

## 7.2.2 Important MEMS Processes

### Cavities

As we have seen, we need to make relatively large cavities. For doing that we have to remove relatively large volumes of **Si**, and this we can do by either wet chemical etching or by plasma etching - [as in microelectronics](#).

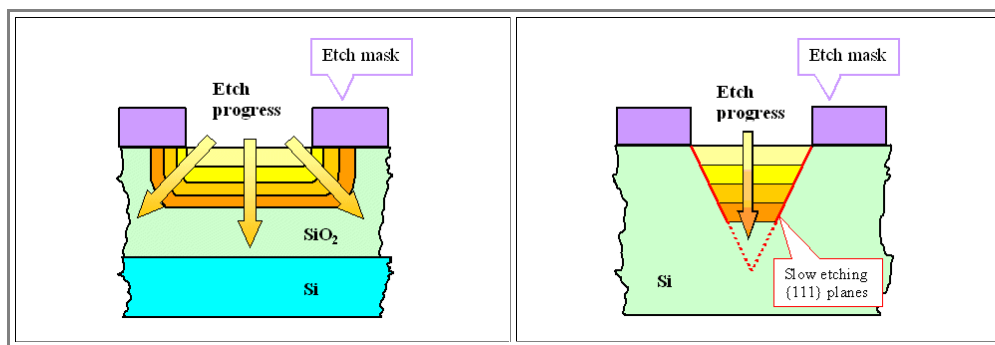
However, plasma etching in microelectronics has not been engineered to remove a lot of **Si**. If we would use existing processes for etching cavities, it would take forever and probably would run out of producing the structures desired with increasing depth.

So let's modify those processes! Easier said than done. In fact there is one special process for deep etching with plasma, that we will look at below, Here we must now resort to the still rather **black art** of wet etching.

Using isotropic chemical etchants thus as **HF** / **HNO<sub>3</sub>** / **CH<sub>3</sub>COOH** (very unhealthy!) would just produce a big hole with lateral dimensions no smaller than twice the etching depth and some semiglobular shape.

**Fortunately**, there is such a thing as **anisotropic chemical etching**; the most prominent chemical for this is **KOH** (at somewhat elevated temperatures like **80 °C**. **Unfortunately**, alkali metals like **potassium (K)** are **anathema** for **Si** microelectronics (and thus for MEMS with some electronics integrated on the chip).

The figures compare schematically isotropic and anisotropic etching.



We cannot use **KOH** for the reason given above, but its anisotropic etching properties gives us the idea to try other basic (instead of acidic) etchants.

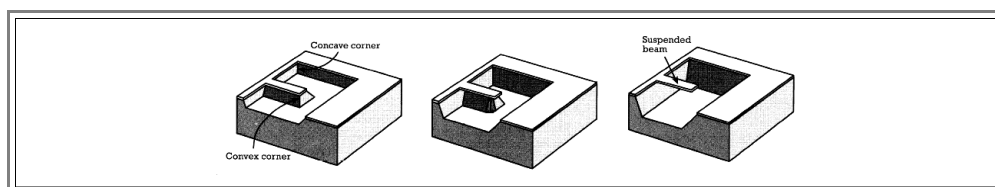
What works and what we can use is, for example, aqueous **TMAH** (=tetra-methyl ammonium hydroxide; **(CH<sub>3</sub>)<sub>4</sub>NOH**) solutions in water (**5 % - 30 % TMAH in H<sub>2</sub>O**), a harmless chemical from the viewpoint of easily-contaminated **Si**.

Often (but not always) the **{111}** planes act as "**stop planes**" for anisotropic etchants, meaning that the etching or dissolution rate on this plane is far slower than on other planes.

In fact, the etching rate with these anisotropic etchants on **{111}** planes can be about **1 000** times smaller than on the other low-indexed planes like **{100}** or **{110}**; i.e. the **selectivity** is very large.

If you start in a square opened in some chemistry-resistant mask on typical **(100) Si**, you will start to etch a perfect inverse truncated pyramid. If you etch long enough, a perfect pyramid with **{111}** sides will result. as shown below - we have already seen [nice examples](#).

In the case of cantilever production as shown below, some more time would be needed to remove the "bridges" below the larger cantilevers. A time sequence would look like this:



[Electrochemical etching](#) may also be highly anisotropic, but is not really used so far in mainstream MEMS technology.

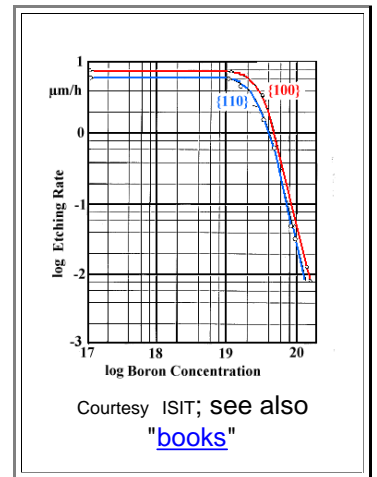
Why is anisotropic (electro)chemical etching of **Si** a [black art](#)? There is an easy simple answer, and a more general one. The simple answer is simple: Nobody, for example, can calculate the selectivity of the etchants named above (as a function of concentration, temperature, doping, and so on; of course) from "first" or even second principles.

Now let's be more general: If you consider the **parameter space** of an etching experiment - it is very large! You can have many compositions of the basic etchant (including those "magical" drops of this or that); the temperature and the doping of the **Si** are parameters of importance. If you actually do an **electrochemical** experiment, you have voltage and current as additional parameters. The results may even depend on the state of illumination of the the **Si**.

Let's assume you have found a set of parameters, i.e. one "point" in this large parameter space, that works. Now change one of the many parameters somewhat. Can you (or anybody else) predict theoretically (i.e. without trying out, or reverting to experiments already done before) what is going to happen? The answer is: Most likely not. Sometimes, predictions can be made with a high level of confidence, quite often unforeseen and amazing things happen. Understanding etching of semiconductors in general therefore is still a wide open field of research

Nevertheless, with anisotropic chemical etching we now have a way to make cavities in principle. This is great, but not good enough. We do not only need a large selectivity with respect to crystallographic planes, we also need selectivity with respect to the doping of the **Si**. We will see why in the next sub-chapter.

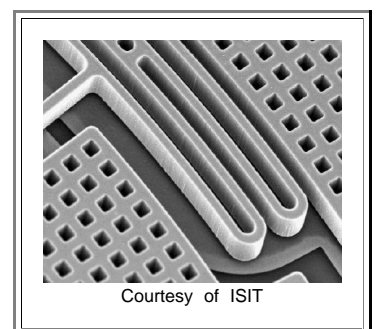
- Fortunately (i.e. not predicted), for highly Boron (**B**) doped **Si**, i.e. **p<sup>+</sup>-Si**, the etching rates in all direction become very small, i.e. **p<sup>+</sup>-Si** does not dissolve anymore in basic etchants.
- The effect is quite dramatic: As soon as some minimum **B**-concentration [**B**] has been reached, the etching rate decreases with the fourth power of [**B**] as shown in the figure.



## The Bosch Process

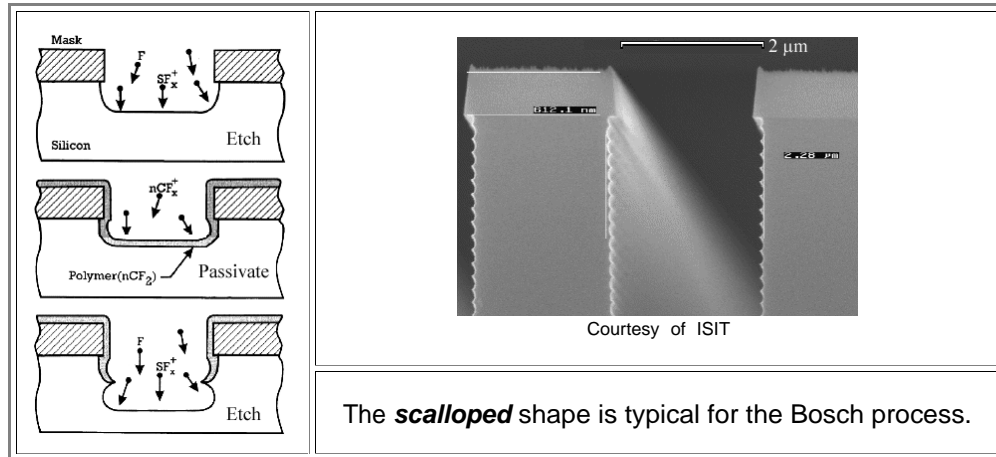
As mentioned above, the [large variety](#) of [plasma etching](#) (also called **dry etching**) process developed for microelectronics are, of course, used in MEMS technology in many ways, but are simply not meeting all etching needs.

- Look at the free moving part of a gyro in the picture. It consists of poly-**Si** about **10 μm** thick that was "etched out" of a solid poly-**Si** layer with very high precision. It has all those little square holes in it for reason we will encounter later - but they need to be made.
- The solution to many of the dry etching or plasma etching problems encountered in MEMS technology is the **high-rate plasma etching** or the "**Bosch process**", patented in **1994** by Bosch
- The Bosch process essentially decouples the two necessary ingredients for anisotropic etching: **1.** dissolving **Si** at the bottom, and **2.** passivating the side walls, i.e. protecting them against etching.



The process alternates every **10 s** or so between etching **Si** and depositing some polymer on the walls. The polymer deposited on the bottom of the area to be etches is taken off much faster than the coating of the side walls, allowing some net etching into the depth at rather high speeds and for (almost) arbitrary depth.

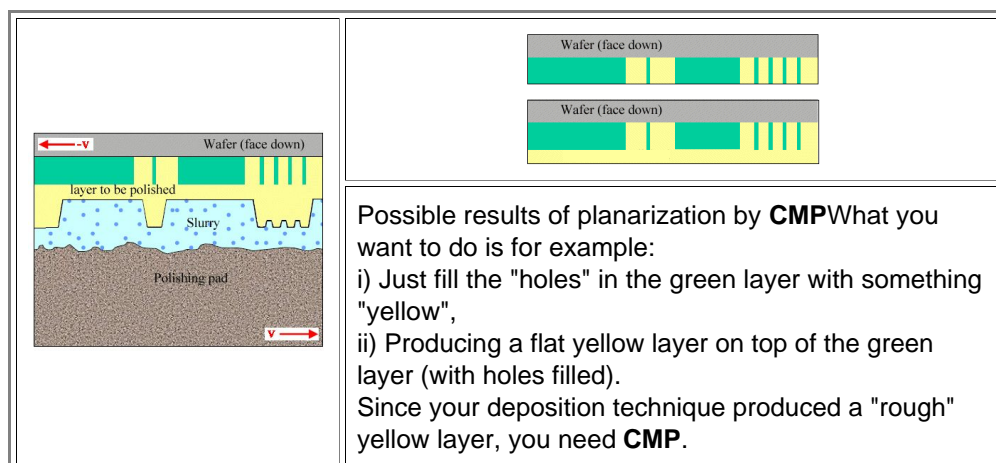
- The etching process uses essentially **SF<sub>6</sub>** plasma chemistry.
- Passivation uses a fluorocarbon process, using gases like **C<sub>4</sub>F<sub>8</sub>** , **C<sub>3</sub>F<sub>6</sub>** or **CHF<sub>3</sub>**.
- Etching rates of **> 10 μm/min** per minute are possible, allowing cost-effective processing even for deep etching, including all the way through a (thinned to about **300 μm**) wafer.
- The frequent switching of gases and etching parameters makes the system quite complicated (and expensive). But the Bosch process is one of the key processes for MEMS and probably will **spawn** many variants in due time.



## Chemical Mechanical Polishing

We have encountered **chemical-mechanical polishing** or **CMP** before - [in one sentence](#).

- It is a key process for both microelectronics and MEMS. Microelectronics, however, thrived without **CMP** for more than **20** years, while many MEMS products would just not be possible without **CMP**.
- How is it done? In principle, it is simply grinding down the surface with an abrasive. This is the **mechanical** part of polishing. The **chemical** part comes in because your **slurry** - the liquid or viscous goo that contains fine particles of the abrasive materials, also contains chemicals that etch off **only** the mechanically damaged **Si**. **KOH**, for an example, would be good at that but is not used for the reason [given before](#).
- The amazing (and at the time of its introduction quite unbelievable) thing is that you can use **CMP** with nanometer precision on huge **Si** wafers - if you know all the tricks of the trade, of course. Recipes for slurries, the kind of polishing pads to use for the kind of material you want to polish off, and everything else are closely guarded secrets.



The figure shows the basic principle and what one can achieve.

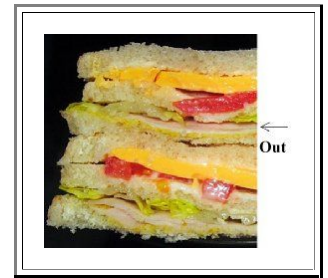
- **CMP** may also be counted to some extent among the **"black arts"**. Not only does it have a chemical etching components; its [parameter space](#) is extremely large, too, and hard to consider in theory.

## Other Typical MEMS Processes

There is much more, but here we can only give a few hints.

### Etching of sacrificial layers.

- Free standing structures like in the [gyro](#) first are made on top of some *sacrificial layer* that is etched off at an opportune time. These sacrificial layers must meet a number of requirements, the most important one being that they must be compatible with whatever is under and above it, and that there must be a way of etching them off selectively without dissolving anything else.
- This is tough, man! If you just consider the etching process, it's like taking out the ham from your sandwich without opening it up and without doing *anything* to the bread, butter, mayonnaise, salad leaf etc., above and below. This means you can attack the ham only from the side, and that will take a long time during which a lot can go wrong.
- One common trick is to make the top layer (the one to be free standing after the process) full of small holes, so you can attack the sacrificial layer from more places. This is why we always see "perforated" structures, e.g. [further up](#).
- There is no way, of course, to do the etching with plasma. You *must* use "black art" chemistry. On occasion, even gases are being used (e.g. **HF** vapor, an extremely dangerous chemical) because pulling filigree structures out of a liquid may make them stick together (think about your [hair in the bathtub!](#)) - even after drying.



### Wafer bonding

- Consider an atomically flat **Si** wafer with a perfectly clean (non-oxidized) surface. Now put another wafer with an equally perfect surface on top. The two wafers would instantly bond into one piece of **Si** (with a grain boundary at the interface if they were not exactly at the same orientation).
  - Real **Si** wafers are neither atomically flat nor perfectly clean - but they come rather close. If you join them, they will stick together by [van-der-Waals bonding](#). Add a little heat and some mild pressure, and they will bond to one piece of **Si**. Now you have a well-established process that was developed for microelectronics, but is not much used (too expensive).
  - In MEMS technology, wafer bonding might be a crucial process. Gyros or optical micro mirrors operating at their resonance frequency, for example, must be kept in a cavity with a precisely established low pressure because that determines the [damping](#) of the system. The tough part is to keep that pressure constant for **> 20** years.
  - So bond another **Si** wafer (with a cavity) on top. Better hermetically sealing with no long-time stability problem at the interface is not possible, but there are many ways how to do worse.
- There are many tricks to achieve efficient bonding without having to be ultra-pure and ultra-precise. Moreover, in MEMS technology not only **Si**-wafer to **Si**-wafer bonding is used, but also, for example **Si**-wafer to glass-wafer bonding.
- This is easy to understand. Hermetically sealing your micro-mirrors with a **Si** wafer on top is fine as long as you only want to work only in the infrared. If you want to use *visible* light, your top better be transparent.
  - Going through the various material combinations and the technologies used to bond them is far beyond the scope of this lecture course. Suffice it to state that bonding technologies are a major part of MEMS processing.