In the past three chapters we’ve described the sources, lenses, and detectors that make up a TEM. The only other parts of the instrument you need to know about in detail are those that, if you are not careful, can seriously degrade the quality of the information you generate even if the rest is perfect. These two parts are the holder in which you put your specimen and the vacuum that surrounds it. While there isn’t much you can do to improve the vacuum, beyond buying a better microscope, there is a lot you can do that will degrade the quality of the vacuum in the column and, in doing so, contaminate your specimen. So we’ll tell you a few basics about how the vacuum pumps work, and how the vacuum system is put together. Although the vacuum system is under computer control in most TEMs, you still affect the vacuum by what you put in the microscope. Consequently, you need to know what not to do on those occasions when you might otherwise degrade the vacuum.

The vacuum in the stage of a typical TEM is \(10^{-5}\) Pa, compared with atmospheric pressure of \(10^5\) Pa. It is quite remarkable that we can transfer a specimen into the TEM, reducing the ambient pressure at its surface by 10 orders of magnitude in a matter of a few seconds. This rapid transfer is a testament to the skills of TEM designers, and particularly the construction of the specimen holder and the airlock system. Specimen holders are the physical contact between you and your specimen across this extraordinary vacuum range. You must transmit all the experimental variables that you want to inflict on your specimen by way of the holder. The most basic requirement is that you should be able to move the specimen laterally to look at different areas; to optimize the imaging you should also be able to move the specimen vertically. In addition we’ll describe how you can tilt, rotate, heat, cool, strain, and bias the materials that you are studying. Unfortunately, the holder also transmits vibrations, drift, and contamination to the specimen and may be a source of X-rays that can degrade any analysis that you want to perform. Care of your specimen holders is extremely important since damaged or worn holders reduce the quality of the data generated by the microscope. If you are not careful, a $10,000 holder can easily limit the information generated by a million-dollar TEM.

**8.1 THE VACUUM**

You know already that electrons are strongly scattered by atoms, which accounts for the versatility of TEM, and the need for thin specimens. Strong scattering also occurs in gases and we can’t transmit coherent, controlled electron beams very far through air, so all EMs operate under vacuum. This means that your specimen has to go through an airlock into the TEM. Therefore, you can only control your specimen remotely, not directly, and this makes TEMs more expensive to build. In addition to permitting the electron beam to travel through the instrument undisturbed, the vacuum also plays a role in keeping the specimen clean (or making it dirty). Contamination of the specimen by vacuum-borne contaminants such as hydrocarbons and water vapor can be a problem in many aspects of TEM. Generally, the better the vacuum, the less contamination, but it is the partial pressure of contaminants, not the absolute pressure, which is important. Fortunately, the vacuum systems in most TEMs today are reasonably clean, fully automated and their operation is transparent to the user. Despite this, you should have some understanding of vacuums and how to control them, so this chapter will cover, very superficially, the principles of vacuum systems and pumps.
First of all, a word on units which, as usual, are in disarray. The SI unit of pressure is the pascal; other non-SI units are the torr and the bar. You’ll come across all three units in TEM texts and in manufacturers’ handbooks, so you need to know the conversions.

We’ll mainly use the pascal, but since the torr is still very common terminology, we’ll occasionally put approximate torr values in parentheses to remind you of the conversion. Since we deal with very low pressures, the numbers are small, although we perversely use the expression ‘high vacuum’ for these low pressures. We think of vacuum in terms of rough, low, high, and ultrahigh. A roughing pump gives a pressure between 100 and 0.1 Pa; 0.1–10⁻⁴ Pa is low vacuum and 10⁻⁴–10⁻⁷ Pa is high vacuum (HV). If the pressure is <10⁻⁷ Pa you have an ultrahigh vacuum (UHV). These values are approximate, not standardized definitions. A typical modern TEM has a pressure inside the column of ~1.3 × 10⁻⁵ Pa (10⁻⁷ Torr), which is in the HV range. UHV TEMs operate below 10⁻⁷ Pa and the gun region of an FEG TEM operates at ~10⁻⁹ Pa (10⁻¹¹ Torr). To have an electron beam inside the TEM that is not scattered significantly by the air molecules in the column, the pressure must be <~0.1 Pa. This was achievable with simple mechanical pumps in the early days of the instrument, but there are good reasons to operate at much lower pressures (higher vacuums), for which you need more sophisticated and more expensive apparatus.

Generally we use one type of pump to create a rough vacuum and another type to create the higher vacuum. The TEM is kept permanently under vacuum, unless it’s being repaired or serviced. If you need access to the inside of the column to change specimens, electron sources, or photographic plates, you do this via an airlock system, which can be pumped separately, as we’ll explain later. There are many different kinds of pumps used in TEMs, and you often have a choice when purchasing an instrument. As with most things, you get what you pay for; a clean UHV system is very expensive. We can divide pumps into roughing pumps and HV/UHV pumps, as we’ll now discuss.

### 8.2 ROUGHING PUMPS

The most common roughing pump is a mechanical (rotary) pump in which a belt-driven, eccentrically mounted reciprocating mechanism sucks air through an inlet valve into a chamber and expels it through an exit valve, as shown in Figure 8.1. Such pumps are very reliable, relatively inexpensive, noisy, and dirty and only lower the pressure to ~10⁻¹ Pa (~10⁻³ Torr). Mechanical pumps should be housed outside your TEM room, and connected to the column through a line that doesn’t transmit their vibration. These pumps use a hydrocarbon oil as a medium. If you have such a pump, the line from the pump to the vacuum should contain a foreline trap to condense out oil vapor before it is deposited in the column. Also, the exhaust line from the pump must also be trapped to prevent (possibly carcinogenic) oil vapor escaping into the room where you are working. There are alternative ‘dry’ roughing pumps which do not use oil. These are more expensive and somewhat less reliable; they do not pump to such low pressure.

**PRESSURE**

1 Torr is ~ 130 Pa
1 Pa is 7.5 × 10⁻³ Torr

One bar is atmospheric pressure (~760 Torr) and is equivalent to ~10⁵ Pa.

The name is the torr; the unit is the Torr, but either way the torr is not an accepted SI unit.

100–0.1 Pa (~1–10⁻³ Torr) is a rough vacuum.

0.1–10⁻⁴ Pa (~10⁻³–10⁻⁶ Torr) is low vacuum.

10⁻⁴–10⁻⁷ Pa (~10⁻⁶–10⁻⁹ Torr) is high vacuum (HV).

<10⁻⁷ Pa (~10⁻⁹ Torr) is ultrahigh vacuum (UHV).

Be careful when you hear a phrase like “the vacuum in the gun is 10⁻⁸” and remember the pascal unit is Pa and the torr unit is Torr.

**FIGURE 8.1.** A mechanical pump for roughing vacuums. The eccentric motion of the pump creates a vacuum in the RH side when it rotates and the vacuum sucks air into the inlet valve. As the cylinder rotates further, it cuts off the inlet and forces the air through the outlet on the LH side, creating a vacuum again on the inlet side as it does so. Because of the constant contact between the rotating cylinder and the inside of the pump, oil is needed to reduce frictional heating.
8.3 HIGH/ULTRAHIGH VACUUM PUMPS

8.3.A Diffusion Pumps

These pumps use a hot plate to boil oil, which then forms a series of concentric vapor jets. The jets drag air molecules out of the microscope as shown in Figure 8.2, then condense onto a cold surface, freeing the air molecules which are extracted by the mechanical pump ‘backing’ the diffusion pump. While this may seem an inefficient way to move air, diffusion pumps can in fact transport more than a hundred liters of air per second, which is quite sufficient to pump out a TEM column. With no moving parts, diffusion pumps are inexpensive and very reliable, but they need external water cooling to aid condensation of the vapor. Failure of the cooling water supply and burnout of the hot plate are about the only possible causes of failure. The absence of moving parts ensures vibration-free operation. As with the mechanical pump, the oil diffusion pump would contaminate the vacuum in the TEM if oil vapor were to escape into the column. To minimize this you must use synthetic nonhydrocarbon oils with low vapor pressures, such as Fomblin™ or Santovac™. (Never use a silicone-based oil, of course.) A liquid-N₂ cold trap sits on top of the pump and condenses out any residual oil molecules. If you have diffusion pumps you must keep the cold traps full of liquid N₂ to maintain a clean system.

Diffusion pumps are capable of very efficient pumping from \(10^{-1}\) to \(10^{-9}\) Pa \((10^{-11}\) Torr\) and, if properly trapped, will provide a clean UHV system that is very reliable. The VG series UHV DSTEMs use only oil diffusion pumps to attain UHV conditions.

8.3.B Turbomolecular Pumps

Turbomolecular pumps, or turbopumps, as the name implies, use a turbine to force gases from the microscope. They have many parts moving at high speeds (in excess of 20,000–50,000 rpm is common), so they are more liable to fail than diffusion pumps. The mechanics of the pump are very simple as you can appreciate from Figure 8.3. They do not use oil so they don’t introduce hydrocarbons to contaminate the microscope, and the best models (unlike earlier versions) are very quiet and almost vibration-free. In fact, modern turbopumps are being used to prepump the specimen chamber when this is critical, as in the cryo-transfer technique (see Section 8.10). If you buy a turbopump, make sure to specify that its use will not transmit vibrations to the TEM column, where it would destroy the image resolution. The turbopump can start (slowly) at

![Figure 8.2](image1.png)

**FIGURE 8.2.** Principles of diffusion pump operation. A heater plate at the base of the pump boils synthetic oil. The expansion of the oil vapor on boiling creates a pressure which forces the vapor up the central column and out of several holes. The stream of oil vapor pulls gas molecules out of the top of the pump down to the base where the oil condenses and the air is pumped out of the base by a mechanical backing pump.

![Figure 8.3](image2.png)

**FIGURE 8.3.** A turbopump (with and without its casing), which is nothing more than a small turbine that rotates at high speed. Like a jet turbine it pulls air in at the front end and forces it out of the back. The blades are designed like airfoils to enhance the flow of gas through the system.
ambient pressures, increasing speed as the pressure is lowered, ultimately providing UHV conditions at high enough speeds. It is usual, however, to back the turbopump with a dry mechanical pump.

### 8.3.C Ion Pumps

Ion pumps do not contain oil, so they cannot contaminate the TEM column. They also have no moving parts, relying solely on the ionization process to remove air. The ion pump emits electrons from a cathode. These ions spiral in a magnetic field (see Section 6.3) and ionize air molecules, which are then attracted to the cathode. The energetic gas ions sputter Ti atoms from the cathode and they condense throughout the pump chamber, mainly on the cylindrical anode, trapping gas atoms. Thus ion pumps remove gas atoms in two ways: by chemisorption on the anode surfaces and by electrical attraction to the cathodes. The smaller the ion current between the electrodes, the lower the vacuum, so the pump acts as its own vacuum gauge. Ion pumps are only efficient at high vacuums, so they are usually switched on after a diffusion pump has lowered the pressure to $<10^{-4}$ Pa ($10^{-5}$ Torr). It is common to add ion pumps directly to the stage or gun chambers of TEMs to focus their pumping action on these important regions. Since these pumps are very common on TEMs, we include a diagram (Figure 8.4) showing how they operate.

### 8.3.D Cryogenic (Adsorption) Pumps

As the name implies, cryogenic pumps (cryopumps) rely on liquid N$_2$ to cool molecular sieves with large surface areas. The cold surface efficiently removes air molecules from ambient pressure down to $\sim 10^{-4}$ Pa ($10^{-5}$ Torr). Because they are oil-free, cryopumps are also used to back ion pumps and prevent their accidental contamination through backstreaming from oil-bearing pumps.

We also use cold surfaces to enhance vacuums in the stage of most non-UHV TEMs. Such ‘cold fingers’ or ‘anticontaminators’ provide an alternative site (rather than your specimen) for condensation of residual components in the vacuum.

The same is true if the anticontaminator in your stage is allowed to warm up; then it will degrade the vacuum around your specimen. So you must use another pump such as a diffusion or mechanical pump to remove the air molecules as they are released from captivity. Otherwise, this outgassing will degrade the quality of the vacuum around your specimen, increasing contamination.

### 8.4 THE WHOLE SYSTEM

As shown schematically in Figure 8.5 the modern TEM has at least two separate pumping systems: one that evacuates the column and one that pumps the camera and screen chamber. We pump the camera separately because the film is one of the primary causes of vacuum degradation since outgassing occurs from the emulsion that contains the AgI grains. So this part of the TEM is usually pumped by a combination mechanical/diffusion pump. The stage is often pumped by a separate ion pump, turbopump, or cryopump, or some combination of these. If the instrument has an FEG, then there is a separate UHV pumping system for the gun region, which often consists of several ion pumps. Each part of the vacuum system consists of roughing pumps (mechanical or turbo) that pump out the appropriate part of the microscope to give the vacuum where the HV/UHV pumps can start to operate.

Looking at Figure 8.5, there are three valves, which are now all computer controlled.

- #1 connects the mechanical pump to the column (the roughing valve).
&2 connects the mechanical pump to the bottom of the diffusion pump (the backing valve).
&3 connects the diffusion pump directly to the TEM column (the butterfly valve).

If you're pumping down from atmospheric pressure, you first use the mechanical pump to back out the diffusion pump, till it gets to a low enough pressure so its heater can be safely switched on without oxidizing. So close #1, open #2, and close #3.

When the diffusion pump is warmed up, you rough out the column: open #1, close #2, and #3, until the column is at a low enough pressure that the diffusion pump can be used.

At this point, close #1, open #2, and then #3, so the diffusion pump is open to the TEM and may be continuously backed by the mechanical pump. The better approach incorporates a vacuum reservoir between the mechanical and diffusion pumps. When the reservoir is pumped to < 0.1 Pa, the mechanical pump is closed off and the diffusion pump exhausts into the reservoir. When the pressure builds in the reservoir, the mechanical pump will automatically switch on and lower the pressure.

Similar arrangements work for other pumps; e.g., a diffusion pump may be used to lower the pressure in the stage and gun sufficiently for the ion pumps to be switched on, and so on. In most TEMs the stage and gun have significantly better vacuums than the camera region, so the camera/screen is isolated from the rest of the column by a differential pumping aperture (not shown in Figure 8.5). This aperture often coincides with the BFP of the projector lens, since all the electrons have to pass through it and the DP in the BFP localizes all the electron trajectories close to the optic axis. A similar arrangement exists between the stage and gun in FEG systems to preserve the tip in case of a vacuum leak in the stage.

The advent of high-quality digital recording which will remove the need for film in the camera will do more to improve the quality of vacuums in TEMs than any advances in pumping technology.

8.5 LEAK DETECTION

"Nature abhors a vacuum," as Francois Rabelais said in 1534. That's the reason why the pumps must keep pumping: the TEM leaks. But some leaks are too large for the pumps to handle, and then the instrument performance degrades. If you can't run the electron gun, your TEM is useless. Under these circumstances, you have to find the leak, cure it, and repump the instrument, (this is usually a job for your service engineer) but some labs or older TEMs don't have service contracts. Leak detection involves using a mass spectrometer, which can be put into the pumping lines of the microscope. You then release helium gas close to the various parts of the TEM where you suspect a leak (the stage airlock, which sees a lot of use, is a common point of failure). The small He atoms are relatively easily sucked into the column through any leak and register on the mass spectrometer. When a leak is isolated, the TEM may have to be opened to the atmosphere to permit replacement of the defective part, such as the O-ring seals.
The most common cause of a leak is your specimen holder. The O-ring seal on the shaft of a side-entry holder (see the second half of this chapter) is easily contaminated with dust or a hair since it is continually inserted and extracted from the column, and left on the bench while you pore over it. Never touch the O-ring, make sure it doesn’t dry out, but if it does (because you don’t have someone else looking after the TEM), lubricate it with a very thin film of vacuum grease.

After repairing a leak, when you’ve pumped down again, it is often useful to ‘bake’ the column. Baking means heating the internal surfaces to >100°C (or >150–200°C in UHV TEMs) to boil off residual water vapor and hydrocarbons that may have entered the system when it was down to air. Usually, you can achieve the bakeout by leaving the lenses running without their cooling water (check this very carefully with the manufacturer before proceeding). In some cases, special heating panels are constructed around the column. Baking can also introduce other leaks as the whole system expands and then contracts, so sometimes leak detection and cure is an iterative process. For UHV systems, you must bake to reach the ultimate vacuum, and the higher the temperature the better.

Be wary, however, since sometimes the TEM accessories, such as XEDS and EELS systems, are not designed to be baked to the same high temperature as the column.

8.6 CONTAMINATION: HYDROCARBONS AND WATER VAPOR

As we said right at the start of the chapter, the vacuum (or rather what is still present in the column) can be a source of contamination. Residual hydrocarbons from the pump oil crack under the electron beam. Carbo- naceous material then deposits on your carefully thinned specimen, making it difficult to do sensible high-resolution imaging or microanalysis. So a clean vacuum (one in which the hydrocarbon partial pressure is < \(10^{-9}\text{Pa}\)) is essential. Fortunately, most modern TEMs are relatively contamination-free, particularly if you use synthetic oils and appropriate traps on the pumps. (See also Section 8.12.)

However, even if you’ve paid dearly for a clean vacuum system, contamination often occurs; it comes primarily through the airlock with your specimen. You can minimize this by heating the specimen to >100°C in a heating holder or with a halogen lamp in the prepump chamber, or cooling the specimen to liquid-N\(_2\) temperatures in a cooling holder. It may help if the prepump chamber is pumped with an oil-free pump. More recently, plasma cleaning of the specimen holder and specimen prior to insertion in the TEM has proven a very successful way to ensure a clean specimen (more on this in Section 8.12).

Polymers and biological specimens can easily introduce hydrocarbon contaminants, as they outgas in the vacuum, so it is sensible to cool such specimens (since heating or plasma cleaning destroys them). However, when you cool your specimen, it attracts water vapor which condenses as ice on the surface; so load your specimen first, then cool it down in the TEM before you switch on the beam. A low partial pressure of H\(_2\)O in the vacuum is obviously essential. Also, warm up any cooled specimens in the TEM before bringing them out to ambient atmosphere, otherwise they will immediately ice up (unless it’s a very dry winter’s day). There will be more about this in the sections on specimen holders.

In addition to the specimen, you personally can be a major source of contamination. Take care never to touch anything that will enter the vacuum, i.e., the specimen itself, the grids, specimen holder (beyond the O-ring seal on the rod), clamping rings, replacement diaphragms, new filaments, replacement Wehnelt, components of XEDS and EELS systems, etc. Use latex gloves whenever you load a specimen, and don’t breathe on it. Store specimen holders and specimens in a dry box containing a desiccant such as silica gel, which should be replaced regularly. Always prepump fresh film in a vacuum desic- cator (which is sometimes integrated into the TEM itself but probably best done elsewhere). Better still, never use film if you can avoid it. Simple precautions like this will minimize contamination of your specimens and the microscope in general and bring a much greater return in terms of good data per TEM session.

8.7 SPECIMEN HOLDERS AND STAGES

To look at your specimen, place it in a specimen holder and insert this assembly into the TEM stage. Therefore, there are two key components which are often not separated, namely, the holder and the stage. In this part of the chapter, we will emphasize the holder but the stage is also critical. Suitable design of the stage is the essential precursor to computer-controlled, remotely accessed TEM, which is already happening.

The cold trap, cold finger, or cryo-blades are a critical part of the stage. Ideally, this cold finger will completely surround the specimen: it cryo-pumps the region around the specimen. However, the cold surfaces, usually brass, provide a source of stray electrons and X-rays which is undesirable for AEM (see Chapter 33), so these blades should be removable for AEM.

X-ray diffractometers use goniometers to hold and tilt the specimen; so do TEMs. Conventional SEMs use a stub on which you mount the specimen so that you can bring the specimen close to the objective lens. However, some high-resolution SEMs use a specimen holder which
is very similar to those used in the TEM, because the specimen is inserted inside the lens, rather than underneath and outside it.

The reason the specimen holder is so important in TEM is that your specimen must invariably be located within the objective lens and the aberrations associated with the objective lens determine the resolution of the TEM.

Historically, microscopists have used two different designs and a lot of what you’ll read here or elsewhere has a strong historical background.

- The traditional side-entry holder is a rod with a motor attached to tilt and/or rotate the specimen and a lead connecting it to a power supply and control box, or liquid-N₂ dewar.
- The traditional top-entry holder is a cartridge which you load into the TEM but is detached from the outside world when you use the microscope.

The actual cup that holds your specimen is either 2.3 or 3.05 mm in diameter, so the specimen disk or support grid has to be the same dimension, as we’ll see in Chapter 10. The reasons for these dimensions are again partly historical. In the top-entry holder the specimen and part of the holder fit through the bore of the upper polepiece (see Figures 6.7 and 6.8). Clearly, the specimen must be smaller than the bore diameter. So the original top-entry holders used small specimens.

### The Word ‘Traditional’

Holders are changing; new manufacturers and new capabilities are emerging. Ideally, the holder should not be moved after it is inserted in the TEM.

Side-entry holders are more versatile and larger specimen dimensions first appeared when they were introduced. However, side-entry holders connect the specimen directly to the outside world via a long lever arm, which is undesirable, unstable, and also not necessary in many cases! Ideally, the side-entry holder should leave the specimen in the stage, not connected to the outside world, and all manipulations should be conducted through the stage itself, not the holder. This ideal is being approached as stages become more computer-controlled.

### 8.8 Side-Entry Holders

Side-entry holders are now the standard, although their design has changed quite radically in recent years. The traditional design is shown in Figure 8.6. The key parts of this holder are:

- **The O-ring**, which is one mechanical link to the microscope column. Some holders have two O-rings and the gap between the O-rings is pumped separately to improve the vacuum.
- **The jewel bearing**, which is the other mechanical link to the microscope column. You push on this bearing to move your specimen back and forth and from side to side. Like the O-ring, you must keep the bearing clean otherwise the specimen will not be stable.
- **The cup**, which actually holds your specimen and thus provides the immediate environment which is seen by stray electrons and any X-rays coming down the column. So cups in holders for AEM are made of Be to minimize the generation of X-rays that would interfere with microanalysis.
- **The clamping ring or screw**, which holds the specimen in the cup. This ring (not shown in the figure), which may also be Be, must be carefully designed. It must hold your specimen firmly (so, e.g., magnetic disks cannot be pulled out of the cup by the lens field). However, the ring must not be so difficult to tighten that you put undue pressure on your specimen. Brittle disks may break as you are loading them. There are two kinds of retaining rings: screw-thread rings, which are easier to control and do not damage metals, but you’ll find they may break ceramics because they transfer shear stresses to the disk; spring clips are difficult for the novice to master, but with practice you’ll find they offer more control over the load that you put on the specimen, so we recommend them for the experienced ceramist. Unfortunately, no one we know makes Be spring clips!

In a more modern design, the jewel bearing is omitted so that the holder is supported at just one pivot point.
8.9 TOP-ENTRY HOLDERS

Top-entry holders are becoming less common because they essentially preclude XEDS analysis in the TEM. Also, it is more difficult to design such holders so that the specimen can be manipulated (e.g., rotated or strained). Their great advantage was that they were much less susceptible to drift since they were not connected directly to the outside, so early HRTEM required top-entry holders. Today, however, nearly all TEMs up to 400 kV use side-entry holders; only the DSTEMs retain the top-entry (bottom entry?) design.

Another drawback of such holders is that the bore of the objective lens must be asymmetric (using an upper and a lower polepiece), which actually limits the ultimate resolution by constraining the lens designer. Figure 8.7 shows a schematic diagram of such a holder.

8.10 TILT AND ROTATE HOLDERS

One feature of TEMs which may surprise you if you are a new user is that a wide variety of holders is available for the TEM. Figure 8.8 shows illustrations of these different designs for the side-entry holder:

- **Single-tilt holder**: This is the basic holder with which any novice should start practicing. You can only tilt around the axis of the rod. It is relatively cheap, robust, and can at least give you some idea of the usefulness of tilting a specimen for diffraction-contrast studies.

- **Quick change holder**: This is also a single-tilt holder that clamps the specimen with a lever arm which you raise and lower onto your disk or grid. It doesn’t put a high stress on the specimen, but it doesn’t hold it very strongly either. Don’t use it for magnetic specimens, but it can be great for ceramics. Different retainers can be substituted for the clamp as shown in Figure 8.8 (bottom), creating a more versatile multipurpose holder.

- **Multiple-specimen holder**: This is usually a single-tilt holder, but you can load up to five specimens into the column at one time as shown in Figure 8.9A. A two-specimen, double-tilt version is also available (Figure 8.9B). Such holders can be useful if you are not very good at specimen preparation, or you want to compare different specimens under identical conditions without turning off the beam. However, in

---

**FIGURE 8.7.** Top-entry holder: (A) cross section; (B) top view. The cartridge has a cone shape which fits into the tapered bore of the objective lens polepiece. The specimen sits in a cup at the base of a column through the cone down which the incident beam travels. Simple manipulations such as tilting or rotating require complex micromechanical design, since the specimen is at the base of the cartridge and completely surrounded by the polepiece. To tilt, e.g., as shown in (A), push rods are pressed against springs in two orthogonal directions, displacing a central ring around the column (see B), thus tilting the specimen cup.

**FIGURE 8.8.** Examples of different designs for the side-entry holder. From the top, they are: a rotation holder, a heating holder, a cooling holder, a double-tilt holder, and a single-tilt holder.
modern TEMs, specimen exchange is relatively quick, except in UHV instruments where the multi-
holder would probably be more useful although it is less common.

- **Bulk specimen holder**: This holder is used for surface imaging and diffraction, e.g., using SE or BSE in a STEM or for reflection diffraction and imaging in a TEM (see Chapter 29 for more about these techniques). The bulk specimen is larger than the traditional 3-mm disk (usually ~10 mm x 5 mm) so if you can create a thin specimen of these dimensions, the bulk holder will allow you to sample more of your material at one time (Figure 8.10).

  So don’t always think that you are limited to 3-mm specimens!

- **Double-tilt holder**: This is the most popular holder since it gives you the most flexibility in orienting the specimen. It is absolutely essential for imaging and diffraction studies of crystalline specimens. The tilt axes are fixed as two orthogonal directions. In some designs, you can remove the cup while the specimen is in place which means that you can reinset your specimen in the same orientation. This feature is extremely useful if your specimen is robust.

- **Tilt-rotate holder**: You would often like to be able to orient your specimen parallel to the tilt axis (along the rod). This holder lets you do just that. It’s a strength for the side-entry holder: one tilt axis is always parallel to the rod of the holder which also gives the largest tilt angle.

- **Low-background holder**: The cup and clamping ring are made of Be to minimize the generation of bremsstrahlung X-rays and characteristic X-rays. So they are required for XEDS studies. They can be double or single tilt and may be cooled also.

- **Tomography holder**: This is a new design that allows you to tilt the sample through a full 360°. It’s ideal for looking at needles (like an AFM or atom-probe tip).

### 8.11 IN-SITU HOLDERS

Special holders have been developed that allow you to change your specimen while you observe it in the TEM; in other words you can do experiments (heat, cool, strain, twist, compress, etc.) on a specimen in the TEM.

- **Heating holder**: Such holders in a conventional TEM can go to ~1300°C which is measured by a thermocouple attached to the cup. In HVEMs, the temperature can go higher because of the larger gap between the polepieces. You have to be careful to calibrate the temperature and remember that the temperature may be different for different specimens. You should also be sure that the material you are studying does not form a eutectic alloy with the material forming the holder! If the eutectic does form it will have a lower melting point, so you may deposit part of your specimen and the holder on the objective lens, or down onto the screen, if the microscope is well aligned.

- **Cooling holder**: This is available for either liquid-N₂ or liquid-He temperatures. These holders, which can be single or double tilt, are a great asset for XEDS, EELS, and CBED studies since they minimize surface-borne contamination. They are also essential for in-situ studies of superconducting materials and ideal for polymers or biological tissue. However, you should remember that the cold holder can also act as a small cryo-pump so that it actually attracts contamination.
Since you are necessarily changing the temperature at the specimen relative to its surroundings, be prepared for specimen drift. It takes time for the whole system to stabilize. The Polara from FEI is a modification of the cooling holder which fits in a special dedicated stage; the tip of the holder detaches from the rod with obvious advantages.

- Cryo-transfer holder: Certain specimens are prepared at cryogenic temperatures such as liquids, latex emulsions, and tissue in general. This holder permits you to transfer such cold specimens into the TEM without water vapor from the atmosphere condensing as ice on the surface.

- Straining holder: This holder clamps the specimen at both ends then applies a load to one end, via a load cell or screw-thread mechanism, as shown in Figure 8.11. The sample can be in the shape of a small tensile specimen and it is thinned in the middle of the gauge length (see inset). The motion of dislocations, cracks, etc., are then easily monitored, so a video camera is an essential accessory. You can vary the load, to study cyclic as well as tensile loading, and the strain rate is another variable that is easily controlled. In Figure 8.11 a furnace is present, so the specimen can be heated while under load. The use of piezoelectric drives is leading to great improvements in this type of holder.

- Probing holders: These are like AFM holders for the TEM. You can use them to ‘poke’ your specimens—just like an AFM, STM, or an indenter—while you observe the effect in the TEM.

- EBIC and CL holders: The essential feature is the electrical feed-through that allows you to control the charge recombination in a semiconductor or certain mineral specimens by applying a bias across the specimen surface.

Beware: heating and straining holders, in particular, can produce effects in thin foils that are totally uncharacteristic of your bulk sample. So you must use these holders carefully and interpret your results cautiously. Often, surface reactions will dominate internal reactions when you are trying to induce a phase transformation by heating. The surface may also stop grain boundaries from migrating at temperatures where they would do so in the bulk material. Obviously, defect motion under applied stress may also be strongly affected since the 3D stress field will be very different in bulk specimens compared to thin foils.

These problems can be overcome to some extent if you use thicker specimens and examine them in an HVEM, or at least an IVEM, and the whole field of in situ studies, particularly, heating and/or straining, is best performed in such microscopes (Butler and Hale 1981 and Section 29.12). However, the high-energy electrons in these microscopes may introduce lattice defects that affect the very phenomenon that you want to study, e.g., beam-induced vacancies can change diffusional phase-transformation kinetics very easily.

It is also possible, but much more difficult and expensive, to manipulate specimens in top-entry stages. The top-entry holder shown in Figure 8.12 is a heating-
straining holder, which is capable of operating at temperatures up to 2300 K. The heat is provided by a coaxial Ta tube that supports the W heater filament as shown in the figure. The holder is used in a 3-MV microscope where the specimen diameter is 5 mm. The larger specimen diameter means that the disk can be shaped as a small tensile specimen and still be quite robust.

There are also special combinations of holders and stages which have been optimized for particular applications. The example shown in Figure 8.13 has been optimized to combine surface studies using low-energy electron diffraction (LEED) and Auger analysis with TEM. The prechamber is fitted with an ion gun to clean the sample before the surface is analyzed. The specimen can then be moved into the TEM column for transmission studies. A similar prechamber has been

FIGURE 8.13. Schematic diagram showing the Hitachi H900 UHV TEM. This instrument is equipped with a prechamber with LEED, Auger, and an ion gun which can be used to clean the specimen, allowing UHV surface analysis to be carried out on the TEM specimen. The holder has to transfer the specimen through a prepump chamber where it is ion-cleaned before going into the column.

FIGURE 8.14. Plasma cleaner: (A) example and (B) schematic.

FIGURE 8.15. Cleaning the surface of the specimen using a plasma cleaner reduces the contamination produced by a focused electron beam. (A) Set 1 was produced before the specimen was cleaned. Sets 2 and 3 (which you can’t see) were produced after 5 minutes of Ar cleaning and then an additional 5 minutes of oxygen cleaning, respectively. (B) The rate at which the specimen contamination builds up; the additional cleaning in pure oxygen always reduced the contamination rate.
used elsewhere to provide a method to clean the sample before growing thin films on the sample by molecular-beam epitaxy (MBE) or thermal evaporation.

One of the reasons for using higher accelerating voltages is that this gives more room in the specimen-stage region. Thus, even 400-kV microscopes can be fitted with a small, differentially pumped environmental chamber. Such a chamber allows in situ studies of corrosion, degradation of catalysts, etc., especially when combined with a heating holder.

Development of new holders and their use for in-situ studies is going to be one of the most exciting topics in TEM. We’ll expand on this topic in the companion text.

8.12 Plasma Cleaners

Plasma cleaners have been available for removing surface contamination and modifying surfaces (for example, changing wettability) for over 30 years. The unit is now marketed as a small box (e.g., as shown in Figure 8.14A) that can usually accommodate one specimen holder; the rod of the holder is placed inside a plasma chamber, as shown in Figure 8.14B, just before it goes into the TEM. Plasma cleaners have long been used to modify/clean surfaces of glass, semiconductors and other ceramics, metals, and even polymers and biomaterials. The development of plasma cleaners for cleaning both the TEM holder (the part that goes into the vacuum of the TEM) and more specifically the specimen that has already been placed in the TEM holder, is more recent, but the possibilities are clear. This cleaning is generally regarded as essential for small-probe AEM. Most of the contamination that occurs in the electron beam originates on your specimen, not from a ‘dirty’ vacuum.

Although the processes occurring in the plasma are complex (like the plasma itself) the basic idea is illustrated by the removal of hydrocarbon contamination on the specimen shown in Figure 8.15A. The plasma consists of a mixture of energetic electrons and ions that bombard the surface and break the C–H bonds. With short duration exposure, the surface of the specimen itself is essentially unaffected. The hydrocarbon is thus gradually reduced in molecular weight and pumped away in the vacuum of the cleaner. The user has some flexibility in the choice of the plasma gas although the manufacturer usually limits this. Although many gasses can be used in principle, O₂, N₂, and Ar are the most common. The graph in Figure 8.15B shows the effect of using an oxygen plasma—it reduces the rate at which the contamination grows under the electron probe; the effect of the oxygen-reactive gas is similar to that found when using iodine in the ion mill.
HOLDERS AND IN SITU


THE COMPANION TEXT

A full chapter on holders is included in the companion text with particular attention being paid to holders for in-situ experiments.

SELF-ASSESSMENT QUESTIONS

Q8.1 How does a diffusion pump work?
Q8.2 How does a mechanical pump work?
Q8.3 How does a turbomolecular pump work?
Q8.4 How does an ion pump work?
Q8.5 How does a cryogenic pump work?
Q8.6 If you are concerned about contaminants in the TEM from pumps, which pump(s) should you use (consider where and why)?
Q8.7 Where does the most common vacuum leak occur in a TEM and why at this point?
Q8.8 Name one way to localize the source of a leak.
Q8.9 Which types of specimens introduce contaminants and how can such introduction be prevented?
Q8.10 How can you personally introduce contamination into the microscope and what should you do to stop doing it?
Q8.11 Summarize the advantages and disadvantages of side-entry holders.
Q8.12 Why are TEM top-entry holders out of fashion?
Q8.13 What is a high vacuum (give a number with SI units)?
Q8.14 What is an ultrahigh vacuum and why is it worth paying a lot of money to buy a TEM with such a vacuum?
Q8.15 Which type of holder is the most flexible for TEM?
Q8.16 Why is the column baked? Why should this be done carefully?
Q8.17 What size is the typical specimen cup in the holder?
Q8.18 Why don’t we make larger-diameter thin samples?
Q8.19 What are the drawbacks when using a straining holder?
Q8.20 Why would you cool a specimen?
Q8.21 What do we sometimes deliberately insert into a TEM that creates even more contamination and vacuum degradation than the specimen?

TEXT-SPECIFIC QUESTIONS

T8.1 How would you notice that there is a vacuum leak in the TEM (without watching the gauges)?
T8.2 Why can baking both improve the vacuum and cause more leaks?
T8.3 Explain with a schematic diagram (like Figure 8.5) how the combination of low and high vacuum pumps is used in tandem to pump down the microscope after: (a) it has been let down to air (e.g., for repair); (b) after it has been degraded by the insertion of a specimen; (c) after it has been degraded by changing the film.
T8.4 Copy Figure 8.1 and cut out the circle attached to the rotating vane. By placing that circle and the vane in different positions within the pump chamber show how air is alternately pulled out of the microscope (into the inlet port) then expelled from the pump (through the outlet port).
T8.5 Estimate the number of atoms in a cubic millimeter of air at $10^{-3}$ and $10^{-6}$ bar and thus estimate the number of Ti atoms that need to be ionized to reduce the vacuum in that cubic millimeter by that amount in each case. State any assumptions and be brief.
T8.6 Compare and contrast top-entry and side-entry holders from the points of view of stability (mechanical, thermal, etc.) versatility, cost, etc.
T8.7 Under what circumstances might you prefer to use a holder with just one tilt axis but the ability to rotate the specimen?

T8.8 Why do we choose to use Be for analytical holders rather than a lighter metal, or a heavier one such as Mg? What are the potential problems with using Be?

T8.9 What are the problems with manufacturing a stage and holders for UHV operations rather than for the generally poorer vacuum in most TEMs?

T8.10 Why is it difficult to manufacture holders that operate at extremely high or very low temperatures that do not compromise the routine operation of the microscope?