Question 16 Re-arrange Equation 4 to obtain $ln(\frac{e_n}{T^2})$ in terms of $\frac{1}{T}$. How can this be used to calculate the trap energy and capture cross section? Check with your demonstrator as this will be important later in the DLTS analysis.

Question 17 One last quick question - do we mean two different things when we say "trap" rather than "defect"? i.e.: how do we refer to defects that are not electrically active?

You might find it useful to write a list of relevant equations on a single page in your notebook with your notes as to their relevance. This will help later when hunting for relationships between the measured values.

5 Experimental Work

The SULA DLTS system and its associated electronics is a joint MARC group Part 3 laboratory apparatus and is thus also used for research. The research of the MARC group (and perhaps even one of your demonstrators) depends on the correct functionality of the equipment so it is especially important to treat it with respect.

Week 1

- SRIM ion implantation simulation.
- C-V measurement on a n-Si Schottky diode at room temp and 79K. (And C-T on the way back up to room temperature)
- Analyse data to determine active dopant concentration and barrier height etc.

Week 2

- Do DLTS measurement on a ion implanted or irradiated Si sample.
- Analyse data to determine trap energy and capture cross section of defects.
- Identify defects created in the implantation process.

5.1 The racks of electronics

The box at the top of the rack is the Lakeshore **temperature controller**. This is primarily under the remote control of the PC and software. Do not play with it, if there is an error message or other problem consult your demonstrator.

Next comes the digital **CRO** for monitoring the applied bias on the sample.

Then there are the **SULA DLTS electronics**. Further information on these units can be found in the SULA DLTS user manual.

Underneath this is a rack of BNC connectors. Computers dont come with BNC sockets, so this rack is purely to interface the PC with all the electronics.

The last tray at the bottom is the variac for the diaphragm pump (for adjusting the pump speed) and the thermocouple vacuum gauge for the roughing line pressure.

5.2 The cryostat and sample holder

Ask your demonstrator to load your diode. The sample mounting setup is shown in figure 5. The InGa is a eutectic which is liquid at room temperature. We use it because it forms an Ohmic contact with the sample (unlike the gold contact on top, which forms a Schottky contact). The sapphire plate is an electrical insulator to reduce the stray capacitance created around the diode but still allowing good thermal contact to the copper stage.

Question 18 What property of In or Ga makes InGa a suitable Ohmic contact? What about the Au to create a Schottky contact? Hint: It has nothing to do with resistivity/conductivity.

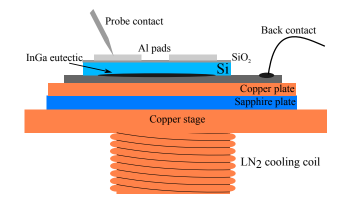


Figure 5: Sample holder schematic

Question 19 There are two temperature sensors in the DLTS setup. Find both of them, label them on a diagram and explain why two are used.

Here we need accurate control of the temperature, for this reason the stage is fitted with a heater and two thermocouples (electric thermometers). Thermocouple A is used for temperature control and is situated at the base of the copper stage in the chamber near the heater, Thermocouple B is situated at the top of the copper stage near the sample and is taken as the measured sample temperature.

The cooling is provided by drawing liquid nitrogen up the tube at the bottom and through a tube around the base of (and in intimate thermal contact with) the cold head and out the side, through the transparent plastic tube. This is achieved by pumping on the transparent plastic tube with a diaphragm pump (the same as used at home in a fish tanks). This diaphragm/N2 pump is controlled by a variable voltage source. 70 V is good for cooling down to 77 K (takes about 15 mins); 20 V will keep it there.

The sample chamber is also evacuated using a rotary roughing pump. It is kept under vacuum for a couple of reasons. It keeps out any moisture, oils and dust in the air which may contaminate the sample and sample chamber. Most importantly it is needed for thermal isolation of the cold head so that lower temperatures can be maintained.

5.3 SRIM simulation

SRIM [10] is a software package written to tell us the Stopping power and Ranges of Ions in Matter. Most of the information about SRIM can be found at www.srim.org, as can the software packages themselves. You will find a hard copy of the SRIM manual in the lab, which contains an excellent introduction to the software. You should read enough of it to feel comfortable with what you are trying to use SRIM for.

We will use SRIM to determine the penetration depth of the implanted ions and the vacancy, or damage, profile created in this process. This is important because you will need to know how to probe for defects later in this practical.

Using SRIM, run a simulation that represents the same implantation parameters as that of the diodes used in the practical. Importantly you will need to set the following parameters:

- Ion element type and energy
- Target element and width (it is important that the width is not too large as there is a limit to the resolution)
- The total number of ions (not too small... try about 100,000 ions)

From this calculation you will be able to find the depth profile over which the vacancies were created when the ions were implanted in the wafer. This vacancy profile can then be used as an approximate for the depth that the deep level traps exist.

5.4 Performing the C-V measurement

Note that, throughout all of the experiment, there may be things that are different from these notes. In these cases, follow the instructions of your demonstrator. Various upgrades are constantly being performed or work-arounds to equipment malfunctions are implemented, so these notes can't always be as accurate as we'd like.

The C-V measurement will allow us to determine the barrier height, depletion width and active dopant density of the sample.

The bias is applied to the top diode contact of the sample via the output BNC connection on the [Pulse generator] unit and the capacitance is read from the back contact which is connected to the [In] of the [Capacitance meter] unit.

Electronics setup

Typically the output of the pulse generator is triggered to the CRO so that the applied bias can be observed. The applied DC bias can be controlled in two ways, with the [Offset] knob and via an input into the [Ext. Bias] connection. The software will use the digital to analogue converter (DAC) in the PC to apply a DC analogue bias to the sample via this input from [DACOUT0] of the BNC adaptor.

Running the experiment

A voltage interval of around 0.1 V will be needed for a nice resolution data set for analysis. Set the [Initial temp] to something around room temp, turn on the diaphragm pump and set the voltage to around 10 V.

The following is a 'general' overview of what you should do. The software controlling the DLTS electronics has recently received an update so more specific instructions will be required from your demonstrator. That is, **don't worry that the following step-by-step guide doesn't match up exactly with the GUI on the computer**.

Step-by-step:

- 1. Flip the [Experiment] type to C-V
- 2. Click [Idle at initial temp] to set the temperature controller.
- 3. Set [Initial bias] and [Final bias] to 0V and -8V (or something like that)

- 4. The [Capacitance range] should be set to be set to 100 pF
- 5. Set the [Diode area]
- 6. Click [Run experiment] to start the measurement.Note: the measurement will not start until the temperature of thermocouple A is at the [Initial temp].

The screen should (eventually) show a trace of the capacitance with voltage. Check with your demonstrator that it is going in the right direction and that it looks reasonable.

5.5 Analysing the C-V data

You will need to re-examine your answers to earlier questions in the capacitance and depletion width section. You will get the best results if you fit the data over biases close to zero. This is because you will find that the active dopant concentration is not uniform over depth. (Why, again?) Furthermore you should not calculate values for each data point and it is much more accurate to perform a linear fit.

Question 20 To determine the built-in voltage V_{bi} you should only fit the data over biases close to zero. This is because the free carrier concentration (as you will see) is not uniform over depth and there is more leakage current at larger biases, making the capacitance measurements less accurate. Do not calculate V_{bi} for each data point; this is not an accurate method. Compare your answer to any answer the software may have given you.

Question 21 Calculate the barrier height which can be expressed as:

$$\phi_b = V_{bi} + E_f \tag{12}$$

Where E_f is the position of the Fermi level relative to the conduction band and is given by:

$$E_f = \frac{k_b T}{q} Log_e[\frac{n_c}{n}] \tag{13}$$

 n_c is the Density of states in the conduction band of silicon:

$$n_c = 2 \times 10^{-6} (2\pi m^* \frac{k_b T}{h^2})^{\frac{3}{2}}$$
(14)

Note: Your answer should be expressed in eV and greater than the built–in voltage but less than the band gap of silicon.

Question 22 Now that you have the built-in voltage and the barrier height, sketch a labeled band diagram for the diode showing the values for V_{bi} and ϕ_b

Question 23 Calculate the active dopant concentration n as a function of depth. You will need to re-examine your answers to questions in the 'capacitance and depletion width' section and convert the capacitance to a corresponding depletion width (depth). You can use Excel or Genplot for data analysis, this may take a bit of time. You will need these results for DLTS section.

Question 24 We have simulated the ion implantation process in SRIM which suggests there is a region in the silicon which has been damaged. You will have also created a profile of the active doping concentration, what important conclusion can you draw from these profiles?

Question 25 What voltages will you use for your DLTS measurement window (V_{offset} and V_pulse) and why?

MAKE SURE you understand how exactly the C-V scan is relevant to your later DLTS scan. Make a table with relevant information if necessary.

5.6 DLTS measurement

Electronics setup

The electronics can now be setup for a DLTS measurement. The first thing to do is set the [Off-set](Voffset) on the [Pulse generator] to place the diode under reverse bias. This should be set so that the depletion region is at the end of range you decided in the previous section. Monitor the voltage with the [V] setting on the [Capacitance meter]. Since the bias output of the pulse generator should also be connected to the CRO you should also be able to observe this on the CRO. To ensure proper operation check the leakage current by pressing the [i] setting on the [Capacitance meter]. The reading is in A, it should not be more than 5 μ A.

Next turn on the pulse generator with the [On/Off] toggle switch. Press the [-] button on the [Capacitance meter]. The display now indicates the voltage at which the bias is pulsed to. Set the pulse [Amplitude](Vpulse) on the pulse generator so that the depletion width will be at the start of the region of uniform free carriers. You should be able to now see the pulse on the CRO. If not, ensure that the CRO is set to the DC mode and external trigger. If the trigger is set to normal you may need to adjust the trigger level.

The pulse [Width] should be set to fill as many traps as possible. The pulse width affects how many of the defects will be filled during the pulse since they have a characteristic capture crosssection (and hence capture rate). For the defects in ion implanted silicon this needs to be about 10-20 ms. The electronics does not monitor this and the CRO needs to be used.

The [Initial delay] of the correlators need to be set to examine the capacitance transient over the correct rate windows. There are four correlators so we can simultaneously look at four different rate windows. The correlators work such that:

- $t_1 = 2.7x$ [Initial delay], and
- $t_2 = 6.9x$ [Initial delay],

hence r = 2.56, where as mentioned earlier $r = t_2/t_1$.

The [Initial Delay] settings make take an aborted scan or two to get right. ASK YOUR DEMON-STRATOR. Always have the first correlator as the longest time, decreasing in time from there. e.g.:

Correlator 1: 10 ms Correlator 2: 2 ms Correlator 3: 0.5 ms Correlator 4: 0.1 ms The [TC] value of the correlators should be set to 10 seconds. The [Period] of pulse repetition needs to be carefully set to ensure the electronics have enough time to recover, I recommend 200 ms. The [Pre-Amp] should be set to 100. The pulse generator is now setup for DLTS and the rest of the measurement electronics can be set up.

Software setup

Change the front panel [Experiment] to [DLTS] and enter all the hardware settings into the window. The [Initial Temp.] should be set to 78 K and the [Final Temp.] to room temperature. The mode should be on [Step]. The stray capacitance is irrelevant and the [Offset] should be set to [use current valses] and the should be zero. Once you have completed this you can click [Run experiment] and the scan will take approximately 2-3 (or longer!) hours to complete.

5.7 Perfoming a C-T measurement

You will need to know C_o at various temperatures to determine the defect concentrations (Eq. 10).

For this you need a C-T scan. You can choose when you'd like to perform it. (Personally I like to do it on the way up from a cold scan, so maybe after the low-temperature CV, but you'll have to decide quickly on your offset voltage!)

You need to know what V_{offset} you are using for your DLTS before you perform the C-T, as the sample needs to be held at this offset (with the pulse off) during the C-T scan.

6 Analysing the DLTS data

Okay, at this point you've had your hand held a little bit. This is third year, and next year won't get easier, so you need to prepare. To that end, though this section humbly titled 'Analysing the DLTS data' looks short, it will take a long time, and made drive you around in both physical and mathematical circles. Keep an eye on the time and take regular breaks to review your work and make sure you're both accurate and heading down the right path...

Oh, and INCLUDE ERRORS.

Question 26 Calculate the defect concentration for each of the defect peaks observed.

Question 27 Use Excel to find the location of each defect peak in terms of temperature. Use this information to find the trap energy E_T and capture cross section σ of each defect. Look to earlier exercises for help here, dont forget to also determine the errors in your energy levels. (This is the crux of the entire experiment.)

Question 28 Which defect do you think your peaks correspond to?

In fact, DLTS offers rather poor energy resolution and hence identifying defects from a single DLTS studies is not that accurate. A more detailed study would involve examining how the observed defect peaks change as the sample is subjected to thermal annealing. In your case however please refer to the literature. There should be several papers provided to you in the lab somewhere. How do your results compare with the literature?

7 Overview

Okay, so just to summarise all of that neatly and give you a page to refer back to, this is the state of affairs.

- 1. You'll have thought about the theory and discussed with your demonstrator any questions.
- 2. An SRIM implantation simulation is performed to determine approximate depth of implanted ions.
- 3. Ion-implanted, Schottky-diode semiconducting sample goes in to sample chamber with appropriate connections.
- 4. A capacitance-voltage (C-V) scan is taken at room temperature and liquid nitrogen temperature. From these C-V scans, you are able to determine:
 - a) The depletion width of the sample with applied voltage (and thus the voltage ranges for DLTS)
 - b) The doping concentration of the sample with applied voltage
 - c) The changes to the doping of the sample near the surface
 - d) The average doping concentration, n, of the sample, used in equations for DLTS
 - e) The built-in voltage V_{bi} and barrier height ϕ_b of the junction
- 5. A capacitance-temperature (C-T) scan is taken on returning to room temperature from liquid nitrogen temperature. This is C_0 .
- 6. The sample is again cooled and a DLTS scan is taken with appropriate voltage, correlator and TC settings. From the DLTS scan, you are able to determine:
 - a) The number of different types of strong defects present in your sample
 - b) The temperatures at which different defects are most active
 - c) The amount of *each species* of defect that are electrically active in your sample
 - d) (Using an Arhennius plot): The location in the bandgap, E_c-E_t , of each of the defect species
 - e) (Using an Arhennius plot): The capture cross section, σ , of each of the defect species

8 Important Numbers

Diameter of diode = 780 μ m with an uncertainty of 20 μ m				
$q = 1.60218 \times 10^{-19}$	Electron charge [C]			
$\epsilon = \epsilon_0 \times \epsilon_s$	Permittivity of sample			
$\epsilon_0 = 8.85418 \times 10^{-14}$	Permittivity of free space [F/cm]			
$\epsilon_s = 11.9$	Dielectric factor for Si			
$k_b = 1.38066 \times 10^{-23}$	Boltzmann's constant [J/K]			
$h = 6.62617 \times 10^{-34}$	Planck's constant [J.s]			
$m^* = \mathbf{m}_{e0} \times \mathbf{m}_e$	Effective electron mass [kg]			
$m_{e0} = 9.1095 \times 10^{-31}$	Electron rest mass [kg]			
$m_e = (0.98 \times 0.19^2)^{1/3}$	Effective electron mass factor in Si at 300K			
Calculating $\mathbf{r} = 2.56$	$r = t_2/t_1$ where $t_2 = 6.9 \times [Initial Delay]$ and $t_1 = 2.7 \times [Initial Delay]$			

References

- [1] D. V. Lang. Deep-level transient spectroscopy: A new method to characterize traps in semiconductors. *Journal of Applied Physics*, 45(7):3023–3032, 1974.
- [2] Ben G Streetman and Sanjay Kumar Banerjee. *Solid state electronic devices; 4th ed.* Prentice-Hall, Englewood Cliffs, NJ, 1995.
- [3] D.K. Schroder. *Semiconductor material and device characterization*. Wiley-International Publications, USA, New York, 2nd edition, 1998.
- [4] B. G. Svensson, B. Mohadjeri, A. Hallén, J. H. Svensson, and J. W. Corbett. Divacancy acceptor levels in ion-irradiated silicon. *Physical Review B*, 43(3):2292–2298, Jan 1991.
- [5] P. Hazdra, V. Haslař, and M. Bartoš. The influence of implantation temperature and subsequent annealing on residual implantation defects in silicon. *Nuclear Instruments and Methods in Physics Research Section B: Beam Interactions with Materials and Atoms*, 55(1-4):637 – 641, 1991.
- [6] S. Coffa and F. Priolo. Electrical properties of ion implanted and electron irradiated c-si. page 746, 1999. In Robert Hull et al., editors *Properties of crystalline silicon*.
- [7] B. G. Svensson, C. Jagadish, A. Hallén, and J. Lalita. Generation of vacancy-type point defects in single collision cascades during swift-ion bombardment of silicon. *Physical Review B*, 55(16):10498–10507, Apr 1997.
- [8] A. Hallen, B. U. R. Sundqvist, Z. Paska, B. G. Svensson, M. Rosling, and J. Tirn. Deep level transient spectroscopy analysis of fast ion tracks in silicon. *Journal of Applied Physics*, 67(3), 1990.
- [9] P. Blood and J. W. Orton. *The Electrical Characterization of Semiconductors: Majority Carriers and Electron States*. Academic Press, 1992.
- [10] J. F. Ziegler. Srim-2003. Nucl. Instrum. Methods Phys. Res., Sect. B, 219:1027, 2004.