6. Materials and Processes for Silicon Technology

6.1 Si Oxides and LOCOS Process

6.1.1 Si Dioxide

The Importance of Silicon Dioxide

Silicon would not be the "miracle material" without its oxide, <u>SiO2</u>, also known as ordinary <u>quartz</u> in its bulk form, or as **rock crystal**if you find it in single crystal form (and relatively pure) somewhere out there in the mountains.

- Not only the properties of Si especially the achievable crystalline perfection in combination with the bandgap and easy doping but also the properties of SiO₂ are pretty much as one would have wished them to be for making integrated circuits and other devices.
- From the beginning of integrated circuit technology say 1970 to the end of the millennium, SiO₂ was a key material that was used for many different purposes. Only now (around 2004), industry has started to replace it for some applications with other, highly "specialized" dielectrics.
- What is so special about SiO₂?
 - First of all, it comes in many forms: There are several *allotropes* (meaning different crystal types) of SiO₂ crystals; the most common (stable at room temperature and ambient pressure) is "low quartz" or α-quartz, the quartz crystals found everywhere. But SiO₂ is also rather stable and easy to make in amorphous form. It is amorphous SiO₂- homogeneous and isotropic that is used in integrated circuit technology. The link provides the <u>phase</u> diagram of SiO₂ and lists some of its allotropes.
 - SiO₂ has excellent dielectric properties. Its dielectric constant ∈_r is about 3.7 3.9 (depending somewhat on its structure). This was a value large enough to allow decent capacitances if SiO₂ is used as capacitor dielectric, but small enough so that the time constant *R* · *C* (which describes the time delay in wires having a resistance *R* and being insulated by SiO₂, and thus endowed with a parasitic capacitance *C* that scales with ∈_r) does not limit the maximum frequency of the devices. It is here that successors for SiO₂ with larger or smaller ∈_r values are needed to make the most advanced devices (which will hit the market around 2002).
 - It is among the best insulators known and has one of the highest break-through field strengths of all materials investigated so far (it can be as high as 15 MV/cm for very thin layers; compare that with the values given for normal "bulk" dielectrics).
 - The electrical properties of the Si SiO₂ interface are excellent. This means that the interface has a very low density of energy states (akin to surface states) in the bandgap and thus does neither provide recombination centers nor introduce fixed charges.
 - SiO₂ is relatively easy to make with several quite different methods, thus allowing a large degree of process leeway.
 - It is also relatively easy to structure, i.e. unwanted SiO₂ can be removed selectively to Si (and some other materials) without many problems.
 - It is very stable and chemically inert. It essentially protects the Si or the whole integrated circuit from rapid deterioration in a chemically hostile environment (provided simply by humid air which will attack most "unprotected" materials).

What are the uses of **SiO₂**? Above, some of them were already mentioned, here we just list them a bit more systematically. If you do not quite understand some of the uses - do not worry, we will come back to it.

Gate oxides: As we have <u>seen before</u>, we need a thin dielectric material to insulate the gate from the channel area. We want the channel to open at low threshold voltages and this requires large dielectric constants and especially no charges in the dielectric or at the two interfaces. Of course, we always need high break through field strength, too. No dielectric so far can match the properties of **SiO₂** in total.

Dielectrics in integrated capacitors. Capacitors with high capacitance values at small dimensions are needed for socalled **dynamic random access memories** (*DRAM*), one of the most important integrated circuits (in terms of volume production). You want something like **30 fF** (femtofarad) on an area of **0.25 μm²**. The same issues as above are crucial, except that a large dielectric constant is even more important. While **SiO₂** was the material of choice for many **DRAM** generations (from the **16 kbit** **DRAM** to the **1 Mbit DRAM**), starting with the **4 Mbit** generation in about **1990**, it was replaced by a triple layer of $SiO_2 - Si_3N_4 - SiO_2$, universally known as "<u>ONO</u>" (short for oxide - nitride - oxide); a compromise that takes not only advantage of the relatively large dielectric constant of silicon nitride (around **7.5**) while still keeping the superior quality of the Si - SiO₂ interface, but has a few added benefits - at added costs, of course.

- Insulation: Some insulating material is needed between the transistor in the Si as well as between the many layers of wiring on the chip; cf. the many pictures in chapter five, starting with the one accessible via the link. SiO₂ was (and still is) the material of choice. However, here we would like to have a material with a *small* dielectric constant, ideally 1, minimizing the parasitic capacitance between wiring and SiO₂ may have to be replaced with a different kind of dielectric around 2003.
- Stress relieve layer: SiO₂ becomes "viscous" at high temperatures - it is a glass, after all. While it is a small effect, it is large enough to absorb the stress that would develop between unwielding materials, e.g. Si₃N₄ on Si, if it is used as a "buffer oxide", i.e. as a thin intermediary layer.
- Masking: Areas on the Si which are not be exposed to dopant diffusion or ion implantation must be protected by something impenetrable and that also can be removed easily after "use". It's SiO₂, of course, in many cases,
- "Screen oxides" provide one example of so-called sacrificial layers which have no direct function and are disposed off after use. A screen oxide is a thin layer of SiO₂ which stops the low energy *debris* that comes along with the high-energy ion beam consisting, e.g., of metal ions that some stray ions from the main beam banged off the walls of the machine. All these (highly detrimental) metal and carbon ions get stuck in the screen oxide (which will be removed after the implantation) and never enter the Si. In addition, the screen oxide scatters the main ion beam a little and thus prevents "*channeling*", i.e. deep penetration of the ions if the beam happens to be aligned with a major crystallographic direction.
- Passivation: After the chip is finished, it has to be protected from the environment and all bare surfaces need to be electrically passivates - its done with SiO₂ (or a mixture of oxide and nitride).
- Enough reasons for looking at the oxide generation process a little more closely? If you think not well there are more uses, just consult the list of processes for a **16 Mbit DRAM**: You need **SiO**₂ about **20** times!

How is **SiO₂** made - in thin layers, of course? There are essentially *three* quite distinct processes with many variations:

- Thermal oxidation. This means that a solid state reaction (Si + O₂ ⇒ SiO₂) is used: Just expose Si to O₂ at sufficiently high temperatures and an oxide will grow to a thickness determined by the temperature and the oxidation time.
- CVD oxide deposition. In complete analogy to the production of poly-Si by a CVD (= chemical vapor depositions) process, we can also produce SiO₂ by taking the right gases and deposition conditions.
- Spin-on glass (SOG). Here a kind of polymeric suspension of SiO₂ dissolved in a suitable solvent is dropped on a rapidly spinning wafer. The centrifugal forces spread a thin viscous layer of the stuff on the wafer surface which upon heating solidifies into (not extremely good) SiO₂. Its not unlike the stuff that was called "water glass" or "liquid glass" and that your grandma used to conserve eggs in it.

There is one more method that is, however, rarely used - and never for mass production: Anodic oxidation.

- Anodic oxidation uses a current impressed on a Si electrolyte junction that leads to an oxidation reaction. While easy to understand in principle, it is not very well understood in practice and an object of growing basic research interest.
- In the module **2.5.1** "Porous semiconductors more information to this point can be found.

- Gate oxide for Transistors
- Dielectric in Capacitors
- Insulation
- Stress relieve layer
- Masking layer
- Screen oxide during
 Implantation
- Passivation

Thermal Oxidation

In this paragraph we will restrict ourselves to **thermal oxidation**. It was (and to a large extent still is) one of the key processes for making integrated circuits. While it may be used for "secondary" purposes like protecting the bare **Si** surface during some critical process (remember the "screen oxide" from above?), its major use is in three areas:

- **Gate oxide** (often known as "*GOX*")
- Capacitor dielectric, either as "simple" oxide or as the "bread" in an "ONO" (= oxide-nitride-oxide sandwich)
- Field oxide (FOX) the lateral insulation between transistors.

We can use a picture from <u>chapter 5.1.4</u> to illustrate **GOX** and **FOX**; the <u>capacitor dielectric</u> can also be found in this chapter.



We must realize, however, that those drawing are *never to scale*. The gate oxide is only around **10 nm** thick (actually, it "just" (**2007**) petered out at **1.2 nm** accoding to Intel and is now replaced by a thicked **HfO**₂), whereas the field oxide (and the insulating oxide) is in the order of **500 nm**. What it looks like at atomic resolution in an electron microscope is shown in <u>this link</u>.

There are essentially two ways to do a thermal oxidation.

"Dry oxidation", using the reaction

- This is the standard reaction for thin oxides. Oxide growth is rather slow and easily controllable.
- To give an example: Growing 700 nm oxide at 1000 °C would take about 60 hr far too long for efficient processing. But 7nm take only about 15 min - and that is now too short for precise control; you would want to lower the temperature.
- "Wet oxidation", using the reaction

Si + 2 H₂O ⇒ SiO₂ + 2 H₂ (800 °C - 1100 °C)

- The growth kinetics are about **10x** faster than for dry oxidations; this is the process used for the thick field oxides.
- Growing **700 nm** oxide at **1000 °C** now takes about **1.5 hr** still pretty long but tolerable. Thin oxides are never made this way.

In both cases the oxygen (in the form of **O**, **O**₂, **OH**⁻, whatever,...) has to diffuse through the oxide already formed to reach the **Si** - **SiO**₂ interface where the actual reaction takes place.

- This becomes more difficult for thick oxides, the reaction after some time of oxide formation is always diffusion limited. The thickness dox of the growing oxide in this case follow a general "square root" law, i.e. it is proportional to the diffusion length L=(Dt)^{1/2} (D=diffusion coefficient of the oxygen carrying species in SiO₂; t= time).
- We thus have a general relation of the form .



For short times or thin oxide thicknesses (about < 30 nm), a linear law is found

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d_{\text{thin-ox}} = \text{const.} \cdot t
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In this case the limiting factor is the rate at which O can be incorporated into the Si - SiO₂ interface.

This kind of behavior - linear growth switching to square root growth - can be modelled quite nicely by a not too complicated theory known as the **Deal-Grove model**. Some results of experiments and modeling are shown below.



The left diagram shows dry, the right hand one wet oxidation. The solid curves were calculated with the Deal-Grove model after parameter adjustment, the circles indicate experimental results. *The theory seems to be pretty good !*.

However, for very thin oxides - and those are the ones needed or **GOX** or capacitors - things are even more complicated as shown below.



- The red points are data points from an experiment (at an unusually low temperature); the blue curve is the best Deal-Grove fit under the (not justified) assumption that at *t=0* an oxide with a thickness of about 20 nm was already present.
- The Deal-Grove model is clearly inadequate for the technologically important case of very thin oxides and experimentally an exponential law is found for the dependence of the oxide thickness on time for very short times
- Moreover, the detailed kinetics of oxide growth are influenced by many other factors, e.g. the crystallographic orientation of the Si surface, the mechanical stress in the oxide (which in turn depends on many process variables), the substrate doping, and the initial condition of the Si surface.

And this is only the thickness! If we consider the properties of the oxide, e.g. the amount of fixed charge or interface charge, its etching rate, or - most difficult to **assess** - how long it will last when the device is used, things become most complicated. An oxide with a nominal thickness d_{ox} can be produced in many ways: dry or wet oxidation, high temperatures and short oxidation times or the other way around - its properties, however, can be very different.

Learning about microelectronic processes involves very little Math; and "theory" is needed only at an elementary to medium level! But this does *not* make the issue trivial - quite the contrary. If you would have a theory - however complicated - that predicts all oxide properties as a function of all variables, process development would be easy. But presently, even involved theories are mostly far too simple to come even close to what is needed. On an advanced level of real process development, it is the interplay of a solid foundation in materials science, lots of experience, usage of mathematical models as far as they exist, and finally some luck or "feeling" for the job, that will carry the day.

So do not consider microelectronic processes "simple" because you do not find lots of differential equations here. There are few enterprises more challenging for a materials scientist then to develop key processes for the next chip generation! How is a thermal oxidation done in real life? Always inside an oxidation furnace in a **batch process**; i.e. many wafers (usually **100**) are processed at the same time.

Oxidation furnaces are complicated affairs, the sketch below does not do justice to the intricacies involved (nowadays they are usually no longer horizontal as shown below, but vertical). For some pictures of real furnaces use the <u>link</u>.



- First of all, temperature, gas flow etc., needs not only to be very constant but precisely adjustable to your needs. Generally, you do not load the furnace at the process temperature but at some lower temperature to avoid thermal shock of the wafers (inducing temperature gradients, leading to mechanical stress gradients, leading to plastic deformation, leading to the generation of dislocations, leading to wafers you have to throw away). After loading, you "ramp up" the temperature with a precisely defined rate, e.g. **15** °C/min, to the process temperature selected.
- During loading and ramping up, you may not want to start the oxidation, so you run N₂ or Ar through the furnace. After the process temperature has been reached, you switch to O₂ for dry oxidation or the right mixture of H₂ and O₂ which will immediately burn to H₂O if you want to do a wet oxidation.
- After the oxidation time is over, you ramp down the temperature and move the wafers out of the furnace.
- Moving wafers in and out of the furnace, incidentally, is not easy.
 - First you have to transfer the wafers to the rigid rod system (usually SiC clad with high purity quartz) that moves in and out, and than you have to make sure that the whole contraption easily 2 m long moves in and out at a predetermined speed v without ever touching anything because that would produce particles.
 - There is quite a weight on the rods and they have to be absolutely unbendable even at the highest process temperature around 1150 °C.

Of course, the rods, the quartz furnace tube and anything else getting hot or coming into contact with the Si, must be ultrapure - hot Si would just lap up even smallest traces of <u>deadly fast-diffusing metals</u>.

And now to the *difficult* part: After the process "works", you now must make sure that it works exactly the same way (with tolerances of < 5%) on all 100 wafers in a run in, from run to run, and independently of which one of the 10 or so furnaces is used.

So take note: Assuring stable process specifications for a production environment may be a more demanding and difficult job than to develop the process in the first place.

You see: At the final count, a "simple" process like thermal oxidation consists of a process recipe that easily calls for more than **20** precisely specified parameters, and changing anyone just a little bit may change the properties of your oxide in major ways.

All these points were emphasized to demonstrate that even seemingly simple processes in the production of integrated circuits are rather complex.

The processes to be discussed in what follows are no less complex, rather more so. But we will not go into the finer points at great depth anymore.

There are two more techniques to obtain **SiO₂** which are so important that we have to consider them in independent modules:

Local oxidation - this will be contained in the following module, and

CVD deposition of oxide - this will be part of the CVD module.