# Focused Ion Beam Microscopy and Micromachining

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### Abstract

The fairly recent availability of commercial focused ion beam (FIB) microscopes has led to rapid development of their applications for materials science. FIB instruments have both imaging and micromachining capabilities at the nanometer–micrometer scale; thus, a broad range of fundamental studies and technological applications have been enhanced or made possible with FIB technology. This introductory article covers the basic FIB instrument and the fundamentals of ion–solid interactions that lead to the many unique FIB capabilities as well as some of the unwanted artifacts associated with FIB instruments. The four topical articles following this introduction give overviews of specific applications of the FIB in materials science, focusing on its particular strengths as a tool for characterization and transmission electron microscopy sample preparation, as well as its potential for ion beam fabrication and prototyping.

### Introduction

The focused ion beam (FIB) microscope has gained widespread use in fundamental materials studies and technological applications over the last several years because it offers both high-resolution imaging and flexible micromachining in a single platform.

The FIB instrument is similar to a scanning electron microscope (SEM), except that the beam that is rastered over the sample is an ion beam rather than an electron beam. Secondary electrons are generated by the interaction of the ion beam with the sample surface and can be used to obtain high-spatial-resolution images. In most commercially available systems, Ga ions are used, and their sputtering action enables precise machining of samples. In conjunction with the gas-injection capabilities on these systems, which enable ion-beam-activated deposition and enhanced etching, a range of sample fabrication schemes are possible.

During the last 25 years, FIB instrumentation has become an important technology for a wide array of materials science applications, from circuit editing and transmission electron microscopy (TEM) sample preparation to microstructural analysis and prototype nanomachining. Most modern FIB instruments supple-

ment the FIB column with an additional SEM column so that the instrument becomes a versatile "dual-beam" platform (FIB–SEM, see Figure 1) for imaging, material removal, and deposition at length scales of a few nanometers to hundreds of microns. The FIB instrument becomes a powerful tool for nanomanipulation and fabrication through the augmentation of an FIB instrument with micromanipulators and gas injection for local chemical vapor deposition (CVD).

The first FIB instruments evolved from advances in field ion microscopes<sup>1</sup> and through the development of highresolution liquid metal ion sources (LMISs).<sup>2-4</sup> In the 1980s, FIB instruments were embraced by the semiconductor industry as offline equipment for mask or circuit repair. It was not until the 1990s that FIB instruments began to be used in research laboratories, and today there are commercial instruments available from multiple manufacturers.5 With the popularity of FIB instruments for TEM sample preparation, microstructural analysis, and nanomachining, dual-beam FIB instruments are becoming a versatile and powerful tool for materials researchers.

This introductory article focuses on the FIB instrument itself and the basic ion–solid interactions that lead to the various functionalities of FIBs. In the topical articles that follow, the major subspecialties of FIB research are discussed.

# The FIB Instrument

The basic functions of the FIB, namely, imaging and sputtering with an ion beam, require a highly focused beam. A consistent tenet of any focused beam is that the smaller the effective source size, the more current that can be focused to a point. Unlike the broad ion beams generated from plasma sources, high-resolution ion beams are defined by the use of a field ionization source with a small effective source size on the order of 5 nm, therefore enabling the beam to be tightly focused.

The ion source type used in all commercial systems and in the majority of research systems designed with micromachining applications in mind is the liquidmetal ion source (LMIS).<sup>6,7</sup> Of the existing ion source types, the LMIS provides the brightest and most highly focused beam (when connected to the appropriate optics). There are a number of different types of LMIS sources, the most widely used being a Ga-based blunt needle source. Ga has decided advantages over other LMIS metals such as In, Bi, Sn, and Au because of its combination of low melting temperature (30°C), low volatility, and low vapor pressure. The low melting temperature makes the source easy to design and operate, and because Ga does not react with the material defining the needle (typically W) and evaporation is negligible, Ga-based LMISs are typically more stable than other LMIS metals. During operation, Ga flows from a reservoir to the needle tip (with an end radius of about 10 μm), where it is extracted by field emission. A large negative potential between the needle and an extraction electrode generates an electric field of magnitude  $10^{10}$  V/m at the needle tip. The balance between the electrostatic forces and the Ga surface tension wetting the tapered W needle geometry results in the formation of a single Taylor cone at the needle tip. For typical emission currents used in FIB microscopes (~2 µA), a cusp forms at the tip of the Taylor cone with a tip radius of approximately 5 nm.

The simplest and most widely used ion beam columns consist of two lenses (a condenser and objective lens) to define the beam and then focus it on the sample, beam-defining apertures to select the beam diameter and current, deflection plates to raster the beam over the sample surface, stigmation poles to ensure a spherical beam profile, and a high-speed beam blanker to quickly deflect the beam

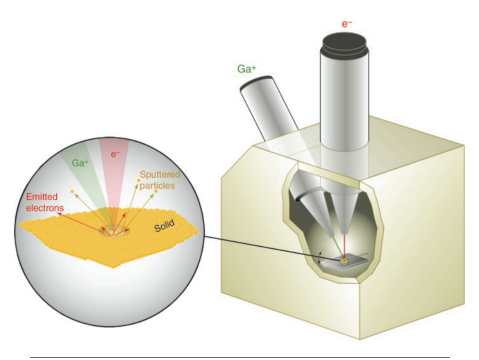


Figure 1. Schematic illustration of a dual-beam FIB-SEM instrument. Expanded view shows the electron and ion beam sample interaction.

off the sample and onto a beam stop such as a Faraday cup. Because the focusing strength of an electromagnetic lens is directly related to the charge/mass ratio of a particle, it is impractical to build electromagnetic lenses for ions (which would weigh thousands of kilograms); thus, focusing and steering are performed using electrostatic components rather than the electromagnetic components used for electrons.

The size and shape of the beam intensity profile on the sample determines the basic imaging resolution and micromachining precision. Generally, the smaller the beam diameter, the better the achievable resolution and milling precision, although the requirements for the two applications are not exactly the same.8 For the energies, currents, and acceptance angles used in typical FIB systems, the beam spot size is limited mostly by the chromatic aberration that results primarily from the energy spread of the beam due to space charge effects at the ion source and secondarily from the spherical aberration of the lenses. However, the ultimate spatial resolution for FIB imaging is, in fact, limited by sputtering and is thus sample-dependent.9 In modern FIB systems, the imaging resolution determined by the sputter-limited signal/noise usually is about 10 nm.

The sample is mounted on a grounded stage with three-axis translation, rotation, and tilt capabilities. The stage is designed

to have a eucentric point (i.e., a well-centered point such that the field of view is maintained when tilting the specimen) at the location where the two beams cross (or at the working distance of the ion beam, in the case of a single-beam FIB). The region of interest on the sample is moved to the eucentric point using translation and rotation and then tilted for the desired angle of beam incidence. The total current on the sample (sum of the incoming ion or electron beam and all emitted charged particles) is measured at the stage.

Depending on the application, the various emitted particles or radiation can be detected with appropriate detectors in the sample chamber. Traditional detectors such as those in an SEM can be used to detect the electrons or x-rays created by the interaction of the ion beam with the sample. The ions sputtered from the sample can also be detected using a variety of detectors such as charge electron multipliers, and mass selection of the sputtered charged particles is also possible (secondary ion mass spectrometry). FIBs derive an important additional functionality through the use of gas-injection sources to deliver gas locally to either enhance the etching rate or result in site-specific CVD. Secondary electrons generated by the incident ion beam (or, alternatively, the incident electron beam in dual-beam systems) can crack hydrocarbon precursor

gases, leading to local deposition of the conducting material (W, Pt, or C) or insulating material (SiO<sub>2</sub>); see the article by MoberlyChan et al. in this issue. The local deposition of material also enables sophisticated micromanipulation within the FIB chamber, made possible through micromanipulation accessories and the ability of the FIB to cut (sputter), paste (deposit), and watch (image) during a manipulation process within the chamber. The result is a system that can image, analyze, sputter, and deposit material all with very high spatial resolution and controlled through one software program. In a large part, it is this multifunctional versatility that has made FIB instruments popular among materials researchers.

# Ion-Solid Interactions

Ion–solid interactions play an important role in many different endeavors, ranging from fabrication of microelectronic devices to understanding distributions of cosmic gases. This brief introduction will be limited to the processes and conditions relevant to the use of FIB systems in materials science. For more detailed descriptions of ion–solid interactions, the reader is referred to books and review articles on the subject 6,10–12 as well as to literature specifically on FIB. 13–15

When an ion impinges on a solid, it loses kinetic energy through interactions with the sample atoms. This transfer of energy from the ion to the solid results in a number of different processes (see Figure 2):

- ion reflection and backscattering
- electron emission
- electromagnetic radiation
- atomic sputtering and ion emission
- sample damage
- sample heating

The ion typically comes to rest in the solid, leading to implantation of the ion. With the possible exception of electromagnetic radiation generation, all of these processes are important to FIB and FIB–SEM system applications and are described in the next sections.

### **Collision Cascade**

Ion kinetic energy and momentum are transferred to the solid through both inelastic and elastic interactions. In inelastic interactions (called electronic energy loss), ion energy is lost to the electrons in the sample and results in ionization and the emission of electrons and electromagnetic radiation from the sample. In elastic interactions (called nuclear energy loss), ion energy is transferred as translational energy to screened target atoms and can result in damage (displacement of sample

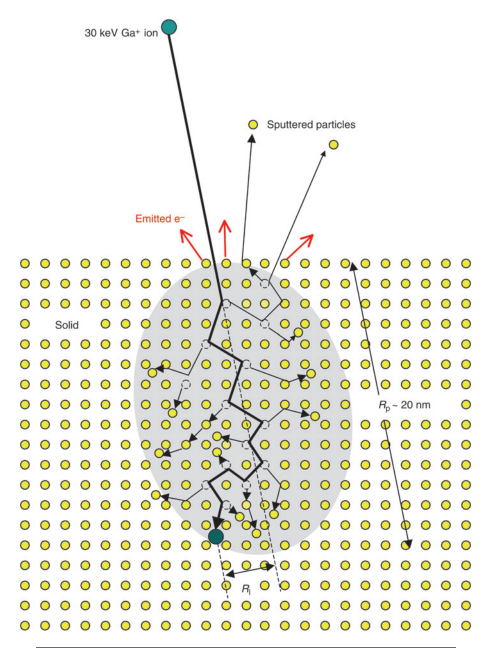


Figure 2. Schematic illustration of a collision cascade generated by a 30 keV  $Ga^+$  ion incident on a crystal lattice, showing the damage created in the collision cascade volume, and the projected range  $R_0$  and lateral range  $R_1$  of the implanted ion.

atoms from their initial sites) and sputtering from the sample surface.

The most widely accepted concept for ion–solid interactions is the collision cascade model<sup>16,17</sup> (Figure 2). For the case of 5–30 keV Ga impinging on most solids, the collision cascade involves a series of independent binary collisions (the linear collision cascade regime). If the translational energy transferred to a target atom during a collision exceeds a critical value called the displacement energy, the atom will be knocked out of its original site, for

example, creating an interstitial–vacancy pair in a crystalline sample. This primary recoil atom may have sufficient energy to displace further sample atoms (secondary recoils), thus generating a volume where large numbers of atoms have excess kinetic energy. If a displacement collision occurs near the surface, the recoil atom may be emitted from the solid and lead to sputtering. The displacement energy (typically on the order of 20 eV) is much larger than the binding energy for the atoms (of the order of 1 eV), reflecting the fact that

the collisions are nonadiabatic, because of the very short time scale.

After approximately 10<sup>-11</sup> s, the 5–30 keV Ga ion comes to rest in the solid, and the energies of all particles participating in the cascade have decreased below the displacement energy. At this point, the collision cascade has ended. What remains are the emitted particles and radiation, and ion beam damage such as lattice defects, incorporated Ga, and heat, all of which may continue to interact and evolve.

Molecular dynamics calculations are ideally suited for simulating collision cascades, because of the short length and time scales. Monte Carlo calculations are also well suited to simulating ion-solid interactions by including the "frictional" electronic stopping and stochastic elastic collisions. The most widely used Monte Carlo simulation is the program TRIM or SRIM (transport, or stopping range, of ions in matter). 18 Such calculations for 30 keV Ga into elements from Li to Bi show that roughly two times as much of the ion energy is lost to nuclear energy losses than to ionization energy losses (the former from interactions with the nucleus of an atom and the latter from interactions with the electrons). Of the nuclear energy losses, the majority is lost through sample atom vibrations or heating rather than through vacancy formation.

The projected and lateral ranges of the 30 keV Ga scale inversely with the sample density and are between 10 nm and 100 nm (projected) and between 5 nm and 50 nm (lateral). TRIM sputtering yields (sputtered target atoms per incoming ion) are between 1 and 20 for normalincidence 30 keV Ga and increase somewhat with atomic number.6 However, sputtering yield predictions depend critically on surface binding energies, which are not well known and are sensitive to surface structure and chemistry. TRIM vacancy generation predictions between 300 and 1000 vacancies per incoming ion are overestimates because defect diffusion and interactions are ignored. In addition, discrepancies between experimental and simulated ranges and collision cascade shapes in crystals can be expected, because TRIM samples are isotropic and cannot capture channeling effects. Despite these limitations, such calculations are invaluable in predicting trends and in estimating the effects of ion-solid interactions.

### Ion Beam Imaging

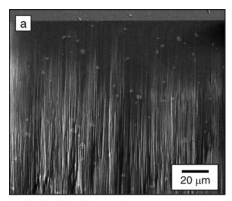
In the same manner that images are generated in an SEM, the ion beam can be rastered over a sample surface and the emitted electrons, particles (atoms and

ions), and electromagnetic radiation can be detected. Conventional SEM imaging is based on detecting the secondary electrons (SEs). To date, most imaging in a FIB is based on detecting the low-energy electrons, often referred to as ion-induced secondary electrons (ISEs). Typically, 1-10 electrons with energies below 10 eV are generated per incoming 5-30 keV Ga ion. These electrons are created by both kinetic and potential emission from the top few atomic layers where the primary ion impacts the solid as well as where backscattered or sputtered particles exit the sample. The total low-energy electron yield depends strongly on surface oxidation and contamination and thus will change as the surface is sputter-cleaned and Ga is incorporated.

Ion beams are not as finely focused as electron beams and, partly for this reason, they generally offer lower resolution. However, the contrast mechanisms for ISE generation are different from those for SE generation and can offer complementary information about a sample surface. SE and ISE images of the same sample are shown in Figure 3. Both the SE (Figure 3a) and the ISE (Figure 3b) images show contrast due to surface topography and material differences. However, ISE imaging typically delivers stronger channeling contrast from crystals than SE imaging. The contrast due to crystal orientation is easily distinguished from material contrast, because crystal contrast changes with the incidence angle of the ion beam and material contrast does not. In ideal samples such as Cu or Au, ISE channeling contrast can reveal twin lamellae as narrow as 20 nm and grains as small as 50 nm.

The different contrast mechanisms are illustrated in Figure 4. A comparison of Figures 4a and 4b shows that when the crystal is oriented so that the ion "channels" along crystal planes, there are fewer ion interactions with sample atoms near the surface and thus fewer electrons are emitted. Figure 4c illustrates that heavier samples typically result in more ISEs (and SEs). Figure 4d shows that surface topography can lead to increases in the number of ISEs (and SEs), because of the increase in the number of ion-solid interactions near the sample surface. The SE and ISE images are also often distinguished by the amount of charging generated in insulating samples.

Presumably because of differences in the low-energy/secondary electron yields and to the fact that the Ga implantation creates a thin conducting layer at the sample surface, the FIB can often be used to image uncoated samples that are difficult to image even with low-voltage SEM. However, it is important to remember that ion beam imaging always results in some Ga implantation and sputtering of the sample surface.



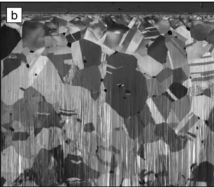


Figure 3. (a) Secondary electron (SE) and (b) ion-induced secondary electron (ISE) images of an FIB-cut cross section in brass. Surface topography generated during the FIB milling and phase contrast are visible in both images. The heavier second-phase precipitates are bright in the SE image and dark in the ISE image.<sup>37</sup> Channeling contrast showing the grain structure is visible only in the ISE image.

# Ion Beam Sputtering

Because of the sputtering action of the ion beam, the FIB can be used to locally remove or mill away material. For example, a Sn sphere is progressively sputtered over a defined area in Figure 5. For direct milling, the limiting feature size is typically about 10 nm<sup>9,19,20</sup> (see the articles by MoberlyChan et al. and Langford et al. in this issue). Quantitative aspects of sputtering are complicated and depend on the material, crystal orientation, ion beam incidence angle, and the extent of redeposition.

As the incidence angle of the ion beam is increased, the intersection of the collision cascade with the sample surface increases, and the number of sputtered atoms per collision cascade increases (a similar effect of geometry on electron emission is shown in Figure 4d). However, at the same time, the fraction of reflected or backscattered Ga ions increases. The combination of these two effects leads to a maximum in sputtering yield at an incidence angle of approximately 75-80°. This effect has been confirmed for 25-30 keV Ga into a variety of materials, including single-crystal Si,<sup>19–22</sup> amorphous SiO<sub>2</sub>,<sup>21,22</sup> and polycrystalline Au and W<sup>21</sup> and shows good agreement between experiment and theory. Si or amorphous solids are ideal for such a study because the effects of crystal channeling are avoided (the surface region of Si amorphizes under the Ga beam<sup>7</sup>). The behavior is more complicated in crystalline material where both incident angle and channeling effects are present.<sup>21</sup>

The sputtering yield at a given incidence angle can vary by as much as a factor of 10 for strongly channeling crystal orientations in materials such as Cu.<sup>23</sup> This is because for easy channeling orienta-

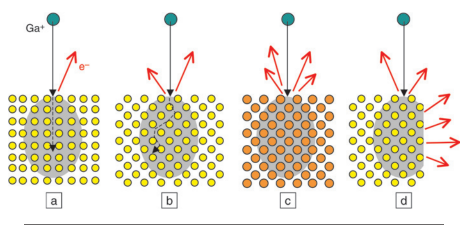
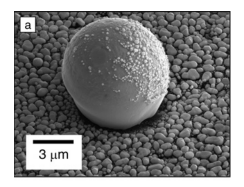
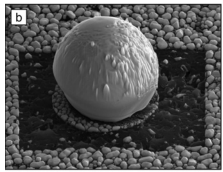


Figure 4. Schematics showing the influence of (a), (b) crystal orientation, (c) atomic mass, and (d) surface geometry on 30 keV Ga<sup>+</sup> collision cascades and ISE image contrast formation. The orange atoms in (c) are more massive than the yellow atoms in (a), (b), and (d). Similar concepts influence sputtering yields.

tions, the ion experiences only inelastic glancing-angle collisions with the atoms lying in a crystal plane and travels deeper into the crystal before causing elastic collisions, so that fewer atoms are sputtered from the surface. This is analogous to the effect of crystal orientation on low-energy electron yields illustrated in Figures 4a and 4b. Channeling effects on sputtering at a vertical grain boundary in Cu are shown in Figure 6. The grain with the smaller electron yield appears dark in Figure 6. Its lower sputter yield results in a shallower sputter profile than that observed for the brighter neighboring grain.

The sputter profiles also depend on the exact sequence in which the ion beam is





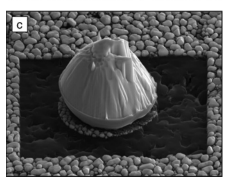


Figure 5. (a)–(c) Secondary electron images during sputtering of a large Sn sphere surrounded by Sn droplets, using a 30 keV Ga beam rastered across a 10  $\mu$ m  $\times$  10  $\mu$ m area.

rastered over the surface.<sup>22</sup> For instance, the sputter profile of a ring cut by rapid and repetitive scanning ("multi-pass" scanning) differs from that obtained by slowly scanning the beam just once over the same area ("single-pass" scanning) (Figure 6). Despite an identical total ion dose (ions per unit area) for both rings in Figure 6, the slow single-pass scan, spiraling from outside to inside, results in a deeper cut, because of effects of ion focusing, incident angle effects, and redeposition.

Redeposition decreases the effective sputter yield and changes sputter profiles. The decreased yield comes about because redeposited material lands in the area being sputtered and must be sputtered a second time. Redeposition is also given as a reason why completely vertical sidewalls cannot be cut with the FIB without over-tilting the sample, 13,24 but it is certainly also partly because of the intensity tails of the ion beam profile and of the decrease in sputter yield at high incidence angles. 19,20,25 Many details of redeposition effects remain open, such as the development of crystal orientation and channeling effects seen in the redeposited material (e.g., Figure 6, single-pass ring).

In addition to redeposition, surface roughening and shadowing effects are prevalent during sputtering. An example of a shadowing effect on topology during cross-sectioning with the FIB, the so-called "curtain effect," is seen in the lower half of the images in Figure 3. Surface roughening, specifically ripple formation, is widespread during ion bombardment 9.26 and is attributed to competition between smoothing by surface diffusion or viscous flow and roughening because of surface-curvature-dependent sputter yields. (The sputter yield depends on local curvature for the same reasons it depends on angle of incidence.)

Even during normal-incidence sputtering, surface roughening can occur and is dependent on the crystal orientation, as shown in the multi-pass ring cut across two differently oriented grains in Figure 6. Such crystal-orientation-dependent rippling is attributed to anisotropic surface diffusion.<sup>27</sup> Other unusual effects of sputtering and redeposition on surface evolution continue to be discovered.<sup>28,29</sup> Nonetheless, progress is being made toward models that can predict surface evolution during sputtering and can be used to achieve desired sputter profiles (see the article by Langford et al. in this issue).

# **Ga Incorporation**

Imaging and milling with Ga ions always result in Ga incorporation near the sample surface. As the sample surface is sputtered away at a rate proportional to the sputtering yield and the ion flux (ions per area per time), the Ga is implanted

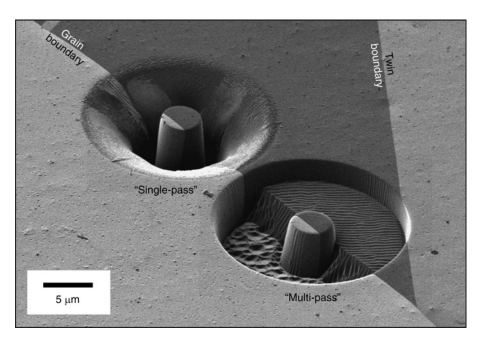


Figure 6. Ion-induced secondary electron image of rings milled with 30 keV Ga at a grain boundary in Cu. The single-pass ring shows an enhanced sputtering yield (deeper trough) and redeposition (sloped, thicker sidewalls); the multi-pass ring shows channeling effects and surface roughening.

further into the sample, and a steady-state profile of Ga is reached. The maximum Ga concentration occurs at steady state after the removal of target material to a depth roughly equal to the ion range and is constant over a depth also roughly equal to the ion range. Ignoring the effects from possible diffusion of Ga in the target material, differences in molar volume, and preferential sputtering, the Ga atom fraction at steady state is

$$f_{Ga} = 1/(\alpha Y), \tag{1}$$

where  $\alpha$  is the fraction of ions that are not reflected or backscattered from the sample surface, Y is the sputtering yield (number of sputtered atoms/ions per incoming ion)<sup>10</sup> and  $\alpha Y$  is the sputtering yield per implanted ion. Based on this equation and typical TRIM sputtering yields (from 1 to 20 sputtered atoms per incoming ion), atom fractions from 1 at.% to 50 at.% Ga are expected near the sample surface. Because of the increase in sputter yield with incident angle, the steady-state Ga concentration is expected to be roughly 5-10 times smaller during glancing-angle sputtering than normalincidence sputtering. In contrast, channeling is expected to raise the steady-state Ga concentration.

Most studies report Ga concentrations in good agreement with simulations and theory, although a few claim to see almost no Ga. Unless the Ga diffuses away or is reflected, it must end up in the surface region of the sample. Some studies were performed with energy-dispersive spectroscopy in an SEM, which for typical electron beam parameters will look through the thin Ga-doped layer at the surface and underestimate the actual Ga concentration. Other studies have used surface Auger analysis (without sputtering) that may also underestimate the Ga concentration, because of the presence of carbonaceous or oxide surface layers. Carbon-based surface layers are particularly prevalent in samples that have been imaged in the SEM after FIB milling.30 In contrast, Auger depth profiles or chemical analysis in the TEM give reasonable agreement with predictions for both Ga concentration and range (allowing for channeling effects).

All of these considerations become much more complicated in alloys or if Ga diffusion or reactions take place. One unusual example is the formation of Gacontaining surface phases during imaging of certain fcc metals (see the article by Mayer et al. in this issue). This phenomenon, which is coupled with sample crystal structure and channeling effects, limits the ability to obtain high-quality images from certain materials.

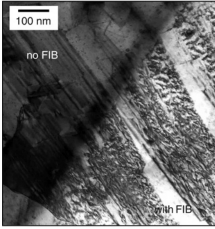
# Ion Beam Damage

A major drawback of FIB imaging and machining, particularly for TEM samples, is the damage created by the ion beam (again, see Mayer et al. in this issue). As the ion dose increases, the individual disordered cascade regions overlap and a damaged surface layer is formed. Depending in particular on the sample material and temperature, the ion beam damage can take the form of sample surface amorphization, point defect creation, dislocation formation, phase formation, grain modification, or other unusual effects. With the exception of Si amorphization, systematic investigations of FIB damage are just beginning. Nonetheless, several trends can be identified based on literature results for broad beam ions and on anecdotal FIB observations.

Ion beam amorphization is a wellknown phenomenon and has been extensively studied for covalently bound materials such as Si, Ge, GaAs, and C (diamond). Because of the highly directional nature of the atomic bonds in covalent materials and in certain alloys, the atomic rearrangements necessary to heal the disorder created by the ion beam are often hindered. In contrast, pure metals have nondirectional bonding and do not amorphize. Thus, thin amorphous layers sometimes observed at the edge of pure metal TEM samples made by FIB presumably contain impurities such as Ga, C, or O. Some oxygen gets in because of the relatively poor quality of the vacuum.

Point defects and dislocation loops can also be created during FIB imaging and machining. Systematic studies of FIBinduced defects have not been undertaken, although there are many isolated observations, both published and unpublished. For example, Cu is prone to extensive FIB damage,<sup>31</sup> as shown in the TEM image in Figure 7, but Al is not and even provides reasonable-quality high-resolution TEM samples.<sup>32</sup> Tooth enamel (hydroxyapatite) is resistant to FIB-induced damage, 33 but other apatite crystals that decompose at lower temperatures amorphize easily. Several additional and unusual types of damage have been observed in FIB-milled samples. These include the formation of Gacontaining surface phases (see Mayer et al., this issue) as well as ion-beaminduced grain growth in fine-grained Ni and Ni alloys.34 Preferential sputtering, which is prevalent during FIB milling of materials that decompose at low tempera-





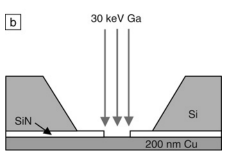


Figure 7. (a) Transmission electron microscopy (TEM) image of a Cu film, showing that the existing dislocation structure was modified by FIB milling. (b) Schematic illustration of the milling geometry. (TEM image courtesy of G.P. Zhang.)

tures, can lead to chemical changes in the surface region and influence the ease of damage formation and amorphization.

# Ion Beam Heating

During ion implantation, almost all of the ion kinetic energy is eventually converted to heat, with only a small fraction stored as defects in the sample or emitted as energetic particles or radiation.14 For times longer than approximately a nanosecond and distances larger than around 100 nm, the ion beam can be approximated as a continuous heat source. At shorter times, there are large temporal variations in heating, and at times of less than  $10^{-12}$  s, the atoms barely have time to interact with each other, and the temperature of the solid is not well defined. 15 The maximum temperature reached in a sample depends on the beam power P, sample thermal conductivity  $\kappa$ , sample geometry, and contact to a heat reservoir. 15,35 Beam powers in commercial systems have maximum values of 1 mW.

When the ion beam is incident on a flat surface, the transfer of heat away from the incidence point is so effective that even in the absence of a heat reservoir (e.g., for a semi-infinite sample), a finite steady-state temperature increase is reached given by<sup>36</sup>

$$T = P/(\pi a \kappa), \tag{2}$$

where a is the radius of the circular ion beam profile on the sample surface. For values of P/a available in commercial FIBs, between 1 W/m and 1000 W/m, the temperature rise predicted by Equation 2 can range from entirely negligible for samples with good thermal conductivity to huge for poor conductors. For example, for Si ( $\kappa = 148 \text{ W/m K}$ ) the temperature increase is <2°C even for the most extreme beam conditions. In contrast, for polymeric or biological materials ( $\kappa \sim 0.1 \text{ W/m}$ K), such small temperature rises are only achieved with P/a values of <2 W/m, which is at the low end of what is available in commercial FIBs.

Beam heating can be diminished by placing samples in good contact with a heat reservoir. On the other hand, when imaging or machining TEM lamellae, membranes, or other structured samples, much higher temperatures may be reached when the sample geometry limits the transfer of heat.<sup>35</sup> An extreme example of this is when the heat transfer is reduced to one dimension, such as for a cylindrical pillar with a diameter equal to the beam diameter. In this case, the temperature rise is given by Equation 2 multiplied by the height-to-diameter ratio of the pillar. Thus, by determining the length-to-width ratio of the thermal path in a structured sample, a worst-case estimate of the actual temperature rise can be obtained and used to select appropriate beam conditions. However, imaging or cutting of highaspect-ratio features in low-conductivity materials may lead to unacceptably high temperatures, even for the most mild beam conditions in commercial systems.

# In This Issue

Four articles follow this introduction and are intended to cover the most widely used FIB applications as well as the current state of the art in FIB materials research. In the first article, by Mayer et al., one of the most important applications of the FIB is presented. The FIB not only enables the preparation of site-specific samples of uniform thickness, but

it also facilitates the fabrication of lamellae from composite samples consisting of materials with very different properties. The strengths of the method as well as the problems, such as FIB-induced damage and Ga contamination, are illustrated with examples. The second article, by Uchic et al., describes how the FIB can be used as a tool for three-dimensional characterization by complementing 2D imaging or mapping with serial sectioning at a submicron level. The third article, by Langford et al., is organized according to the general technological area, such as microelectronics, photonics, microelectromechanical systems, and rapid prototyping. The strengths and limitations of FIB micro- or nanostructuring are illustrated through representative examples, and the article ends with a brief summary and a look to the future of FIB nanostructuring. In the fourth and final article, by MoberlyChan et al., advanced topics such as single-ion implantation, surface morphology during ion-induced erosion, and ion-induced CVD are discussed.

We hope that this issue of MRS Bulletin will inform readers about the fundamental science and current applications of focused ion beam microscopy and micromachining and give them a sense of the future potential of FIB in materials science.

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