Advanced Plasma Processing: Etching, Deposition, and Wafer Bonding Techniques for Semiconductor Applications

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1. Introduction

Plasma processing techniques are one of the cornerstones of modern semiconductor fabrication. Low pressure plasmas in particular can achieve high radical density, high selectivity, and anisotropic etch profiles at low temperatures and mild voltages. This gentle processing environment prevents unwanted diffusion and degradation of materials due to heat and lattice damage from ion bombardment. Plasma treatments have a minimal effect on existing wafer structure, which is a key requirement for large scale integration schemes such as CMOS. In addition, recent progress in plasma-assisted wafer bonding has demonstrated low temperature, low pressure recipes utilizing O₂ plasma surface treatment for joining dissimilar semiconductor materials, such as silicon (Si) and indium phosphide (InP) (Fang et al., 2006).

In this chapter, we will cover the applications of plasmas to etching and depositing materials, as well as novel processing modalities such as surface treatments in preparation for wafer bonding. High aspect ratio nanopillars have recently been fabricated using this technique, with features as small as 22 nm etched over 1.25 μm deep. In particular, our use of Al₂O₃ as a mask material along with cryogenic wafer temperatures has demonstrated to increase the etch selectivity of silicon over mask to more than 5000:1, enabling ultrathin masks for nanoscale pattern transfer (Henry et al., 2009a). After patterning, in situ deposition can encapsulate these structures in preparation for further processing.
We will also discuss our work in hybrid integration of Si/III–V materials for on-chip optics applications using plasma etching and bonding techniques. Plasma-assisted wafer bonding has realized hybrid Si/III–V structures, using the III–V material as an efficient gain medium while maintaining the economic and electronic integration benefit of a silicon platform (Sun et al., 2009). In addition, smooth etches have realized low loss waveguides in silicon. Finally, unique planar microcoils have been made using the deep etching and in situ deposition capabilities in the ICP-RIE.

2. Plasmas in Electronics Processing

Plasmas are found in a wide range of industrial applications, including ashing, sputtering, etching, and chemical vapor deposition. However, the types of plasmas used in each process vary greatly. In this section, we will describe the important parameters of plasmas for understanding ICP-RIE etching and deposition, and establish how they relate to our desired processing results.

2.1 Figures of Merit

Central to semiconductor processing is the high fidelity transfer of a pattern onto a substrate through addition, modification, or removal of material. In order to quantify the ability to accomplish this selective processing, it is useful to have a few figures of merit to describe the process, namely:

- **Etch rate** – controllable and robust to small deviation in processing conditions. Depending on the application, one may want a higher etch rate for increased throughput or a lower etch rate for precision
- **Uniformity** – both at each feature and across the wafer
- **Selectivity** – the ability to etch only the desired material, relative to the etching of mask and other substrate materials
- **Anisotropy** – the verticality of the etch profile. Also, the nanostructure of this vertical surface is important in many applications, particularly waveguides
- **Damage** – any surface or substrate damage acquired from the processing technique.

In addition, the processing environment is important, as some conditions will have a deleterious effect on existing wafer structure. Chief among these is the temperature of processing. Heating (and cooling) can cause many problems, including thin film delamination due to thermal expansion coefficient mismatch, unwanted dopant diffusion, and other negative effects (Schmidt, 1998). In the CMOS industry, the wafer’s tolerance to temperature fluctuations is often captured as a thermal budget (Takeuchi et al., 2005), which means that minimizing use during one process step can give more latitude in other steps.

2.2 Plasma Characteristics

Plasma is a partially ionized gas with a combination of free electrons, ions, radicals, and neutral species. To create and sustain a plasma in the laboratory, energy input is required. Generally, this energy is transferred via coupling of an external electromagnetic field to the plasma constituents. Different coupling methods generate plasmas with different characteristics. Most useful to reactive ion etching (RIE) are those generated by
glow discharge plasmas (GDP), capacitive coupled plasmas (CCP), inductively coupled plasmas (ICP), or some combination thereof.

In a GDP process, electromagnetic energy is delivered as a voltage applied between two conducting plates, known as the cathode and the anode. The applied voltage is usually constant or in the low frequency regime, such that the characteristic time of field variation is longer than the response time of the system. The potential difference generates an electric field across the gases in the chamber. Plasma initiation occurs when a small initial population of charged species is accelerated through the electric field and collides with other molecules, causing them to ionize. A relatively high voltage is required to initiate and sustain the plasma, which is a severe processing drawback. High voltage will cause the resultant energy of incident ions on the cathode to be high, favoring rough, physical processes (sputtering) over smooth, chemical processes (surface reactions). This will lower the selectivity to masking materials and cause sidewall roughening due to mask erosion. For these reasons, GDP sources are often used to sputter materials rather than etch anisotropically (Chapman, 1980).

In a CCP process, energy is again supplied as a voltage between an anode and a cathode plate, but in a time-varying fashion. Most commonly, a radio frequency (RF) voltage is applied to the plates. The frequency of operation is often at 13.56 MHz, which is a band reserved for industrial use by the Federal Communications Commission in the United States. In this time-varying field, electrons in the plasma oscillate between the anode and the cathode plates. Collisions of rapidly moving electrons with the slowly moving ions cause further ionizations. However, massive ions are less mobile and cannot track the rapidly oscillating electric field changes. By placing a capacitor between the anode plate and the RF supply, negative charge accumulates on the plate (typically referred to as the table). The resulting potential difference between the plasma and the negatively charged plate is called the self-bias $V_b$. The electric field due to $V_b$ drives the positive ions in plasma towards the negatively charged table. This is the basis for traditional reactive ion etching.

In an ICP process, the excitation is again a time-varying RF source, but is delivered inductively, instead of capacitively, resulting in a changing magnetic field. This changing magnetic field, through the Maxwell-Faraday equation, induces an electric field that tends to circulate the plasma in the plane parallel to the CCP plates. Similarly to a CCP, collisions of the rapidly moving electrons with the slowly moving ions cause further ionizations. Loss of electrons from the plasma through the grounded chamber walls tends to create a static voltage, deemed the plasma voltage $V_{\text{plasma}}$. This is distinct from the self bias $V_b$, as will be examined later. Inductive coupling is generally realized through a large 4 to 5 turn coil encircling the plasma chamber. In the typical geometry, this means that one is able to change ion density and other plasma parameters without significantly perturbing the incident energy of the ions.

The experimental results discussed in this chapter are realized on Oxford Systems Plasma Lab 100 ICP-RIE 380 systems, which utilize a CCP and an ICP power source, as seen in Fig. 1. Throughout the text, we will frequently refer to the CCP power as the “forward power” in order to distinguish it from the ICP power and to emphasize its role in driving ions toward the sample surface. This dual plasma powering affords the greatest flexibility in altering plasma characteristics such as ion density and bias voltage independently of each other. These systems have been extensively studied, particularly for silicon etching (Jansen et al., 2009).
2.3 Processing Parameters

There are a few important features of an ICP-RIE plasma that have an effect on etching. Most noticeable during operation is the region of glow discharge, where visible light emission occurs from a cloud of energetic ions and electrons. As the gas particles move in the plasma, some collisions occur which transfer energy to bound electrons. When these electrons return to their ground state, a photon may be emitted. The color of the plasma is characteristic of the excited gas species, because the photon energy is a function of the electronic structure of the gas molecules and their interactions with surrounding molecules (Hodoroaba et al., 2000). This can be a good diagnostic for incorrect plasma striking conditions or other adverse changes in your plasma. For example, in a multiple gas recipe, sometimes the emission looks like only one of your gas species, instead of the average of the colors. This happens when the other species are not being ionized, and thus will cause the process to take on a completely different character from a calibrated recipe.

Beneath the glow discharge region is a dark space, where atoms are no longer excited into emitting photons due to the depletion of electrons. This dark space is also the part of the plasma that most directly affects the paths of incoming ions that will accomplish the etching. Neutral atoms and other ions will tend to scatter the otherwise straight path of the ions from the edge of the glow discharge to the cathode.

We can characterize this spreading in both energy and trajectory into probability distributions, called the ion angular distribution function (IADF) and the ion energy distribution function (IEDF) (Jansen et al., 2009). These distributions, depicted in Fig. 2, describe the likelihood that an incident ion has a given energy and trajectory. IADF strongly affects the sidewall profile, as a wider IADF corresponds to a higher flux of ions reaching the sidewall. Similarly, the IEDF controls the types of processes the ions can be engaged in when they reach the surface, including removing passivating species, overcoming activation energies for chemical reactions, and enhancing sputtering yield. These processes determine the performance characteristics of an etch, so understanding these effects and recognizing associated faults are paramount to optimizing a recipe. Parameters controlling the IADF
and IEDF include the bias voltage $V_b$, the ion density, the gas composition, and the mean free path (which also depends on the aforementioned parameters).

![Fig. 2. Illustration of the ion angular and ion energy distribution functions, with hypothetical resultant etched profile distortion. Points in IEDF correspond to different ion kinetic energies, while points in the IADF correspond to different angles of incidence](image)

### 2.4 Etch Reaction Dynamics

In wet chemical processes, etching is accomplished through physical dissolution or reaction-specific dissolution (Reinhardt & Kern, 2008). This takes place at any exposed surface and thus results in isotropic etching, although the etch rate can vary along different crystalline orientations due to the bonding state variation of the surfaces. A good example of crystalline anisotropy in Si wet processing is potassium hydroxide (KOH) etching, which is widely used for making MEMS structures that capitalize on the direction-dependent etch rate of KOH (Wolf & Tauber, 2000). However, in a myriad of planar processes that are utilized in the semiconductor industry, an anisotropic etching profile with sidewalls perpendicular to the wafer surface is frequently required for effective pattern transfer.

In order to prevent the isotropic or crystalline anisotropic behavior of our processing gases, the sidewalls must be protected from further etching. This is accomplished by forming a passivating or inhibiting layer on the sidewall, in one of the following ways:

- **Surface passivation**
  
  - inserting gases in the plasma which react with wafer materials and forming involatile compounds (Legtenberg et al., 1995)
  
  - freezing volatile reaction products at the structure’s walls using, e.g., cryogenic wafer cooling (Aachboun et al., 2000)

- **Inhibitor deposition**
  
  - using polymer precursor gases to form physical barrier layers (e.g., C₄F₈) (Kenoyer et al., 2003)
  
  - eroding and redepositing inert mask materials

All of these processes are important to consider when evaluating an etch, as there may be problems with the etch profile related to the deleterious effect of one of these regimes. We
use both surface passivation and inhibitor deposition techniques in the following etch descriptions.

2.5 Time-Dependent Processes
In addition to the previously discussed processing parameters, we have one additional variable at our disposal: time. A notable example of using time as an etching parameter is the Bosch silicon etch process, which occurs in a time-multiplexed manner, or “pulsed mode,” using an etching plasma followed immediately by a deposition plasma. Alternatively, we can try to accomplish the etching and deposition simultaneously by using a plasma that contains both etching and deposition gases. This is called a “mixed mode” process. Finally, we can also tune our processes to change continuously over time in response to the changing surface condition of our wafer, or to compensate for a negative effect due to the initial conditions of the wafer.

2.6 Conclusion
As we have seen in this section, plasma processes depend on a large number of variables, which accounts for both their sensitivity and their flexibility. By having basic knowledge of the underlying physical processes, diagnosing your processes becomes more intuitive and makes recipe invention and refinement much easier. In the following sections, we will refer to many of the concepts covered here to explain results and understand how we arrived at a given recipe. However, there is still no replacement for hands-on experimentation for building an even greater understanding of ICP-RIE processing.

3. Deep Silicon Etching
Silicon is the workhorse of the semiconductor industry, and thus etching of Si is one of the most frequent processes used in a fab. In order to achieve deep etches in silicon using an ICP-RIE, three basic etch requirements must be met. First, the etch must have a relatively high etch rate. A slow etch rate is cost prohibitive in a high throughput, industrial process and has the potential for the introduction of process variations, leading to etch failures. Second requirement is that the etch must have a high selectivity, or preference, to etch the silicon as compared to the etch mask. Insufficient selectivity limits the maximum etch depth or requires complicated thick masks to compensate for erosion, limiting the minimum feature size. Finally, the etch must remain anisotropic throughout the etching process. If lateral etching occurs, pattern transfer begins to fail as the etching continues vertically. To date, only two etching modalities have the potential to stand up to these rigorous requirements: pulsed mode and mixed mode silicon etches. Both etch schemes employ forms of etching combined with passivation that actively protect sidewalls during etching and improve anisotropy. Each has their own advantages and disadvantages which will become clear during the discussion. To illustrate the differences between the two modes of etches, two widely used etches will be discussed here. For the pulsed mode etch we describe the chopping Bosch silicon etch, which uses gas “chopping” to alternately etch and deposit inhibitor on your surface, and for the mixed mode etch, we demonstrate the cryogenic silicon etch, which uses a different gas chemistry to form passivating compounds
at the sidewalls at the same time as etching. Note that both gas chemistries reviewed here can be used in either pulsed or mixed mode.

As mentioned, the chopping Bosch etch requires two alternating plasma steps. The first step etches the silicon for a short period then rapidly shuts off the gas and plasma. The second step then initiates a plasma that deposits an inhibitor film on exposed surfaces. This alternating sequence continues as the etch progresses. Inherent in the discreteness of the etching is notching on the sidewalls that occurs every step. The duty cycle between steps controls the etch angle and the total length of the combined steps controls the depth of the notching.

In contrast, cryogenic silicon etching combines the discrete etch and passivation steps into a single continuous etch. By using cryogenic temperatures from -80 °C to -140 °C, improvements in etch mask selectivity and passivation effects are enabled. Both of these etching chemistries, mask selections, and characteristics will be reviewed here along with their applications or demonstrations.

3.1 Gas Chemistries

Chopping Bosch etching utilizes sulfur hexafluoride, $\text{SF}_6$, as the etching gas and octafluorocyclobutane, $\text{C}_8\text{F}_8$, as the passivation gas. As described earlier, when the $\text{SF}_6$ is injected into the chamber, the plasma ionizes and radicalizes the gas molecules to create a mixture of $\text{SF}_x$ and $\text{F}_y$ ions and neutrals, where $x$ and $y$ range from 0 to 6 and 1 to 2, respectively (Cliteur et al., 1999). The potential established between the plasma and the substrate, due in part to the ICP and the CCP power, causes the electric field that drives the ions down to the substrate. The unmasked silicon then bonds to the fluorine atoms to create the volatile tetrafluorosilane ($\text{SiF}_4$) etch product which is then pumped away from the chamber. The etch becomes a combination of chemical bonding and mechanical milling; the milling is established from the momentum imparted to the ions from the electric field. While the chemical etching is essentially isotropic in nature, the mechanical milling is anisotropic. After a few seconds of etch time, the $\text{SF}_6$ flow is rapidly terminated and the $\text{C}_8\text{F}_8$ gas is then injected into the chamber for the passivation step. During this step, the $\text{C}_8\text{F}_8$ fragments into smaller $\text{CF}_x$ ions which act as film precursors (Takahashi et al., 2000). A Teflon-like film forms on the substrate, on both the vertical and horizontal surfaces. The thickness of the protective layer is dependent on the passivation step time. Once the deposition is complete and the subsequent etch step begins, the ions first mill away the horizontal passivation layers and then begin again with the silicon etching. This cyclic process of etching followed by passivating continues on until the etching is terminated, leaving the etched silicon structures coated with the passivation polymer.

The cryogenic silicon etch also utilizes the $\text{SF}_6$ chemistry similar to that of the chopping Bosch. However, by lowering the substrate’s temperature, and by simultaneously injecting $\text{SF}_6$ and oxygen gas, $\text{O}_2$, a passivation layer is created simultaneously as the silicon is etched. The current understanding of the chemical process is that oxygen ions combine with the fluorine bonded to the silicon surface prior to the silicon’s removal and forms a $\text{SiO}_x\text{F}_y$ layer. The exact composition of this layer is a topic of current research (Mellhaoui et al., 2005). In a manner similar to the chopping Bosch passivation, the $\text{SiO}_x\text{F}_y$ passivation layer protects the exposed vertical silicon while the unmasked horizontal silicon is etched way. To make this passivation process as energetically favorable as the chemical reaction of making $\text{SiF}_4$, the substrate temperature is required to be cooler than approximately -80 °C. When the silicon
is warmed back up to room temperature, the $\text{SiO}_x\text{F}_y$ becomes volatile and leaves the sample (Pereira et al., 2009).

### 3.2 Mask Selection

The ultimate test of a mask is the fidelity of pattern transfer into the silicon over the entire etching period. Since the mask interacts with the etching process parameters, it is vital to understand which masks to use for different etches. As stated earlier, if the selectivity is too low a thicker mask is required to achieve the desired etch depths. Furthermore, as the edge of the mask erodes it will impart undesired slope or features to the sidewalls of the etched structure, often referred to as mask-induced roughness. For these reasons, deep silicon etching requires higher selectivity masks. Conventional silicon etch masks are metal, oxides, and resist.

Metal masks, such as chrome, offer the advantage of high selectivity as high as thousands to one. This is primarily due to their lack of chemical reactivity with the etch gas molecules and their mechanical strength. However, metal masks typically induce detrimental effects such as notching at the top of the etched structures, due to image forces, and unwanted masking due to redeposited metal introduced by ion sputtering. A particular problem with chrome during the cryogenic etch is that oxygen radicals appear to be locally deactivated around the mask reducing the silicon passivation layer near the top of the mask (Jansen et al., 2009). Silicon dioxide masks typically offer high selectivity (150:1 for Bosch and 200:1 for cryogenic etching) with the added cost of more complicated patterning. The oxide layer must be grown or deposited, followed by pattern transfer from another material or resist into the oxide mask. Increasing the number of processing steps increases the effort needed for accurate pattern transfer as well as the potential for reduction in mask fidelity. Resist masks offer the simplicity of a single processing step along with good selectivity (approximately 75:1 for Bosch and 100:1 for cryogenic etching). These selectivity values highly depend on process conditions and are seen to widely vary. Jansen et al. have commented that sidewall protection using resist is better than that using oxide masks due to the erosion of the resist mask providing additional material to protect the etched walls (Jansen et al., 2009).

Several new masks have recently demonstrated improvements both in selectivity and in ease of integration. Sputtered aluminum oxide, or alumina, provides mask selectivity greater than 5000:1 for cryogenic etching. Because of the extremely high selectivity, only a thin layer is required for masking. This makes the film easily patterned via resist lift-off, instead of traditional ion milling for hardmask pattern transfer. Patterning difficulty is only slightly increased as compared to traditional resist processing. Starting with a photoresist mask, a thin layer of alumina is sputtered onto the sample. This is followed by lift-off of the undesired alumina and the resist using acetone. Due the brittle nature of the material, the alumina cleanly fractures and easily lifts off. Furthermore, since the alumina is electrically insulating, image force effects and undercuts seen in metal masks are not seen with this mask. Removal of the alumina mask is easily achieved using buffered hydrofluoric acid or ammonium hydroxide combined with hydrogen peroxide, both of which do not significantly etch silicon.

A second new mask innovation is using gallium (Ga) to mask silicon (Chekurov et al., 2009). The Ga mask is implanted by a focused ion beam (FIB), where a gallium beam is focused on a silicon sample and writes out the pattern in a similar way to other direct-write lithography.
techniques. The dose can be accurately controlled by manipulating the time the beam spends focused on the silicon as well as the accelerating voltage of the beam. This offers the advantage of high mask resolution on small feature sizes (~40 nm) without needing a polymer to be patterned or a developer to be used. Typical dosing creates masks around 30 nm in thickness and offers greater than 1000:1 selectivity in a cryogenic etch. Unfortunately, mask removal poses a problem since the gallium atoms are implanted in the silicon and damage to the silicon surface has not yet been characterized. Since using the Ga mask is, in a sense, a maskless and resistless technique, pattern definition can take place on any surface upon which the beam can be focused. This presents the opportunity of multidimensional patterning, such as patterning on a pre-etched sidewall to create a lateral mask.

3.3 Etching Conditions and Optimization

Control over the etch rates, selectivity, sidewall profile, and etch roughness is achieved through tuning process parameters. The major controllable parameters include ICP power, forward power, temperature, chamber pressure, and gas flow rates. While this list is not all inclusive, these parameters directly control the state of the chamber and therefore the plasma. Many subtleties also play an important role in the etch process. This list would include silicon loading, chamber conditioning, and chemical interactions in the gas chemistry and with the mask. Each etch process will have optimization parameters that will be reactor specific, but this section will assist in building intuition for both the Bosch and the cryogenic etch.

The ICP power controls the amount of ionization occurring for a given gas flow rate and chamber pressure. Typically, as the ICP power is increased, the amount of ions created will also increase. This will increase the chemical etch rate, both vertically and laterally, increase the milling etch rate, reduce the selectivity by milling the mask away faster, and reduce the effect of passivation by bombarding the sidewalls more due to the ion angular dispersion effect. If the vacuum pumping rate does not change, e.g., when controlling the throttle valve position instead of the chamber pressure, then when increasing the ICP power one can measure the fact that more gas is ionized by measuring the chamber pressure. It should be noted that increasing the ICP power does not increase the etch rate infinitely. In fact there is an optimum ICP power for a given etch gas flow rate. These trends apply for both Bosch and cryogenic etching for the SF$_6$ chemistry. Increasing the ICP power for the passivation step of chopping Bosch, similarly to the etching, will increase the thickness of the passivation for a given passivation time. A subtle effect of increasing the ICP power is that it also slightly increases the bias between the plasma and the electrode. For the Bosch etch, the bias from the forward power is typically much greater in magnitude than plasma potential increase from the ICP change and the effect is generally unnoticed. Since the cryogenic etch uses very little forward power, applying more ICP power can significantly increase the amount of milling occurring. Another subtle effect is that a higher etch rate also increases the substrate’s temperature. For the cryogenic etch, it is estimated that the exothermic formation of SiF$_4$ releases 2 W/cm$^2$ for an 8 μm/min etch rate. For an unmasked 6” Si wafer, this results in approximately 360 W of exothermic heating.

Increasing the forward power establishes a larger electric field between the plasma and the table electrode. By imparting more momentum to the ions, the silicon milling rate increases. This usually increases just the vertical etch rate, but due to the IAD (ion angular distribution) effect the lateral etching does slightly increase. Since the milling action increases, the
erosion rate of the mask also increases, thereby reducing the selectivity. Similar to the
temperature effect from increasing ICP power, increasing the forward power increases the
rate and energy of ion bombardment to the substrate. This effect is easily calculated from
the potential difference and the ion flux for the cryogenic etch and is estimated around 0.5
W/cm².
The Bosch etch is typically insensitive to temperature effects, while the cryogenic etch is
extremely responsive to any temperature changes. Since the Bosch etch is performed at 20
°C, the polymer passivation layer is far from both the melting and freezing regimes.
However, the high temperature dependence of the passivation reaction during the cryogenic
etch means even small temperature fluctuations change the etching profile. Heating by as
little as 5 °C during the cryogenic etch reduces the passivation rate and thereby induces
undercutting due to image force effects. Passivation during the cryogenic etch roughly
begins to occur around -85 °C. However, if the wafer is too cold, SF₆ etch gases and SiFₓ
product gases can freeze on the sample sidewalls, adding to the SiOₓFᵧ passivation layer.
Variations in table temperature by 5 °C due to oscillations in the table temperature
controller have been seen to change the profile of deep etches adding a sinusoidal curvature
to the sidewalls. Temperature is typically controlled by cooling the stage with liquid
nitrogen or water and thermally connecting the wafer to the table by flowing helium
between them. When silicon samples smaller than a full wafer are etched, they require
thermal conductivity to the carrier wafer. This is accomplished by using thermal grease or
Fomblin pump oil on the backside of the wafer to the substrate. Removal of the thermal
grease is done with trichloroethylene and the Fomblin is easily removed by isopropanol.
Chamber pressure controls the amount of gas in the chamber for ionization. As noted
during the ICP power discussion, changing the amount of incident ions controls both etch
rate and selectivity. For a given ICP power, there is an optimum gas flow rate for SF₆.
Increasing the pressure can be accomplished by shutting the throttle valve or by injecting
more gas. A subtle effect of increasing chamber pressure is that it also increases the
scattering collisions of ions traversing the Faraday dark space. This creates a larger angular
spread in incident ions to the substrate, or increases the IAD. This increases the amount of
undercut or lateral etch.
Other parameters which can alter both the Bosch and the cryogenic etch are not necessarily
due to changing a mechanical feature on the reactor. Changing the amount of exposed
silicon can also change etch results. Increasing the ratio of exposed silicon to masked silicon
changes the amount of ions needed for etching and will significantly reduce the etch rate.
As explained earlier, the exothermic nature of etching more silicon also induces an increase
in substrate heating. A positive effect, however, is that for large silicon loading, slight
changes in mask patterning have relatively minor effects in etch results. This is a convenient
feature for establishing an etch for a wide range of users. It also reduces the effect of
changing the etch as the etch goes deeper into the silicon and effectively exposes more
silicon surface. Cleanliness of the chamber can also change the effects of etches. Since the
plasma interacts with the sidewalls as well as the substrate, residual molecules on the
sidewalls can be redeposited on the etched surface, causing micromasking, or can
chemically react with the etch gas. For this reason, it is highly recommended that good
chamber cleans followed by chamber conditioning be performed prior to etching.
3.4 Application: High Aspect Ratio Pillars and Metallization Liftoff

Using the high selectivity of photoresist for the cryogenic silicon etch, fabrication of high aspect ratio micropillars was demonstrated (Henry et al., 2009a) and serves as an example of achievable profiles using the mixed mode etching process. These pillars were utilized for validating theories concerning radial $p$–$n$ junctions for applications of solar cells (Kayes et al., 2008). The patterns transferred to a 1.6 $\mu$m thick photoresist on a silicon substrate were groups of disks 5, 10, 20, and 50 $\mu$m in diameter in a hexagonal array. The spacing between each disk grouping was equivalent to the diameter of the disks, i.e., each 5 $\mu$m disk was separated from its nearest neighbor by 5 $\mu$m, the 10 $\mu$m disks by 10 $\mu$m, etc.

Fig. 3. High aspect ratio silicon micropillars: This cross-sectional SEM of 5 $\mu$m wide and 83 $\mu$m tall silicon micropillars demonstrates the cryogenic silicon etch using a 110 nm thick alumina etch mask. The very tops of the pillars indicate that mask erosion is beginning.

Concluding etch profile optimization, multiple samples of the patterns were etched for varying times. Since each etch had an array of the four different diameters, a direct study of aspect ratio, i.e., ratio of the etched depth to width, dependence upon etch depth was made. Assuming that the etch rate was comprised of the etch rate of silicon with no structures (zero aspect ratio) minus a linear dependence on aspect ratio, a simple differential equation may be solved to yield the following:
Here $E_0$ and $b$ are the zero aspect ratio etch rate and the aspect ratio dependent etch rate, respectively. The equation solves for the etch depth $d$ given the etched trench width $w$ and the etching time $t$. Using this equation, etches were performed achieving an aspect ratio of 17.5:1. The angle of the micropillars’ sidewalls was controlled by varying the oxygen flow rate, which allowed for passivation rates to be controlled and consequently changing the angle of the profile up to $6^\circ$. This number appears small at first but when deep etches are being performed, controlling the angle can prove critical to not etching the base of the pillars to a point.

![Graph showing etch rates and aspect ratio dependence](https://www.intechopen.com)

**Fig. 4.** Etch rates and aspect ratio dependence: This graph contains data points taken from etches creating the silicon micropillar arrays of 5, 10, 20, and 50 μm diameter pillars as well as the solutions to the solved differential equation for the various widths. It becomes evident that as the aspect ratio of the etched trench increases, the etch rate slows down. This is the so-called “Aspect Ratio Dependent Etching” or ARDE effect.

A second use of the cryogenic etch is based on the high selectivity of the etch mask. Since very little resist is eroded away during etching, the remaining etch mask becomes useful as a layer for metallization liftoff. This fabrication sequence was employed for creating silicon...
microcoils (Henry et al., 2009b). Using a 1.6 \( \mu \text{m} \) thick patterned photoresist, a cryogenic etch was used to etch highly doped silicon. The structures then had varying thicknesses of chemically vapor deposited (CVD) amorphous silicon dioxide. Following the deposition, copper was thermally evaporated into the trenches with thickness up to 15 \( \mu \text{m} \). Lift-off of the silicon dioxide and metal using acetone was then performed. Typically for conventional metallization, resist heights are required to be 3–4 times thicker than the metal being deposited with necessary rigorous sidewall profile control. Here, since the metal is approximately 10 times thicker than the resist, the depth of the cryogenic etch can replace the thick resist requirements as well as reliably accomplishing the profile requirements needed for the thick metal deposit. This fabrication sequence created planar copper microcoils embedded in silicon and insulated from the substrate using silicon dioxide.

![Planar copper microcoils in cross section. Coils are copper 10 \( \mu \text{m} \) thick embedded in silicon and insulated using a 1 \( \mu \text{m} \) thick CVD oxide. Using the high selectivity of the cryogenic silicon etch, thick copper metallization is possible with lift-off achieved using the etch mask.](image)

**4. Nanoscale Silicon Etching**

Unlike deep silicon etching, nanoscale etching requires neither extraordinary selectivity nor large etch rates. On the contrary, moderate selectivity of 5:1 is acceptable and slower etch rates, 100–200 nm/min, are more useful for accuracy of etch depths. Further, Bosch etching and cryogenic etching prove to be unsuitable for very small structures due to the notching and lateral etching of the two chemistries respectively. In general, nanoscale etch properties should include smooth and highly controllable sidewalls, slow etch rates, and low undercutting effects. To meet the first two requirements, mixed mode gas chemistries become useful due to the simultaneous etching and passivating. Proper choices in masks can reduce undercutting effects. This section will discuss several emerging mask
technologies and demonstrate nanoscale etching using \( \text{SF}_6 / \text{C}_4\text{F}_8 \), termed as the Pseudo Bosch etch here.

### 4.1 Gas Chemistries

Although the cryogenic etch creates very smooth sidewalls, its inherent undercut is typically too much for the nanoscale regime. Furthermore, the etch rates are too high for accurate control on the nanoscale. A combination of the Bosch gases introduced in a mixed mode process creates an ideal etch recipe which has allowed silicon nanopillars with an aspect ratio of 60:1 and diameters down to 20 nm. To etch the silicon, \( \text{SF}_6 \) is again used while \( \text{C}_4\text{F}_8 \) is used to passivate simultaneously. Since ions are constantly needing to mill the continuously deposited fluorocarbon polymer layer, the etch rate significantly reduces to 200–300 nm/min. Etch recipe parameters are similar to the cryogenic etch and are around 1200 W for the ICP power and 20 W for the forward power. The advantage of using the \( \text{C}_4\text{F}_8 \) as the passivation gas also extends to not requiring cryogenic temperatures.

### 4.2 Mask Selection

Typical masks for nanoscale etches are based on the difficult patterning requirements. To define structures down to 20 nm, e-beam resists such as polymethylmethacrylate (PMMA) are employed with thicknesses ranging from 500 nm down to 30 nm. The advantage of using this as the etch mask is the simplicity in pattern transfer: once the e-beam patterning is complete, the resist can be developed leaving the patterned etch mask. The disadvantage is that typical selectivity values range from 4:1 to 0.5:1. This implies that only very shallow etches can be performed on the very small structures since thicker e-beam resists are difficult to expose for small structures. However, a great advantage is achieved by using alumina etch masks with this etch. A thin layer of alumina, approximately 30 nm thick, can serve as an etch mask yielding selectivity of better than 60:1. This allows for e-beam resists, with thickness to be patterned and developed, followed by having the alumina sputter deposited. After lift-off in acetone, the alumina pattern remains on the silicon. Another common etch mask is nickel, which is patterned similarly to that of sputtered alumina. Sputtered nickel offers good selectivity with the disadvantage of increased mask undercutting due to image forces. We recently have also demonstrated using implanted gallium as an etch mask for silicon nanostructures. With this method, Ga ions are implanted in the silicon substrate using a focused ion beam. The dwell time of the beam combined with the current determines the dosage while the beam accelerating voltage determines the depth and spread of the mask. Typical threshold dosages are about \( 10^{16} \) ions/cm\(^2\) or 2000 \( \mu \text{C/cm}^2 \). For comparison, typical resist sensitivities range from 200–1200 \( \mu \text{C/cm}^2 \) when exposed on a 100 keV electron beam lithography system. Using a 30 kV beam, we estimate the Ga layer to be approximately 20 nm thick. Using the Pseudo Bosch etch, selectivities greater than 50:1 have been demonstrated using a Ga mask with resolution of better than 60 nm. At this point, we suspect that the resolution has not reached the intrinsic limit imposed by the implantation process, and is instead limited by our beam optics.
4.3 Etching Conditions and Optimization
By changing the ratio of the etch gas to passivation gas, SF\textsubscript{6}:C\textsubscript{4}F\textsubscript{8}, the sidewall profile can be controlled. A typical ratio is 1:3 with the absolute gas flow rates dependent upon chamber volume, as sufficient flow is required to establish a chamber pressure of 10 mTorr; a good starting point is roughly 30 and 90 sccm respectively. Increasing the ratio improves the etch rate, reduces the selectivity, and drives the sidewall to be reentrant. Typical ICP power is around 1200 W combined with a slightly higher forward power than that of the cryogenic etch of around 20 W. Increasing the forward power again reduces the selectivity with a slight improvement in etching rates. Unlike cryogenic mixed mode, this etch is typically performed at room temperature or 15–20 °C.

4.4 Application: Waveguides and Nanopillars
Since passivation occurs during etching, very straight and smooth sidewalls can be fabricated on nanoscale structures. In particular, combining this feature of the Pseudo Bosch etch with the high selectivity of the alumina etch mask, impressive 60:1 aspect ratio nanopillars have been demonstrated. Pillars were created by first patterning PMMA using a 100 kV electron beam and developing the pattern using methyl isobutyl ketone (MIBK) and isopropanol solution. A 30 nm thick alumina layer was then sputtered and lifted off leaving the alumina mask on silicon. The Pseudo Bosch etch was then performed with an etch rate of 250 nm/min leaving well defined arrays of silicon nanopillars. The smallest diameter created was 22 nm for a pillar standing 1.26 μm tall.
Fig. 7. High aspect ratio silicon nanopillars: These cross-sectional SEMs, taken at 45°, of a) 73 nm diameter and 2.8 μm tall silicon nanopillar, b) 40 nm diameter and 1.75 μm tall silicon nanopillar, c) tungsten probe contacting a 200 nm diameter and 1.25 μm tall silicon nanopillar, and d) array of alternating 40 nm and 65 nm diameter pillars 1 μm tall demonstrate the Pseudo Bosch silicon etch using a 30 nm thick alumina etch mask

5. Nanoscale Indium Phosphide Etching

In contrast to the previously discussed fluorine-based etch recipes, many III–V materials require the use of chlorine-based chemistries. This is due to the difference in chemical properties of the etch products. As seen in the previous section, the proposed mechanism for Ga masking of Si is the formation of involatile GaF$_x$ compounds that prevent further etching. Thus, for etching of Ga and other similar compounds, we expect that a Cl$_2$-based etch will result in faster etching rate and smoother sidewalls from the readily removed etch products. In this section, we will discuss an InP etch that uses a hybrid gas mixture of Cl$_2$, CH$_4$, and H$_2$.

5.1 Gas Chemistries

The gas composition of this etching recipe is a hybrid between two established InP recipes. Specifically, high etch rate recipes with Cl$_2$- and Cl$_2$/Ar-based plasmas are well known but suffer from sidewall roughness and require high processing temperatures to volatilize InCl$_x$ species (Yu & Lee, 2002), as illustrated in Fig. 8. Smooth etch recipes with CH$_4$/H$_2$ plasmas have also been studied but have prohibitively slow etch rates. In this case, the smoothness is a result of two factors. Firstly, the likely etching mechanism of InP is the evolution of volatile products PH$_3$ and In(CH$_3$)$_3$, which can be controlled by adjusting the gas flow rates (Feurprier et al., 1998). Secondly, the deposition of CH films from the source gases serves to protect the sidewalls (von Keudell & Möller, 1994). In our etch, we utilize a precise ratio of source gases that balances all these properties and takes interactions into account, such as removal of H and Cl ions by formation of HCl.
Fig. 8. Micromasking due to insufficient heating (left). By increasing ICP power and thus raising sample temperature, micromasking is removed (right)

5.2 Mask Selection
Appropriate masks for the InP etch are metals and dielectrics. This is due to the high rate of mask erosion inherent in the etching conditions. The forward bias and thus bias voltage that drive ions toward the wafer surface are much higher than those found in the SF$_6$-based silicon etches in previous sections. This will make the etch more milling, and will help to maintain the same etch characteristics in other stoichiometries of interest, such as InGaAsP compounds. We utilized masks of silicon dioxide spheres and evaporated Au layers in the etching experiments. The selectivity of oxide was approximately 10:1; however, faceting occurred before the mask was completely eroded, limiting the useful selectivity to a more modest 4:1. In deeper nanoscale etching applications, a silicon nitride or metal mask is preferred as it has high selectivity and does not suffer from faceting as readily as oxide. As seen in Fig. 9, the metal hardmask has eliminated most pattern-induced roughness.

Fig. 9. Anisotropic InP etch using a metal hardmask. Smoothness is only limited by mask irregularities
5.3 Etching Conditions and Optimization

Our etch had Cl₂:H₂:CH₄ ratio of 8:7:4 with actual gas flows of 32 sccm Cl₂, 28 sccm H₂, and 16 sccm CH₄ and a chamber pressure of 4 mTorr. The table was heated to 60 °C to reduce polymer deposition, and no helium backing was applied in order to have the plasma heat the sample. This heating is key to proper etch characteristics, as too little heat will cause micromasking due to involatile gas products such as InClₓ. The forward power was 180 W, found experimentally by varying until an anisotropic profile was achieved without excessive mask erosion. This resulted in a cathode bias of approximately 200 V. ICP power was 2200 W, also found experimentally by monitoring the transition of “black” InP to smooth InP due to the cessation of micromasking during etching. The etch rate of pure InP was measured to be 1.2 μm/min.

During some etches with identical conditions, a roughening of the bottom surface was noticed due to chamber cleanliness. The sensitivity of this etch to chamber condition is not as high as the cryogenic Si etch described earlier, but reproducible results require a regular cleaning schedule to return the chamber to a known “clean” state. This is best implemented by running a short, minute-long SF₆ cleaning plasma just before etching to remove any contaminants that are readily incorporated into the plasma. For long term cleanliness, periodic hour-long SF₆/O₂ plasma is run. The frequency depends on what other etches have been performed previously, but is typically one hour of cleaning per three to four hours of etching. In an industrial setting, this could be done in shorter periods between each wafer to maintain a constant chamber state.

6. Inductively Coupled Plasma Chemical Vapor Deposition

ICP-RIE systems have been demonstrated in this chapter to be a gentle environment for etching both silicon and III–V materials. Over the last decade, research has extended this useful environment to the deposition of thin dielectric films and conductive silicon layers.
A common film deposition method is low pressure chemical vapor deposition, which is typically performed at temperatures around 600 °C. The low pressure limits unwanted gas phase reactions, while the high temperature ensures that there are adequate diffusion and energy to overcome any activation barrier in the desired reactions. By using plasma enhanced chemical vapor deposition (PECVD), typical deposition temperature can be lowered to the range of 300 °C to 400 °C. By adding a gas ring to improve gas uniformity in an ICP-RIE, the new operation of ICP chemical vapor deposition, or ICP-CVD, is added to the machine. In ICP-CVD, depositions may proceed with temperatures in the range of 50 °C to 150 °C. Densities of layers deposited at 70 °C using ICP-CVD have now become comparable to PECVD at 350 °C. This illustrates the key advantage of ICP-CVD over traditional CVD processes: high density films deposited at lower temperatures.

A typical PECVD reactor has a single power source establishing both the plasma density and the ion flux in a method similar to that of an RIE (although new PECVD reactors can have a 13.56 MHz source combined with a kHz source, where both of the sources create RIE plasmas albeit at different frequencies). A unique advantage of ICP-CVD reactors over PECVD is that both the ion density and the ion flux can be independently controlled. In this case, the ICP power changes the ion density while the forward power controls the ion flux (Lee et al., 2000). This provides another method for tuning film parameters such as optical index, deposition rate, and density.

Since the ICP-RIE has improved efficiency ionizing the gas over an RIE-only reactor, and due to the lower operating pressures of ICP-RIEs over that of PECVD, significantly less gas is required for depositions. Typical flow rates for gases are 20 to 100 sccm. However, typical deposition rates for ICP-CVD are significantly lower than PECVD, ranging from 6–30 nm/min whereas PECVD rates range from 60–250 nm/min. The slower rates in the ICP-CVD allow for precise control over thin films with the intended use as dielectrics. Using the same method for creation of thicker films for etch masks becomes impractical.

### 6.1 Gas Chemistries

Types of films available using ICP-CVD include SiO$_2$, SiN$_x$, SiON$_{x}$, a-Si, and SiC. All of these silicon containing compounds require the source gas silane (SiH$_4$). Because silane is pyrophoric and will combust spontaneously in air, it is typically diluted to 2-10% levels using inert carrier gases such as nitrogen, helium, or argon. The silane flows into the reactor through a gas ring surrounding the table. A second gas is injected from the top of the chamber where the ICP ionizes the gas, creating various radicals and ions. In the same manner as in etching processes, the electrostatic potential difference between the plasma and the table drives the ions to the wafer surface, where they chemically combine with silane to create the film. Typical additive gases are N$_2$O for silicon dioxide, NH$_3$ and N$_2$ for silicon nitride, both N$_2$O and NH$_3$ for silicon oxy-nitride, and CH$_4$ for silicon carbide. The ratio of additive gas flow rates to the silane flow rate into the chamber can be used to change the index of the deposited material by making nonstoichiometric films. For a silicon dioxide film with refractive index of 1.44, the required ratio of pure silane to N$_2$O ranges from 0.3 to 0.4. For a silicon nitride film with refractive index of 2.0, a ratio of pure silane to NH$_3$ ranges from 1.18 to 1.28.
6.2 Process Tuning

A deposited film can be evaluated by several basic material properties: refractive index (indicative of stoichiometry), density, and stress levels. Evaluation of refractive index is easily performed using an ellipsometer which usually can also determine thickness simultaneously. This type of measurement also allows for deposition rate to be directly measured. Measurement of the density of the film is performed by wet etching in buffered hydrofluoric acid with the figure of merit being the etch rate; slower rates imply denser films. Stress is usually measured using an interferometer to determine the radius of curvature of wafer bowing.

Control over the refractive index is coarsely adjusted by setting the ratio of flow rates of gases into the reactor. Although increasing the chamber pressure can raise the index of the film, a more linear increase in index can be achieved by increasing the forward power of the deposition. Film density is most directly controlled through the plate’s temperature, where higher temperatures correspond with higher densities. However, increasing the plate temperature above 150 °C for silicon dioxide deposition and above 70 °C for silicon nitride has little additional effect on film density. Another method for increasing density is by increasing the forward power and thus ion flux. Although control over stress in silicon dioxide is difficult, silicon nitride stress can be tuned using chamber pressure. Increasing chamber pressure can move stress values from negative to positive, and thus zero stress silicon nitride films are possible. For further quantization of process trends of silicon nitride and silicon dioxide, see (Lee et al., 2000).

6.3 Applications of ICP-CVD films

As seen earlier, in situ deposition of these films immediately after etching is possible. Deposition of a cladding layer or even a dielectric stack after etching an optical structure is possible, due to the refractive index tunability of the films. Particularly as the thermal budgets of many wafers shrink due to the use of exotic materials such as HfO2 (He et al., 2005), low temperature processes such as ICP-CVD will become more valuable.

7. Plasma-Assisted Wafer Bonding

In previous sections we have focused individually on the processing of Si-based materials and InP-based III–V materials. Silicon systems make modern computing and data processing possible, while III–V gain materials generate and amplify light for transmission over optical fibers. Silicon is an indirect semiconductor and thus a very poor convertor of electricity to light while III–V’s are direct semiconductors and thus efficient convertors of injected electrons to photons. Now it is widely recognized that major future progress in both computers and optical communication will require chip-scale integration of these two types of materials (Mathine, 1997).

Heteroepitaxy by molecular beam epitaxy (MBE) or metalorganic chemical vapor deposition (MOCVD) has proved to be very successful for lattice-matched material systems such as GaAs- and InP-based ternary or quaternary compounds. However, due to the large lattice mismatch (4.1% between Si and GaAs and 8.1% between Si and InP), heteroepitaxial growth of III–V’s on silicon has not produced high-quality material for practical applications. The high density of threading dislocations in the epitaxial layers greatly reduces the lifetime of fabricated devices (Pearson et al., 1996).
7.1 Direct Wafer Bonding

As an alternative to epitaxial growth, direct wafer bonding provides a way to join together two flat and clean semiconductor surfaces at room temperature without the restriction of matching lattice constants. The intermolecular and interatomic forces bring the two wafers together and the bonds form at the interface. By introducing a superlattice defect-blocking layer, dopants and defects are prevented from migrating from the bonding interface to the active region so that the luminescence from the multiple quantum well structure can be preserved (Black et al., 1999). To increase the bond strength, a high-temperature post-bonding annealing step is usually required. However, this high-temperature annealing step induces material degradation and is incompatible with backend Si CMOS processing. For this purpose, many efforts have been put into reducing the annealing temperature while keeping a strong bonding (Takagi et al., 1996; Krauter et al., 1997; Berthold et al., 1998).

For Si-to-InP wafer bonding, a pre-bonding oxygen plasma treatment for both wafer surfaces has been demonstrated to yield a very spontaneous bonding at room temperature (Pasquariello & Hjort, 2002). Similar to previously discussed etching plasmas, the pre-bonding plasma aims to have a high density of chemically active species arrive at the surface with a low incident power to minimize surface damages, such as dislocations, that work against bond formation. The post-bonding annealing temperature can be below 200 °C while the interface strength can be as high as the bulk fracture energy of InP. As mentioned in previous sections, the plasma affects the bonding surfaces both physically and chemically. The oxygen plasma is used to remove hydrocarbon and water molecules so as to reduce the probability of the formation of interfacial bubbles and voids during post-bonding annealing. Additionally, the plasma treatment generates a very smooth and reactive thin oxide layer which helps in bonding process.

7.2 Bonding Procedure

We have succeeded in transferring InGaAsP epifilms to Si substrates using oxygen plasma assisted wafer bonding technique. We start with a Si wafer and an InP wafer with InGaAsP epitaxial film. The InGaAsP epifilm consists of an InGaAs contact layer at the top, a $p$-InP upper cladding layer at middle followed by an InGaAsP active layer and then an $n$-InP lower cladding layer at the bottom. The total thickness of the epifilm is ~2 μm. The bonding procedure begins with solvent cleaning of both surfaces. A 10-nm-thick oxide layer is grown on top of the Si wafer to enhance the bonding strength. The surfaces of the two wafers are then activated through exposure to oxygen plasma, and bonded together under a pressure of 0.1 MPa at 150 °C for 2 h. Following the bonding process, the InP substrate is removed by HCl wet etching. Fig. 11 clearly shows the cross-sectional structure consisting of the remaining InGaAsP epifilm bonded onto the Si substrate. The bonding interface between the epifilm and Si is thin and smooth. Fig. 11(a) focuses on one end of the epifilm: it is evident that the top InGaAs layer is protrusive at the end due to its different composition from the $p$-InP layer below. Fig. 11(b) focuses on a middle part of the epifilm. The pyramids at the top of $p$-InP layer are results of HCl wet etching during InP substrate removal due to the damaged outer InGaAs layer.
Fig. 11. SEM cross-sectional view of InGaAsP epifilm on Si by wafer bonding after the step of InP substrate removal. The InGaAsP epifilm consists of an InGaAs contact layer at the top, a $p$-InP upper cladding layer at middle followed by an InGaAsP active layer and then an $n$-InP lower cladding layer at the bottom. (a) Zoom at one end of the epifilm. The top InGaAs layer is protrusive at the end due to its different composition from the $p$-InP layer below. (b) Zoom at middle of the epifilm. The pyramids at the top of $p$-InP layer are results of HCl wet etching during InP substrate removal due to the damaged outer InGaAs layer.

7.3 Application: Hybrid Si/III–V Optoelectronic Devices

With this epilayer transferring technology, the two disparate materials Si and III–V’s now can be brought together to realize a variety of active devices on Si, such as lasers (Fang et al., 2006; Fang et al., 2008; Sun et al., 2009), amplifiers (Park et al., 2007a), modulators (Chen et al., 2008; Kuo et al., 2008), and detectors (Park et al., 2007b). To take the hybrid Si lasers for an example: as seen in Fig. 12, the hybrid Si/III–V structure consists of a prepatterned SOI wafer and a III–V epilayer bonded together. The III–V epilayer has been reported to be AlGalnAs (Fang et al., 2006) or InGaAsP (Sun et al., 2009) quaternary semiconductor compounds, either of which can be epigrown onto an InP substrate with very high crystal quality with current state of the art. In our work (Sun et al., 2009) the thicknesses of the buried SiO$_2$ layer and the undoped Si device layer are respectively 2.0 $\mu$m and 0.9 $\mu$m. The Si waveguide is defined using e-beam lithography and SF$_6$/C$_4$F$_8$ plasma reactive ion etching as described in detail in previous sections. The Si to the two sides of the waveguide is entirely etched down to the SiO$_2$ layer, and the waveguide width ranges between 0.9 $\mu$m and 1.3 $\mu$m. A 5-$\mu$m-wide center current channel by means of proton implantation on its two sides is created to enable efficient current injection. In a working device, the injected current starts from the top $p$-side contact, passes through the center current channel in the $p$-InP cladding and the InGaAsP active region, then bifurcates in the $n$-InP layer until it reaches the $n$-side contacts on both sides (not shown in Fig. 12). This hybrid structure is designed to support a joint optical mode, whose profile overlaps both materials. The modal gain is obtained by the evanescent penetration of the joint mode into the III–V active region. The devices are referred to as “hybrid Si evanescent lasers” (Fang et al., 2006). Single facet output power from the Si waveguide can exceed 10 mW at room temperature, making these devices ready for practical use (Sun et al., 2009).
Advanced Plasma Processing: Etching, Deposition, and Wafer Bonding Techniques for Semiconductor Applications

8. Conclusion

In this chapter, we have described and explored some of the emerging plasma applications of etching, deposition, and surface modification to semiconductor materials. The process latitude available in modern ICP-RIE systems has enabled these novel processes. This is a direct consequence of the multitude of changes one can effect on a plasma by changing the pressure, driving fields, gases, temperature, and other parameters as discussed in the introduction. In particular, temperature-controlled stages capable of cryogenic cooling have given process engineers the ability to tune the types and rates of chemical reactions that occur on their samples. This was strongly illustrated in our discussion of cryogenic silicon etching, but a similar scheme could be imagined for other materials, given appropriate gas selection. We demonstrated both deep Si etching appropriate for MEMS as well as nanoscale Si etching, and discussed the difference in processing details between these two regimes.

New applications of these etches were discussed. By combining a chemically inert mask (Al₂O₃) with the nanoscale Pseudo Bosch silicon etch, high aspect ratio nanopillars were demonstrated, along with the Ga implantation masking concept that also relies on the chemical inertness of the mask material to etch gases. Similarly, by taking an established mask technology (photoresist) and combining it with the highly selective, fluoropolymer-free deep cryogenic silicon etch and in situ deposition, novel inductive elements were fabricated.

Next, applications of III–V etching and wafer bonding were discussed. By using a plasma treatment of both surfaces, we were able to join together two dissimilar semiconductors because this method is not affected by lattice constant mismatch. This bonding technique showed room temperature spontaneous bonding which can be annealed at 200 °C or less while keeping a high interfacial strength.

Finally, the culmination of all of the previously discussed processing techniques in Si and III–V’s was the demonstration of a hybrid evanescent Si/III–V laser, made using Si etching
and wafer bonding. The III–V active material was used as an efficient gain medium while the Si provides the guidance and feedback for the lasing modes and also serves to couple the light out to the rest of the on-chip photonic circuit. Lasers had room temperature output of more than 10 mW emitted from a single facet of the Si waveguide. The authors expect further innovations in the plasma processing field and hope these examples have demonstrated the great versatility of this research.

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10. References


Semiconductor technologies continue to evolve and amaze us. New materials, new structures, new manufacturing tools, and new advancements in modelling and simulation form a breeding ground for novel high performance electronic and photonic devices. This book covers all aspects of semiconductor technology concerning materials, technological processes, and devices, including their modelling, design, integration, and manufacturing.

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