When we use an analytical method to solve a problem, there is no guarantee that will obtain accurate or precise results. In designing an analytical method we consider potential sources of determinate error and indeterminate error, and we take appropriate steps—such as reagent blanks and the calibration of instruments—to minimize their effect. Why might a carefully designed analytical method give poor results? One possible reason is that we may have failed to account for errors associated with the sample. If we collect the wrong sample, or if we lose analyte when we prepare the sample for analysis, then we introduce a determinate source of error. If we fail to collect enough samples, or if we collect samples of the wrong size, then the precision of our analysis may suffer. In this chapter we consider how to collect samples and how to prepare them for analysis.
**7A The Importance of Sampling**

When a manufacturer lists a chemical as ACS Reagent Grade, they must demonstrate that it conforms to specifications set by the American Chemical Society (ACS). For example, the ACS specifications for commercial NaBr require that the concentration of iron is less than 5 ppm. To verify that a production lot meets this standard, the manufacturer collects and analyzes several samples, reporting the average result on the product’s label (Figure 7.1).

If the individual samples do not represent accurately the population from which they are drawn—a population that we call the **target population**—then even a careful analysis will yield an inaccurate result. Extrapolating a result from a sample to its target population always introduces a determinate sampling error. To minimize this determinate sampling error, we must collect the right sample.

Even if we collect the right sample, indeterminate sampling errors may limit the usefulness of our analysis. Equation 7.1 shows that a confidence interval about the mean, $\bar{X}$, is proportional to the standard deviation, $s$, of the analysis

$$\mu = \bar{X} \pm \frac{ts}{\sqrt{n}}$$  \hspace{1cm}  (7.1)

where $n$ is the number of samples and $t$ is a statistical factor that accounts for the probability that the confidence interval contains the true value, $\mu$.

Each step of an analysis contributes random error that affects the overall standard deviation. For convenience, let’s divide an analysis into two steps—collecting the samples and analyzing the samples—each of which is characterized by a variance. Using a propagation of uncertainty, the relationship between the overall variance, $s^2$, and the variances due to sampling, $s_{\text{amp}}^2$, and the variance due to the analytical method, $s_{\text{meth}}^2$, is

$$s^2 = s_{\text{amp}}^2 + s_{\text{meth}}^2$$  \hspace{1cm}  (7.2)

Equation 7.2 shows that the overall variance for an analysis is limited by either the analytical method or sampling, or by both. Unfortunately, analysts often try to minimize the overall variance by improving only the method’s precision. This is a futile effort, however, if the standard deviation for sampling is more than three times greater than that for the method.¹

Figure 7.2 shows how the ratio $s_{\text{amp}}/s_{\text{meth}}$ affects the method’s contribution to the overall variance. As shown by the dashed line, if the sample’s standard deviation is $3 \times$ the method’s standard deviation, then indeterminate method errors explain only 10% of the overall variance. If indeterminate sampling errors are significant, decreasing $s_{\text{meth}}$ provides only limited improvement in the overall precision.

---

Example 7.1

A quantitative analysis gives a mean concentration of 12.6 ppm for an analyte. The method’s standard deviation is 1.1 ppm and the standard deviation for sampling is 2.1 ppm. (a) What is the overall variance for the analysis? (b) By how much does the overall variance change if we improve \( s_{\text{meth}} \) by 10% to 0.99 ppm? (c) By how much does the overall variance change if we improve \( s_{\text{amp}} \) by 10% to 1.9 ppm?

**Solution**

(a) The overall variance is

\[
s^2 = s_{\text{amp}}^2 + s_{\text{meth}}^2 = (2.1 \text{ ppm})^2 + (1.1 \text{ ppm})^2 = 5.6 \text{ ppm}^2
\]

(b) Improving the method’s standard deviation changes the overall variance to

\[
s^2 = (2.1 \text{ ppm})^2 + (0.99 \text{ ppm})^2 = 5.4 \text{ ppm}^2
\]

Improving the method’s standard deviation by 10% improves the overall variance by approximately 4%.

(c) Changing the standard deviation for sampling

\[
s^2 = (1.9 \text{ ppm})^2 + (1.1 \text{ ppm})^2 = 4.8 \text{ ppm}^2
\]

improves the overall variance by almost 15%. As expected, because \( s_{\text{amp}} \) is larger than \( s_{\text{meth}} \), we achieve a bigger improvement in the overall variance when we focus our attention on sampling problems.

To determine which step has the greatest effect on the overall variance, we need to measure both \( s_{\text{amp}} \) and \( s_{\text{meth}} \). The analysis of replicate samples provides an estimate of the overall variance. To determine the method’s variance we must analyze samples under conditions where we can assume

**Practice Exercise 7.1**

Suppose you wish to reduce the overall variance in Example 7.1 to 5.0 ppm\(^2\). If you focus on the method, by what percentage do you need to reduce \( s_{\text{meth}} \)? If you focus on the sampling, by what percentage do you need to reduce \( s_{\text{amp}} \)?

Click [here](#) to review your answer to this exercise.
that the sampling variance is negligible; the sampling variance is determined by difference.

Example 7.2

The following data were collected as part of a study to determine the effect of sampling variance on the analysis of drug-animal feed formulations.\(^2\)

<table>
<thead>
<tr>
<th>% Drug (w/w)</th>
<th>% Drug (w/w)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0114</td>
<td>0.0105</td>
</tr>
<tr>
<td>0.0102</td>
<td>0.0087</td>
</tr>
<tr>
<td>0.0100</td>
<td>0.0098</td>
</tr>
<tr>
<td>0.0105</td>
<td>0.0095</td>
</tr>
</tbody>
</table>

The data on the left were obtained under conditions where both \(s_{\text{samp}}\) and \(s_{\text{meth}}\) contribute to the overall variance. The data on the right were obtained under conditions where \(s_{\text{samp}}\) is insignificant. Determine the overall variance, and the standard deviations due to sampling and the analytical method. To which source of indeterminate error—sampling or the method—should we turn our attention if we want to improve the precision of the analysis?

**Solution**

Using the data on the left, the overall variance, \(s^2\), is \(4.71 \times 10^{-7}\). To find the method’s contribution to the overall variance, \(s_{\text{meth}}^2\), we use the data on the right, obtaining a value of \(7.00 \times 10^{-8}\). The variance due to sampling, \(s_{\text{samp}}^2\), is

\[
\begin{align*}
  s_{\text{samp}}^2 &= s^2 - s_{\text{meth}}^2 \\
  &= 4.71 \times 10^{-7} - 7.00 \times 10^{-8} \\
  &= 4.01 \times 10^{-7}
\end{align*}
\]

Converting variances to standard deviations gives \(s_{\text{samp}}\) as \(6.33 \times 10^{-4}\) and \(s_{\text{meth}}\) as \(2.65 \times 10^{-4}\). Because \(s_{\text{samp}}\) is more than twice as large as \(s_{\text{meth}}\), improving the precision of the sampling process will have the greatest impact on the overall precision.

---


See Chapter 4 for a review of how to calculate the variance.

---

Practice Exercise 7.2

A polymer’s density provides a measure of its crystallinity. The standard deviation for the determination of density using a single sample of a polymer is \(1.96 \times 10^{-3}\) g/cm\(^3\). The standard deviation when using different samples of the polymer is \(3.65 \times 10^{-2}\) g/cm\(^3\). Determine the standard deviations due to sampling and to the analytical method.

Click [here](#) to review your answer to this exercise.
Chapter 7 Collecting and Preparing Samples

7B Designing A Sampling Plan

A sampling plan must support the goals of an analysis. For example, a material scientist interested in characterizing a metal’s surface chemistry is more likely to choose a freshly exposed surface, created by cleaving the sample under vacuum, than a surface previously exposed to the atmosphere. In a qualitative analysis, a sample need not be identical to the original substance provided there is sufficient analyte present to ensure its detection. In fact, if the goal of an analysis is to identify a trace-level component, it may be desirable to discriminate against major components when collecting samples.

For a quantitative analysis, the sample’s composition must represent accurately the target population, a requirement that necessitates a careful sampling plan. Among the issues we need to consider are these five questions.

1. From where within the target population should we collect samples?
2. What type of samples should we collect?
3. What is the minimum amount of sample needed for each analysis?
4. How many samples should we analyze?
5. How can we minimize the overall variance for the analysis?

7B.1 Where to Sample the Target Population

A sampling error occurs whenever a sample’s composition is not identical to its target population. If the target population is homogeneous, then we can collect individual samples without giving consideration to where we collect sample. Unfortunately, in most situations the target population is heterogeneous and attention to where we collect samples is important. For example, due to settling a medication available as an oral suspension may have a higher concentration of its active ingredients at the bottom of the container. The composition of a clinical sample, such as blood or urine, may depend on when it is collected. A patient’s blood glucose level, for instance, will change in response to eating and exercise. Other target populations show both a spatial and a temporal heterogeneity. The concentration of dissolved O\textsubscript{2} in a lake is heterogeneous due both to a change in seasons and to point sources of pollution.

If the analyte’s distribution within the target population is a concern, then our sampling plan must take this into account. When feasible, homogenizing the target population is a simple solution, although this often is impracticable. In addition, homogenizing a sample destroys information about the analyte’s spatial or temporal distribution within the target population, information that may be of importance.

Random Sampling

The ideal sampling plan provides an unbiased estimate of the target population’s properties. A **Random Sampling** is the easiest way to satisfy this requirement. Despite its apparent simplicity, a truly random sample is difficult to collect. Haphazard sampling, in which samples are collected without a sampling plan, is not random and may reflect an analyst’s unintentional biases.

Here is a simple method to ensure that we collect random samples. First, we divide the target population into equal units and assign to each unit a unique number. Then, we use a random number table to select the units to sample. Example 7.3 provides an illustrative example.

**Example 7.3**

To analyze a polymer’s tensile strength, individual samples of the polymer are held between two clamps and stretched. To evaluate a production lot, the manufacturer’s sampling plan calls for collecting ten 1 cm × 1 cm samples from a 100 cm × 100 cm polymer sheet. Explain how we can use a random number table to ensure that we collect these samples at random.

**Solution**

As shown by the grid to the left, we divide the polymer sheet into 10000 1 cm × 1 cm squares, each identified by its row number and its column number, with numbers running from 0 to 99. For example, the blue square is in row 98 and in column 1. To select ten squares at random, we enter the random number table in Appendix 14 at an arbitrary point and let the entry’s last four digits represent the row number and the column number for the first sample. We then move through the table in a predetermined fashion, selecting random numbers until we have 10 samples. For our first sample, let’s use the second entry in the third column of Appendix 14, which is 76831. The first sample, therefore, is row 68 and column 31. If we proceed by moving down the third column, then the 10 samples are as follows:

<table>
<thead>
<tr>
<th>Sample</th>
<th>Number</th>
<th>Row</th>
<th>Column</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>76831</td>
<td>68</td>
<td>31</td>
</tr>
<tr>
<td>2</td>
<td>66558</td>
<td>65</td>
<td>58</td>
</tr>
<tr>
<td>3</td>
<td>33266</td>
<td>32</td>
<td>66</td>
</tr>
<tr>
<td>4</td>
<td>12032</td>
<td>20</td>
<td>32</td>
</tr>
<tr>
<td>5</td>
<td>14063</td>
<td>40</td>
<td>63</td>
</tr>
<tr>
<td>6</td>
<td>41701</td>
<td>17</td>
<td>01</td>
</tr>
<tr>
<td>7</td>
<td>38605</td>
<td>86</td>
<td>05</td>
</tr>
<tr>
<td>8</td>
<td>64516</td>
<td>45</td>
<td>16</td>
</tr>
<tr>
<td>9</td>
<td>13015</td>
<td>30</td>
<td>15</td>
</tr>
<tr>
<td>10</td>
<td>12138</td>
<td>21</td>
<td>38</td>
</tr>
</tbody>
</table>

When we collect a random sample we make no assumptions about the target population, which makes this the least biased approach to sampling. On the other hand, a random sample often requires more time and expense.

---

than other sampling strategies because we need to collect a greater number of samples to ensure that we adequately sample the target population, particularly when that population is heterogenous.\(^4\)

**Judgmental Sampling**

The opposite of random sampling is selective, or **judgmental sampling** in which we use prior information about the target population to help guide our selection of samples. Judgmental sampling is more biased than random sampling, but requires fewer samples. Judgmental sampling is useful if we wish to limit the number of independent variables that might affect our results. For example, if we are studying the bioaccumulation of PCB’s in fish, we may choose to exclude fish that are too small, too young, or that appear diseased.

**Systematic Sampling**

Random sampling and judgmental sampling represent extremes in bias and in the number of samples needed to characterize the target population. **Systematic sampling** falls in between these extremes. In systematic sampling we sample the target population at regular intervals in space or time. Figure 7.3 shows an aerial photo of the Great Salt Lake in Utah. A railroad line divides the lake into two sections that have different chemical compositions. To compare the lake’s two sections—and to evaluate spatial variations within each section—we use a two-dimensional grid to define sampling


**Figure 7.3** Aerial photo of the Great Salt Lake in Utah, taken from the International Space Station at a distance of approximately 380 km. The railroad line divides the lake into two sections that differ in chemical composition. Superimposing a two-dimensional grid divides each section of the lake into sampling units. The red dots at the center of each unit represent sampling sites. Photo courtesy of the Image Science and Analysis Laboratory, NASA Johnson Space Center, Photo Number ISS007-E-13002 (eol.jsc.nasa.gov).
locations, collecting samples at the center of each location. When a population's is heterogeneous in time, as is common in clinical and environmental studies, then we might choose to collect samples at regular intervals in time.

If a target population's properties have a periodic trend, a systematic sampling will lead to a significant bias if our sampling frequency is too small. This is a common problem when sampling electronic signals where the problem is known as aliasing. Consider, for example, a signal that is a simple sign wave. Figure 7.4a shows how an insufficient sampling frequency underestimates the signal's true frequency. The apparent signal, shown by the dashed red line that passes through the five data points, is significantly different from the true signal shown by the solid blue line.

According to the Nyquist theorem, to determine accurately the frequency of a periodic signal, we must sample the signal at least twice during each cycle or period. If we collect samples at an interval of \( \Delta t \), then the highest frequency we can monitor accurately is \( (2\Delta t)^{-1} \). For example, if we collect one sample each hour, then the highest frequency we can monitor is \( (2 \times 1 \text{ hr})^{-1} \) or \( 0.5 \text{ hr}^{-1} \), a period of less than 2 hr. If our signal's period is less than 2 hours (a frequency of more than \( 0.5 \text{ hr}^{-1} \)), then we must use a faster sampling rate. Ideally, we use a sampling rate that is at least 3-4 times greater than the highest frequency signal of interest. If our signal has a period of one hour, then we should collect a new sample every 15-20 minutes.

**Systematic–Judgmental Sampling**

Combinations of the three primary approaches to sampling also are possible. One such combination is **SYSTEMATIC–JUDGMENTAL SAMPLING**, in which we use prior knowledge about a system to guide a systematic sampling plan. For example, when monitoring waste leaching from a landfill, we expect the plume to move in the same direction as the flow of groundwater—this helps focus our sampling, saving money and time. The systematic–judgmental sampling plan in Figure 7.5 includes a rectangular grid for most of the samples and linear transects to explore the plume's limits.

---

**Figure 7.4** Effect of sampling frequency when monitoring a periodic signal. Individual samples are shown by the red dots (•). In (a) the sampling frequency is approximately 1.5 samples per period. The dashed red line shows the apparent signal based on five samples and the solid blue line shows the true signal. In (b) a sampling frequency of approximately 5 samples per period accurately reproduces the true signal.

**Figure 7.5** Systematic–judgmental sampling plan for monitoring the leaching of pollutants from a landfill. The sampling sites, shown as red dots, are on a systematic grid that straddles the direction of the groundwater's flow. Sampling along the linear transects that extend out from the grid help establish the plume's limits.

---

---

---
**STRATIFIED SAMPLING**

Another combination of the three primary approaches to sampling is judgmental—random, or **STRATIFIED SAMPLING**. Many target populations consist of distinct units, or strata. For example, suppose we are studying particulate Pb in urban air. Because particulates come in a range of sizes—some visible and some microscopic—and come from many sources—such as road dust, diesel soot, and fly ash to name a few—we can subdivide the target population by size or by source. If we choose a random sampling plan, then we collect samples without considering the different strata, which may bias the sample toward larger particulates. In a stratified sampling we divide the target population into strata and collect random samples from within each stratum. After we analyze the samples from each stratum, we pool their respective means to give an overall mean for the target population. The advantage of stratified sampling is that individual strata usually are more homogeneous than the target population. The overall sampling variance for stratified sampling always is at least as good, and often is better than that obtained by simple random sampling. Because a stratified sampling requires that we collect and analyze samples from several strata, it often requires more time and money.

**CONVENIENCE SAMPLING**

One additional method of sampling deserves mention. In **CONVENIENCE SAMPLING** we select sample sites using criteria other than minimizing sampling error and sampling variance. In a survey of rural groundwater quality, for example, we can choose to drill wells at sites selected at random or we can choose to take advantage of existing wells; the latter usually is the preferred choice. In this case cost, expedience, and accessibility are more important than ensuring a random sample.

**7B.2 What Type of Sample to Collect**

Having determined from where to collect samples, the next step in designing a sampling plan is to decide on the type of sample to collect. There are three common methods for obtaining samples: grab sampling, composite sampling, and in situ sampling.

The most common type of sample is a **GRAB SAMPLE** in which we collect a portion of the target population at a specific time or location, providing a “snapshot” of the target population. If our target population is homogeneous, a series of random grab samples allows us to establish its properties. For a heterogeneous target population, systematic grab sampling allows us to characterize how its properties change over time and/or space.

A **COMPOSITE SAMPLE** is a set of grab samples that we combine into a single sample before analysis. Because information is lost when we combine individual samples, normally we analyze separately each grab sample. In
some situations, however, there are advantages to working with a composite sample.

One situation where composite sampling is appropriate is when our interest is in the target population’s average composition over time or space. For example, wastewater treatment plants must monitor and report the average daily composition of the treated water they release to the environment. The analyst can collect and analyze a set of individual grab samples and report the average result, or she can save time and money by combining the grab samples into a single composite sample and report the result of her analysis of the composite sample.

Composite sampling also is useful when a single sample does not supply sufficient material for the analysis. For example, analytical methods for the quantitative analysis of PCB’s in fish often require as much as 50 g of tissue, an amount that may be difficult to obtain from a single fish. Combining and homogenizing tissue samples from several fish makes it easy to obtain the necessary 50-g sample.

A significant disadvantage of grab samples and composite samples is that we cannot use them to monitor continuously a time-dependent change in the target population. *In situ sampling*, in which we insert an analytical sensor into the target population, allows us to monitor the target population without removing individual grab samples. For example, we can monitor the pH of a solution in an industrial production line by immersing a pH electrode in the solution’s flow.

**Example 7.4**

A study of the relationship between traffic density and the concentrations of Pb, Cd, and Zn in roadside soils uses the following sampling plan. Samples of surface soil (0–10 cm) are collected at distances of 1, 5, 10, 20, and 30 m from the road. At each distance, 10 samples are taken from different locations and mixed to form a single sample. What type of sampling plan is this? Explain why this is an appropriate sampling plan.

**Solution**

This is a systematic–judgemental sampling plan using composite samples. These are good choices given the goals of the study. Automobile emissions release particulates that contain elevated concentrations of Pb, Cd, and Zn—this study was conducted in Uganda where leaded gasoline was still in use—which settle out on the surrounding roadside soils as “dry rain.” Samples collected near the road and samples collected at fixed distances from the road provide sufficient data for the study, while minimizing the total number of samples. Combining samples from the same distance into a single, composite sample has the advantage of decreasing sampling uncertainty.

---

7B.3 How Much Sample to Collect

To minimize sampling errors, samples must be of an appropriate size. If a sample is too small, its composition may differ substantially from that of the target population, which introduces a sampling error. Samples that are too large, however, require more time and money to collect and analyze, without providing a significant improvement in the sampling error.

Let’s assume our target population is a homogeneous mixture of two types of particles. Particles of type $A$ contain a fixed concentration of analyte, and particles of type $B$ are analyte-free. Samples from this target population follow a binomial distribution. If we collect a sample of $n$ particles, then the expected number of particles that contains analyte, $n_A$, is

$$n_A = np$$

where $p$ is the probability of selecting a particle of type $A$. The standard deviation for sampling is

$$s_{amp} = \sqrt{np(1 - p)}$$

To calculate the relative standard deviation for sampling, $(s_{amp})_{rel}$, we divide equation 7.3 by $n_A$, obtaining

$$(s_{amp})_{rel} = \frac{\sqrt{np(1 - p)}}{np}$$

Solving for $n$ allows us to calculate the number of particles we need to provide a desired relative sampling variance.

$$n = \frac{1 - p}{p} \times \frac{1}{(s_{amp})_{rel}^2}$$

Example 7.5

Suppose we are analyzing a soil where the particles that contain analyte represent only $1 \times 10^{-7}$% of the population. How many particles must we collect to give a percent relative standard deviation for sampling of 1%?

Solution

Since the particles of interest account for $1 \times 10^{-7}$% of all particles, the probability, $p$, of selecting one of these particles is $1 \times 10^{-9}$. Substituting into equation 7.4 gives

$$n = \frac{1 - (1 \times 10^{-9})}{1 \times 10^{-9}} \times \frac{1}{(0.01)^2} = 1 \times 10^{13}$$

To obtain a relative standard deviation for sampling of 1%, we need to collect $1 \times 10^{13}$ particles.

Depending on the particle size, a sample of $10^{13}$ particles may be fairly large. Suppose this is equivalent to a mass of 80 g. Working with a sample this large clearly is not practical. Does this mean we must work with a
smaller sample and accept a larger relative standard deviation for sampling? Fortunately the answer is no. An important feature of equation 7.4 is that the relative standard deviation for sampling is a function of the number of particles instead of their combined mass. If we crush and grind the particles to make them smaller, then a sample of $10^{13}$ particles will have a smaller mass. If we assume that a particle is spherical, then its mass is proportional to the cube of its radius.

$$\text{mass } \propto r^3$$

If we decrease a particle’s radius by a factor of 2, for example, then we decrease its mass by a factor of $2^3$, or 8.

### Example 7.6

Assume that a sample of $10^{13}$ particles from Example 7.5 weighs 80 g and that the particles are spherical. By how much must we reduce a particle’s radius if we wish to work with 0.6-g samples?

**Solution**

To reduce the sample’s mass from 80 g to 0.6 g, we must change its mass by a factor of

$$\frac{80}{0.6} = 133 \times$$

To accomplish this we must decrease a particle’s radius by a factor of

$$r^3 = 133 \times$$

$$r = 5.1 \times$$

Decreasing the radius by a factor of approximately 5 allows us to decrease the sample’s mass from 80 g to 0.6 g.

Treating a population as though it contains only two types of particles is a useful exercise because it shows us that we can improve the relative standard deviation for sampling by collecting more particles. Of course, a real population likely contains more than two types of particles, with the analyte present at several levels of concentration. Nevertheless, the sampling of many well-mixed populations approximate binomial sampling statistics because they are homogeneous on the scale at which they are sampled. Under these conditions the following relationship between the mass of a random grab sample, $m$, and the percent relative standard deviation for sampling, $R$, often is valid

$$mR^2 = K$$  \hspace{1cm} 7.5

where $K$ is a sampling constant equal to the mass of a sample that produces a percent relative standard deviation for sampling of ±1%.8

---

Example 7.7

The following data were obtained in a preliminary determination of the amount of inorganic ash in a breakfast cereal.

<table>
<thead>
<tr>
<th>Mass of Cereal (g)</th>
<th>0.9956</th>
<th>0.9981</th>
<th>1.0036</th>
<th>0.9994</th>
<th>1.0067</th>
</tr>
</thead>
<tbody>
<tr>
<td>% w/w Ash</td>
<td>1.34</td>
<td>1.29</td>
<td>1.32</td>
<td>1.26</td>
<td>1.28</td>
</tr>
</tbody>
</table>

What is the value of $K_s$ and what size sample is needed to give a percent relative standard deviation for sampling of $\pm 2.0\%$? Predict the percent relative standard deviation and the absolute standard deviation if we collect 5.00-g samples.

**Solution**

To determine the sampling constant, $K_s$, we need to know the average mass of the cereal samples and the relative standard deviation for the amount of ash in those samples. The average mass of the cereal samples is 1.0007 g. The average % w/w ash and its absolute standard deviation are, respectively, 1.298% w/w and 0.03194% w/w. The percent relative standard deviation, $R$, therefore, is

$$R = \frac{s_{\text{amp}}}{X} = \frac{0.03194\% \text{ w/w}}{1.298\% \text{ w/w}} \times 100 = 2.46\%$$

Solving for $K_s$ gives its value as

$$K_s = mR^2 = (1.007 \text{ g}) (2.46)^2 = 6.09 \text{ g}$$

To obtain a percent relative standard deviation of $\pm 2\%$, samples must have a mass of at least

$$m = \frac{K_s}{R^2} = \frac{6.09 \text{ g}}{(2.0)^2} = 1.5 \text{ g}$$

If we use 5.00-g samples, then the expected percent relative standard deviation is

$$R = \sqrt{\frac{K_s}{m}} = \sqrt{\frac{6.09 \text{ g}}{5.00 \text{ g}}} = 1.10\%$$

and the expected absolute standard deviation is

$$s_{\text{amp}} = \frac{KX}{100} = \frac{(1.10)(1.298\% \text{ w/w})}{100} = 0.0143\% \text{ w/w}$$

7B.4 How Many Samples to Collect

In the previous section we considered how much sample we need to minimize the standard deviation due to sampling. Another important consideration is the number of samples to collect. If the results from our analysis of the samples are normally distributed, then the confidence interval for the sampling error is
**Practice Exercise 7.3**

Olaquindox is a synthetic growth promoter in medicated feeds for pigs. In an analysis of a production lot of feed, five samples with nominal masses of 0.95 g were collected and analyzed, with the results shown in the following table.

<table>
<thead>
<tr>
<th>mass (g)</th>
<th>0.9530</th>
<th>0.9728</th>
<th>0.9660</th>
<th>0.9402</th>
<th>0.9576</th>
</tr>
</thead>
<tbody>
<tr>
<td>mg olaquindox/kg feed</td>
<td>23.0</td>
<td>23.8</td>
<td>21.0</td>
<td>26.5</td>
<td>21.4</td>
</tr>
</tbody>
</table>

What is the value of $K_s$ and what size samples are needed to obtain a percent relative deviation for sampling of 5.0%? By how much do you need to reduce the average particle size if samples must weigh no more than 1 g?

Click [here](#) to review your answer to this exercise.

$$
\mu = \bar{X} \pm \frac{t_{samp}}{\sqrt{n_{samp}}} \qquad 7.6
$$

where $n_{samp}$ is the number of samples and $s_{samp}$ is the standard deviation for sampling. Rearranging equation 7.6 and substituting $e$ for the quantity $X - \mu$, gives the number of samples as

$$
n_{samp} = \frac{t^2 s_{samp}^2}{e^2} \qquad 7.7
$$

Because the value of $t$ depends on $n_{samp}$, the solution to equation 7.7 is found iteratively.

**Example 7.8**

In Example 7.7 we determined that we need 1.5-g samples to establish an $s_{samp}$ of ±2.0% for the amount of inorganic ash in cereal. How many 1.5-g samples do we need to collect to obtain a percent relative sampling error of ±0.80% at the 95% confidence level?

**Solution**

Because the value of $t$ depends on the number of samples—a result we have yet to calculate—we begin by letting $n_{samp} = \infty$ and using $t(0.05, \infty)$ for $t$. From Appendix 4, the value for $t(0.05, \infty)$ is 1.960. Substituting known values into equation 7.7 gives the number of samples as

$$
n_{samp} = \frac{(1.960)^2 (2.0)^2}{(0.80)^2} = 24.0 \approx 24
$$

Letting $n_{samp} = 24$, the value of $t(0.05, 23)$ from Appendix 4 is 2.073. Recalculating $n_{samp}$ gives

$$
n_{samp} = \frac{(2.073)^2 (2.0)^2}{(0.80)^2} = 26.9 \approx 27
$$
When \( n_{\text{samp}} = 27 \), the value of \( t(0.05, 26) \) from Appendix 4 is 2.060. Recalculating \( n_{\text{samp}} \) gives

\[
n_{\text{samp}} = \frac{(2.060)^2 (2.0)^2}{(0.80)^2} = 26.52 \approx 27
\]

Because two successive calculations give the same value for \( n_{\text{samp}} \), we have an iterative solution to the problem. We need 27 samples to achieve a percent relative sampling error of \( \pm 0.80\% \) at the 95% confidence level.

### Practice Exercise 7.4

Assuming that the percent relative standard deviation for sampling in the determination of olaquindox in medicated feed is 5.0% (see Practice Exercise 7.3), how many samples do we need to analyze to obtain a percent relative sampling error of \( \pm 2.5\% \) at \( \alpha = 0.05 \)?

Click [here](#) to review your answer to this exercise.

---

**Equation 7.7** provides an estimate for the smallest number of samples that will produce the desired sampling error. The actual sampling error may be substantially larger if \( s_{\text{samp}} \) for the samples we collect during the subsequent analysis is greater than \( s_{\text{samp}} \) used to calculate \( n_{\text{samp}} \). This is not an uncommon problem. For a target population with a relative sampling variance of 50 and a desired relative sampling error of \( \pm 5\% \), **equation 7.7** predicts that 10 samples are sufficient. In a simulation using 1000 samples of size 10, however, only 57% of the trials resulted in a sampling error of less than \( \pm 5\% \). Increasing the number of samples to 17 was sufficient to ensure that the desired sampling error was achieved 95% of the time.

### 7B.5 Minimizing the Overall Variance

A final consideration when we develop a sampling plan is how we can minimize the overall variance for the analysis. **Equation 7.2** shows that the overall variance is a function of the variance due to the method, \( s_{\text{meth}}^2 \), and the variance due to sampling, \( s_{\text{samp}}^2 \). As we learned earlier, we can improve the sampling variance by collecting more samples of the proper size. Increasing the number of times we analyze each sample improves the method’s variance. If \( s_{\text{samp}}^2 \) is significantly greater than \( s_{\text{meth}}^2 \), we can ignore the method’s contribution to the overall variance and use **equation 7.7** to estimate the number of samples to analyze. Analyzing any sample more than once will not improve the overall variance, because the method’s variance is insignificant.

If \( s_{\text{meth}}^2 \) is significantly greater than \( s_{\text{samp}}^2 \), then we need to collect and analyze only one sample. The number of replicate analyses, \( n_{\text{rep}} \), we need to minimize the error due to the method is given by an equation similar to **equation 7.7**.

---

For an interesting discussion of why the number of samples is important, see Kaplan, D.; Lacetera, N.; Kaplan, C. “Sample Size and Precision in NIH Peer Review,” Plos One, 2008, 3(7), 1–3. When reviewing grants, individual reviewers report a score between 1.0 and 5.0 (two significant figure). NIH reports the average score to three significant figures, implying that a difference of 0.01 is significant. If the individual scores have a standard deviation of 0.1, then a difference of 0.01 is significant at \( \alpha = 0.05 \) only if there are 384 reviews. The authors conclude that NIH review panels are too small to provide a statistically meaningful separation between proposals receiving similar scores.

---

Unfortunately, the simple situations described above often are the exception. For many analyses, both the sampling variance and the method variance are significant, and both multiple samples and replicate analyses of each sample are necessary. The overall error in this case is

\[ e = t \sqrt{\frac{s_{samp}^2}{n_{samp}} + \frac{s_{meth}^2}{n_{samp} n_{rep}}} \]

Equation 7.8 does not have a unique solution as different combinations of \( n_{samp} \) and \( n_{rep} \) give the same overall error. How many samples we collect and how many times we analyze each sample is determined by other concerns, such as the cost of collecting and analyzing samples, and the amount of available sample.

**Example 7.9**

An analytical method has a relative sampling variance of 0.40% and a relative method variance of 0.070%. Evaluate the percent relative error (\( \alpha = 0.05 \)) if you collect 5 samples and analyze each twice, and if you collect 2 samples and analyze each 5 times.

**Solution**

Both sampling strategies require a total of 10 analyses. From Appendix 4 we find that the value of \( t(0.05, 9) \) is 2.262. Using equation 7.8, the relative error for the first sampling strategy is

\[ e = 2.262 \sqrt{\frac{0.40}{5} + \frac{0.070}{5 \times 2}} = 0.67\% \]

and that for the second sampling strategy is

\[ e = 2.262 \sqrt{\frac{0.40}{2} + \frac{0.070}{2 \times 5}} = 1.0\% \]

Because the method variance is smaller than the sampling variance, we obtain a smaller relative error if we collect more samples and analyze each sample fewer times.

**Practice Exercise 7.5**

An analytical method has a relative sampling variance of 0.10% and a relative method variance of 0.20%. The cost of collecting a sample is $20 and the cost of analyzing a sample is $50. Propose a sampling strategy that provides a maximum relative error of \( \pm 0.50\% \) (\( \alpha = 0.05 \)) and a maximum cost of $700.

Click [here](#) to review your answer to this exercise.
Chapter 7 Collecting and Preparing Samples

Implementing a sampling plan usually involves three steps: physically removing the sample from its target population, preserving the sample, and preparing the sample for analysis. Except for in situ sampling, we analyze a sample after we have removed it from its target population. Because sampling exposes the target population to potential contamination, our sampling device must be inert and clean.

Once we remove a sample from its target population, there is a danger that it will undergo a chemical or physical change before we can complete its analysis. This is a serious problem because the sample’s properties will no longer be representative of the target population. To prevent this problem, we often preserve samples before we transport them to the laboratory for analysis. Even when we analyze a sample in the field, preservation may still be necessary.

The initial sample is called the primary or gross sample, and it may be a single increment drawn from the target population or a composite of several increments. In many cases we cannot analyze the gross sample without first preparing the sample for analysis by reducing the sample’s particle size, by converting the sample into a more readily analyzable form, or by improving its homogeneity.

7C.1 Solutions

There are many good examples of solution samples: commercial solvents; beverages, such as milk or fruit juice; natural waters, including lakes, streams, seawater, and rain; bodily fluids, such as blood and urine; and, suspensions, such as those found in many oral medications. Let’s use the sampling of natural waters and wastewaters as a case study in how to sample a solution.

Sample Collection

The chemical composition of a surface water—such as a stream, river, lake, estuary, or ocean—is influenced by flow rate and depth. Rapidly flowing shallow streams and rivers, and shallow (<5 m) lakes usually are well mixed and show little stratification with depth. To collect a grab sample we submerge a capped bottle below the surface, remove the cap and allow the bottle to fill completely, and replace the cap. Collecting a sample this way avoids the air–water interface, which may be enriched with heavy metals or contaminated with oil.\(^\text{10}\)

Slowly moving streams and rivers, lakes deeper than five meters, estuaries, and oceans may show substantial stratification with depth. Grab samples from near the surface are collected as described above, and samples at greater depths are collected using a sample bottle lowered to the desired depth (Figure 7.6).

Wells for sampling groundwater are purged before we collect samples because the chemical composition of water in a well-casing may differ significantly from that of the groundwater. These differences may result from contaminants introduced while drilling the well or by a change in the groundwater’s redox potential following its exposure to atmospheric oxygen. In general, a well is purged by pumping out a volume of water equivalent to several well-casing volumes or by pumping until the water’s temperature, pH, or specific conductance is constant. A municipal water supply, such as a residence or a business, is purged before sampling because the chemical composition of water standing in a pipe may differ significantly from the treated water supply. Samples are collected at faucets after flushing the pipes for 2-3 minutes.

Samples from municipal wastewater treatment plants and industrial discharges often are collected as a 24-hour composite. An automatic sampler periodically removes an individual grab sample, adding it to those collected previously. The volume of each sample and the frequency of sampling may be constant, or may vary in response to changes in flow rate.

Sample containers for collecting natural waters and wastewaters are made from glass or plastic. Kimax and Pyrex brand borosilicate glass have the advantage of being easy to sterilize, easy to clean, and inert to all solutions except those that are strongly alkaline. The disadvantages of glass containers are cost, weight, and the ease of breakage. Plastic containers are made from a variety of polymers, including polyethylene, polypropylene, polycarbonate, polyvinyl chloride, and Teflon. Plastic containers are lightweight, durable, and, except for those manufactured from Teflon, inexpensive. In most cases glass or plastic bottles are used interchangeably, although polyethylene bottles generally are preferred because of their lower cost. Glass containers are always used when collecting samples for the analysis of pesticides, oil and grease, and organics because these species often interact with plastic surfaces. Because glass surfaces easily adsorb metal ions, plastic bottles are preferred when collecting samples for the analysis of trace metals.

In most cases the sample bottle has a wide mouth, which makes it easy to fill and to remove the sample. A narrow-mouth sample bottle is used if exposing the sample to the container’s cap or to the outside environment is a problem. Unless exposure to plastic is a problem, caps for sample bottles are manufactured from polyethylene. When polyethylene must be avoided, the container’s cap includes an inert interior liner of neoprene or Teflon.

**Sample Preservation and Preparation**

After removing a sample from its target population, its chemical composition may change as a result of chemical, biological, or physical processes. To prevent a change in composition, samples are preserved by controlling the sample’s pH and temperature, by limiting its exposure to light or to the atmosphere, or by adding a chemical preservative. After preserving a sample, it is safely stored for later analysis. The maximum holding time
between preservation and analysis depends on the analyte’s stability and the effectiveness of sample preservation. Table 7.1 summarizes preservation methods and maximum holding times for several analytes of importance in the analysis of natural waters and wastewaters.

Other than adding a preservative, solution samples generally do not need additional preparation before analysis. This is the case for samples of natural waters and wastewaters. Solution samples with particularly complex matrices—blood and milk are two common examples—may need additional processing to separate analytes from interferents, a topic covered later in this chapter.

### 7C.2 Gases

Typical examples of gaseous samples include automobile exhaust, emissions from industrial smokestacks, atmospheric gases, and compressed gases. Also included in this category are aerosol particulates—the fine solid particles and liquid droplets that form smoke and smog. Let’s use the sampling of urban air as a case study in how to sample a gas.

### Sample Collection

One approach for collecting a sample of urban air is to fill a stainless steel canister or a Tedlar/Teflon bag. A pump pulls the air into the container and, after purging, the container is sealed. This method has the advantage of being simple and of collecting a representative sample. Disadvantages include the tendency for some analytes to adsorb to the container’s walls, the presence of analytes at concentrations too low to detect with suitable accuracy and precision, and the presence of reactive analytes, such as ozone and nitrogen oxides, that may react with the container or that may otherwise alter the sample’s chemical composition during storage. When using a

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Preservation Method</th>
<th>Maximum Holding Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>ammonia</td>
<td>cool to 4°C; add H₂SO₄ to pH &lt; 2</td>
<td>28 days</td>
</tr>
<tr>
<td>chloride</td>
<td>none required</td>
<td>28 days</td>
</tr>
<tr>
<td>metals—Cr(VI)</td>
<td>cool to 4°C</td>
<td>24 hours</td>
</tr>
<tr>
<td>metals—Hg</td>
<td>HNO₃ to pH &lt; 2</td>
<td>28 days</td>
</tr>
<tr>
<td>metals—all others</td>
<td>HNO₃ to pH &lt; 2</td>
<td>6 months</td>
</tr>
<tr>
<td>nitrate</td>
<td>none required</td>
<td>48 hours</td>
</tr>
<tr>
<td>organochlorine pesticides</td>
<td>1 mL of 10 mg/mL HgCl₂ or immediate extraction with a suitable non-aqueous solvent</td>
<td>7 days without extraction 40 days with extraction</td>
</tr>
<tr>
<td>pH</td>
<td>none required</td>
<td>analyze immediately</td>
</tr>
</tbody>
</table>
stainless steel canister, cryogenic cooling, which changes the sample from a gaseous state to a liquid state, may limit some of these disadvantages.

Most urban air samples are collected by filtration or by using a trap that contains a solid sorbent. Solid sorbents are used for volatile gases (a vapor pressure more than $10^{-6}$ atm) and for semi-volatile gases (a vapor pressure between $10^{-6}$ atm and $10^{-12}$ atm). Filtration is used to collect aerosol particulates. Trapping and filtering allow for sampling larger volumes of gas—an important concern for an analyte with a small concentration—and stabilizes the sample between its collection and its analysis.

In solid sorbent sampling, a pump pulls the urban air through a canister packed with sorbent particles. Typically 2–100 L of air are sampled when collecting a volatile compound and 2–500 m$^3$ when collecting a semi-volatile gas. A variety of inorganic, organic polymer, and carbon sorbents have been used. Inorganic sorbents, such as silica gel, alumina, magnesium aluminum silicate, and molecular sieves, are efficient collectors for polar compounds. Their efficiency at absorbing water, however, limits their capacity for many organic analytes.

Organic polymeric sorbents include polymeric resins of 2,4-diphenyl-$p$-phenylene oxide or styrene-divinylbenzene for volatile compounds, and polyurethane foam for semi-volatile compounds. These materials have a low affinity for water and are efficient for sampling all but the most highly volatile organic compounds and some lower molecular weight alcohols and ketones. Carbon sorbents are superior to organic polymer resins, which makes them useful for highly volatile organic compounds that will not absorb onto polymeric resins, although removing the compounds may be difficult.

Non-volatile compounds normally are present either as solid particulates or as bound to solid particulates. Samples are collected by pulling a large volume of urban air through a filtering unit and collecting the particulates on glass fiber filters.

The short term exposure of humans, animals, and plants to atmospheric pollutants is more severe than that for pollutants in other matrices. Because the composition of atmospheric gases can vary significantly over a time, the continuous monitoring of atmospheric gases such as O$_3$, CO, SO$_2$, NH$_3$, H$_2$O$_2$, and NO$_2$ by in situ sampling is important.$^{11}$

**Sample Preservation and Preparation**

After collecting a gross sample of urban air, generally there is little need for sample preservation or preparation. The chemical composition of a gas sample usually is stable when it is collected using a solid sorbent, a filter, or by cryogenic cooling. When using a solid sorbent, gaseous compounds are released for analysis by thermal desorption or by extracting with a suitable

---


1 m$^3$ is equivalent to $10^3$ L.
solvent. If the sorbent is selective for a single analyte, the increase in the sorbent’s mass is used to determine the amount of analyte in the sample.

7C.3 Solids

Typical examples of solid samples include large particulates, such as those found in ores; smaller particulates, such as soils and sediments; tablets, pellets, and capsules used for dispensing pharmaceutical products and animal feeds; sheet materials, such as polymers and rolled metals; and tissue samples from biological specimens. Solids usually are heterogeneous and we must collect samples carefully if they are to be representative of the target population. Let’s use the sampling of sediments, soils, and ores as a case study in how to sample solids.

Sample Collection

Sediments from the bottom of streams, rivers, lakes, estuaries, and oceans are collected with a bottom grab sampler or with a corer. A bottom grab sampler (Figure 7.7) is equipped with a pair of jaws that close when they contact the sediment, scooping up sediment in the process. Its principal advantages are ease of use and the ability to collect a large sample. Disadvantages include the tendency to lose finer grain sediment particles as water flows out of the sampler, and the loss of spatial information—both laterally and with depth—due to mixing of the sample.

An alternative method for collecting sediments is the cylindrical coring device shown in Figure 7.8. The corer is dropped into the sediment, collecting a column of sediment and the water in contact with the sediment. With the possible exception of sediment at the surface, which may experience mixing, samples collected with a corer maintain their vertical
profile, which preserves information about how the sediment’s composition changes with depth.

Collecting soil samples at depths of up to 30 cm is accomplished with a scoop or a shovel, although the sampling variance generally is high. A better tool for collecting soil samples near the surface is a soil punch, which is a thin-walled steel tube that retains a core sample after it is pushed into the soil and removed. Soil samples from depths greater than 30 cm are collected by digging a trench and collecting lateral samples with a soil punch. Alternatively, an auger is used to drill a hole to the desired depth and the sample collected with a soil punch.

For particulate materials, particle size often determines the sampling method. Larger particulate solids, such as ores, are sampled using a riffle (Figure 7.9), which is a trough with an even number of compartments. Because adjoining compartments empty onto opposite sides of the riffle, dumping a gross sample into the riffle divides it in half. By repeatedly passing half of the separated material back through the riffle, a sample of the desired size is collected.

A sample thief (Figure 7.10) is used for sampling smaller particulate materials, such as powders. A typical sample thief consists of two tubes that are nestled together. Each tube has one or more slots aligned down the length of the sample thief. Before inserting the sample thief into the material being sampled, the slots are closed by rotating the inner tube. When the sample thief is in place, rotating the inner tube opens the slots, which fill with individual samples. The inner tube is then rotated to the closed position and the sample thief withdrawn.

Figure 7.9 Example of a four-unit riffle. Passing the gross sample, shown within the circle, through the riffle divides it into four piles, two on each side. Combining the piles from one side of the riffle provides a new sample, which is passed through the riffle again or kept as the final sample. The piles from the other side of the riffle are discarded.
**Sample Preservation**

Without preservation, a solid sample may undergo a change in composition due to the loss of volatile material, biodegradation, or chemical reactivity (particularly redox reactions). Storing samples at lower temperatures makes them less prone to biodegradation and to the loss of volatile material, but fracturing of solids and phase separations may present problems. To minimize the loss of volatile compounds, the sample container is filled completely, eliminating a headspace where gases collect. Samples that have not been exposed to $O_2$ particularly are susceptible to oxidation reactions. For example, samples of anaerobic sediments must be prevented from coming into contact with air.

**Sample Preparation**

Unlike gases and liquids, which generally require little sample preparation, a solid sample usually needs some processing before analysis. There are two reasons for this. First, as discussed in section 7B.3, the standard deviation for sampling, $s_{samp}$, is a function of the number of particles in the sample, not the combined mass of the particles. For a heterogeneous material that consists of large particulates, the gross sample may be too large to analyze. For example, a Ni-bearing ore with an average particle size of 5 mm may require a sample that weighs one ton to obtain a reasonable $s_{samp}$. Reducing the sample’s average particle size allows us to collect the same number of particles with a smaller, more manageable mass. Second, many analytical techniques require that the analyte be in solution.

**Reducing Particle Size**

A reduction in particle size is accomplished by crushing and grinding the gross sample. The resulting particulates are then thoroughly mixed and divided into subsamples of smaller mass. This process seldom occurs in a single step. Instead, subsamples are cycled through the process several times until a final laboratory sample is obtained.

Crushing and grinding uses mechanical force to break larger particles into smaller particles. A variety of tools are used depending on the particle’s size and hardness. Large particles are crushed using jaw crushers that can reduce particles to diameters of a few millimeters. Ball mills, disk mills, and mortars and pestles are used to further reduce particle size.

A significant change in the gross sample’s composition may occur during crushing and grinding. Decreasing particle size increases the available surface area, which increases the risk of losing volatile components. This problem is made worse by the frictional heat that accompanies crushing and grinding. Increasing the surface area also exposes interior portions of the sample to the atmosphere where oxidation may alter the gross sample’s composition. Other problems include contamination from the materials used to crush and grind the sample, and differences in the ease with which
particles are reduced in size. For example, softer particles are easier to reduce in size and may be lost as dust before the remaining sample is processed. This is a particular problem if the analyte’s distribution between different types of particles is not uniform.

The gross sample is reduced to a uniform particle size by intermittently passing it through a sieve. Those particles not passing through the sieve receive additional processing until the entire sample is of uniform size. The resulting material is mixed thoroughly to ensure homogeneity and a subsample obtained with a riffle, or by CONING AND QUARTERING. As shown in Figure 7.11, the gross sample is piled into a cone, flattened, and divided into four quarters. After discarding two diagonally opposed quarters, the remaining material is cycled through the process of coning and quartering until a suitable laboratory sample remains.

**Bringing Solid Samples Into Solution**

If you are fortunate, your sample will dissolve easily in a suitable solvent, requiring no more effort than gently swirling and heating. Distilled water usually is the solvent of choice for inorganic salts, but organic solvents, such as methanol, chloroform, and toluene, are useful for organic materials.

When a sample is difficult to dissolve, the next step is to try digesting it with an acid or a base. Table 7.2 lists several common acids and bases, and summarizes their use. Digestions are carried out in an open container, usually a beaker, using a hot-plate as a source of heat. The main advantage of an open-vessel digestion is cost because it requires no special equipment.

![Figure 7.11 Illustration showing the method of coning and quartering for reducing sample size. After gathering the gross sample into a cone, the cone is flattened, divided in half, and then divided into quarters. Two opposing quarters are combined to form the laboratory sample or a subsample that is sent through another cycle. The two remaining quarters are discarded.](image-url)
Volatile reaction products, however, are lost, which results in a determinate error if they include the analyte.

Many digestions now are carried out in a closed container using microwave radiation as the source of energy. Vessels for microwave digestion are manufactured using Teflon (or some other fluoropolymer) or fused silica. Both materials are thermally stable, chemically resistant, transparent to microwave radiation, and capable of withstanding elevated pressures. A typical microwave digestion vessel, as shown in Figure 7.12, consists of an insulated vessel body and a cap with a pressure relief valve. The vessels are placed in a microwave oven (a typical oven can accommodate 6–14 vessels) and microwave energy is controlled by monitoring the temperature or pressure within one of the vessels.

A microwave digestion has several important advantages over an open-vessel digestion, including higher temperatures (200–300 °C) and pressures (40–100 bar). As a result, digestions that require several hours in an open-vessel may need less than 30 minutes when using a microwave digestion. In addition, a closed container prevents the loss of volatile gases. Disadvantages include the inability to add reagents during the digestion, limitations on the sample’s size (typically <1 g), and safety concerns due to the use of high pressures and the use of corrosive reagents.

Inorganic samples that resist decomposition by digesting with acids or bases often are brought into solution by fusing with a large excess of an alkali metal salt, called a flux. After mixing the sample and the flux in a crucible, they are heated to a molten state and allowed to cool slowly to room

<table>
<thead>
<tr>
<th>Table 7.2</th>
<th>Acids and Bases Used for Digesting Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Solution</td>
<td>Uses and Properties</td>
</tr>
</tbody>
</table>
| HCl (37% w/w) | • dissolves metals more easily reduced than H₂ (E° < 0)  
• dissolves insoluble carbonate, sulfides, phosphates, fluorides, sulfates, and many oxides |
| HNO₃ (70% w/w) | • strong oxidizing agent  
• dissolves most common metals except Al, Au, Pt, and Cr  
• decomposes organics and biological samples (wet ashing) |
| H₂SO₄ (98% w/w) | • dissolves many metals and alloys  
• decomposes organics by oxidation and dehydration |
| HF (50% w/w) | • dissolves silicates by forming volatile SiF₄ |
| HClO₄ (70% w/w) | • hot, concentrated solutions are strong oxidizing agents  
• dissolves many metals and alloys  
• decomposes organics (Caution: reactions with organics often are explosive; use only in a specially equipped hood with a blast shield and after prior decomposition with HNO₃) |
| HCl:HNO₃ (3:1 v/v) | • also known as aqua regia  
• dissolves Au and Pt |
| NaOH | • dissolves Al and amphoteric oxides of Sn, Pb, Zn, and Cr |
temperature. The resulting melt usually dissolves readily in distilled water or dilute acid. Table 7.3 summarizes several common fluxes and their uses. Fusion works when other methods of decomposition do not because of the high temperature and the flux’s high concentration in the molten liquid. Disadvantages include contamination from the flux and the crucible, and the loss of volatile materials.

Finally, we can decompose organic materials by dry ashing. In this method the sample is placed in a suitable crucible and heated over a flame or in a furnace. The carbon present in the sample oxidizes to CO$_2$, and hydrogen, sulfur, and nitrogen are volatilized as H$_2$O, SO$_2$, and N$_2$. These gases can be trapped and weighed to determine their concentration in the organic material. Often the goal of dry ashing is to remove the organic material, leaving behind an inorganic residue, or ash, that can be further analyzed.

![Figure 7.12 Microwave digestion unit: (a) view of the unit’s interior showing the carousel that holds the digestion vessels; (b) close-up of a Teflon digestion vessel, which is encased in a thermal sleeve. The pressure relief value, which is part of the vessel’s cap, contains a membrane that ruptures if the internal pressure becomes too high.](image)

<table>
<thead>
<tr>
<th>Flux</th>
<th>Melting Temperature ($^\circ$C)</th>
<th>Crucible</th>
<th>Typical Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na$_2$CO$_3$</td>
<td>851</td>
<td>Pt</td>
<td>silicates, oxides, phosphates, sulfides</td>
</tr>
<tr>
<td>Li$_2$B$_4$O$_7$</td>
<td>930</td>
<td>Pt, graphite</td>
<td>aluminosilicates, carbonates</td>
</tr>
<tr>
<td>LiBO$_2$</td>
<td>845</td>
<td>Pt, graphite</td>
<td>aluminosilicates, carbonates</td>
</tr>
<tr>
<td>NaOH</td>
<td>318</td>
<td>Au, Ag</td>
<td>silicates, silicon carbide</td>
</tr>
<tr>
<td>KOH</td>
<td>380</td>
<td>Au, Ag</td>
<td>silicates, silicon carbide</td>
</tr>
<tr>
<td>Na$_2$O$_2$</td>
<td>—</td>
<td>Ni</td>
<td>silicates, chromium steels, Pt alloys</td>
</tr>
<tr>
<td>K$_2$S$_2$O$_7$</td>
<td>300</td>
<td>Ni, porcelain</td>
<td>oxides</td>
</tr>
<tr>
<td>B$_2$O$_3$</td>
<td>577</td>
<td>Pt</td>
<td>silicates, oxides</td>
</tr>
</tbody>
</table>
7D Separating the Analyte from Interferents

When an analytical method is selective for the analyte, analyzing a sample is a relatively simple task. For example, a quantitative analysis for glucose in honey is relatively easy to accomplish if the method is selective for glucose, even in the presence of other reducing sugars, such as fructose. Unfortunately, few analytical methods are selective toward a single species.

In the absence of an interferent, the relationship between the sample’s signal, $S_{\text{samp}}$, and the analyte’s concentration, $C_A$, is

$$S_{\text{samp}} = k_A C_A$$  \hspace{1cm} (7.9)

where $k_A$ is the analyte’s sensitivity. If an interferent, is present, then equation 7.9 becomes

$$S_{\text{samp}} = k_A C_A + k_I C_I$$  \hspace{1cm} (7.10)

where $k_I$ and $C_I$ are, respectively, the interferent’s sensitivity and concentration. A method’s selectivity for the analyte is determined by the relative difference in its sensitivity toward the analyte and the interferent. If $k_A$ is greater than $k_I$, then the method is more selective for the analyte. The method is more selective for the interferent if $k_I$ is greater than $k_A$.

Even if a method is more selective for an interferent, we can use it to determine $C_A$ if the interferent’s contribution to $S_{\text{samp}}$ is insignificant. The selectivity coefficient, $K_{A,I}$, which we introduced in Chapter 3, provides a way to characterize a method’s selectivity.

$$K_{A,I} = \frac{k_I}{k_A}$$  \hspace{1cm} (7.11)

Solving equation 7.11 for $k_I$, substituting into equation 7.10, and simplifying, gives

$$S_{\text{samp}} = k_A (C_A + K_{A,I} \times C_I)$$  \hspace{1cm} (7.12)

An interferent, therefore, does not pose a problem as long as the product of its concentration and its selectivity coefficient is significantly smaller than the analyte’s concentration.

$$K_{A,I} \times C_I << C_A$$

If we cannot ignore an interferent’s contribution to the signal, then we must begin our analysis by separating the analyte and the interferent.

7E General Theory of Separation Efficiency

The goal of an analytical separation is to remove either the analyte or the interferent from the sample’s matrix. To achieve this separation we must identify at least one significant difference between the analyte’s and the interferent’s chemical or physical properties. A significant difference in properties, however, is not sufficient to effect a separation if the conditions that favor the extraction of interferent from the sample also removes a small amount of analyte.
Two factors limit a separation’s efficiency: failing to recover all the analyte and failing to remove all the interferent. We define the analyte’s recovery, \( R_A \), as

\[
R_A = \frac{C_A}{(C_A)_o}
\]

where \( C_A \) is the concentration of analyte that remains after the separation, and \( (C_A)_o \) is the analyte’s initial concentration. A recovery of 1.00 means that no analyte is lost during the separation. The interferent’s recovery, \( R_I \), is defined in the same manner

\[
R_I = \frac{C_I}{(C_I)_o}
\]

where \( C_I \) is the concentration of interferent that remains after the separation, and \( (C_I)_o \) is the interferent’s initial concentration. We define the extent of the separation using a separation factor, \( S_{I,A} \),

\[
S_{I,A} = \frac{R_I}{R_A}
\]

In general, an \( S_{I,A} \) of approximately \( 10^{-7} \) is needed for the quantitative analysis of a trace analyte in the presence of a macro interferent, and \( 10^{-3} \) when the analyte and interferent are present in approximately equal amounts.

Example 7.10

An analytical method for determining Cu in an industrial plating bath gives poor results in the presence of Zn. To evaluate a method for separating the analyte from the interferent, samples with known concentrations of Cu or Zn were prepared and analyzed. When a sample of 128.6 ppm Cu was taken through the separation, the concentration of Cu that remained was 127.2 ppm. Taking a 134.9 ppm solution of Zn through the separation left behind a concentration of 4.3 ppm Zn. Calculate the recoveries for Cu and Zn, and the separation factor.

Solution

Using equation 7.13 and equation 7.14, the recoveries for the analyte and interferent are

\[
R_{Cu} = \frac{127.2 \text{ ppm}}{128.6 \text{ ppm}} = 0.9891 \text{ or } 98.91\%
\]

\[
R_{Zn} = \frac{4.3 \text{ ppm}}{134.9 \text{ ppm}} = 0.032 \text{ or } 3.2\%
\]

and the separation factor is

\[
S_{Zn,Cu} = \frac{R_{Zn}}{R_{Cu}} = \frac{0.032}{0.9891} = 0.032
\]

Recoveries and separation factors are useful tools for evaluating a separation’s potential effectiveness; they do not, however, give a direct indication of the error that results from failing to remove all the interferent or from failing to completely recover the analyte. The relative error due to the separation, $E$, is

$$E = \frac{S_{\text{amp}} - S_{\text{amp}}^*}{S_{\text{amp}}^*}$$

where $S_{\text{amp}}$ is the sample’s signal for an ideal separation in which we completely recover the analyte.

$$S_{\text{amp}}^* = k_A(C_A)_o$$

Substituting equation 7.12 and equation 7.17 into equation 7.16, and rearranging

$$E = \frac{k_A(C_A + K_{AI} \times C_I) - k_A(C_A)_o}{k_A(C_A)_o}$$

$$E = \frac{C_A + K_{AI} \times C_I - (C_A)_o}{(C_A)_o}$$

$$E = \frac{C_A}{(C_A)_o} - \frac{(C_A)_o}{(C_A)_o} + \frac{K_{AI} \times C_I}{(C_A)_o}$$

leaves us with

$$E = (R_A - 1) + \frac{K_{AI} \times C_I}{(C_A)_o}$$

A more useful equation is obtained by solving equation 7.14 for $C_I$ and substituting into equation 7.18.

$$E = (R_A - 1) + \frac{K_{AI} \times (C_I)_o}{(C_A)_o} \times R_I$$

The first term of equation 7.19 accounts for the analyte’s incomplete recovery and the second term accounts for a failure to remove all the interferent.

### Example 7.11

Following the separation outlined in Example 7.10, an analysis is carried out to determine the concentration of Cu in an industrial plating bath. Analysis of standard solutions that contain either Cu or Zn give the following linear calibrations.

$$S_{Cu} = 1250 \text{ ppm}^{-1} \times C_{Cu} \quad \text{and} \quad S_{Zn} = 2310 \text{ ppm}^{-1} \times C_{Zn}$$

(a) What is the relative error if we analyze a sample without removing the Zn? Assume the initial concentration ratio, Cu:Zn, is 7:1. (b) What is the relative error if we first complete the separation with the recoveries determined in Example 7.10? (c) What is the maximum acceptable recovery for Zn if the recovery for Cu is 1.00 and if the error due to the separation must be no greater than 0.10%?
**Solution**

(a) If we complete the analysis without separating Cu and Zn, then \( R_{Cu} \) and \( R_{Zn} \) are exactly 1 and equation 7.19 simplifies to

\[
E = \frac{K_{Cu,Zn} \times (C_{Zn})_o}{(C_{Cu})_o}
\]

Using equation 7.11, we find that the selectivity coefficient is

\[
K_{Cu,Zn} = \frac{k_{Zn}}{k_{Cu}} = \frac{2310 \text{ ppm}^{-1}}{1250 \text{ ppm}^{-1}} = 1.85
\]

Given the initial concentration ratio of 7:1 for Cu and Zn, the relative error without the separation is

\[
E = \frac{1.85 \times 1}{7} = 0.264 \text{ or } 26.4\%
\]

(b) To calculate the relative error we substitute the recoveries from Example 7.10 into equation 7.19, obtaining

\[
E = (0.9891 - 1) + \frac{1.85 \times 1}{7} \times 0.032 = -0.0109 + 0.085 = -0.0024
\]

or –0.24%. Note that the negative determinate error from failing to recover all the analyte is offset partially by the positive determinate error from failing to remove all the interferent.

(c) To determine the maximum recovery for Zn, we make appropriate substitutions into equation 7.19

\[
E = 0.0010 = (1 - 1) + \frac{1.85 \times 1}{7} \times R_{Zn}
\]

and solve for \( R_{Zn} \), obtaining a recovery of 0.0038, or 0.38%. Thus, we must remove at least

\[
100.00\% - 0.38\% = 99.62\%
\]

of the Zn to obtain an error of 0.10% when \( R_{Cu} \) is exactly 1.

---

**7F Classifying Separation Techniques**

We can separate an analyte and an interferent if there is a significant difference in at least one of their chemical or physical properties. Table 7.4 provides a partial list of separation techniques, organized by the chemical or physical property affecting the separation.

**7F.1 Separations Based on Size**

Size is the simplest physical property we can exploit in a separation. To accomplish the separation we use a porous medium through which only the analyte or the interferent can pass. Examples of size-based separations include filtration, dialysis, and size-exclusion.
In a **filtration** we separate a particulate interferent from soluble analytes using a filter with a pore size that will retain the interferent. The solution that passes through the filter is called the **filtrate**, and the material retained by the filter is the **retentate**. Gravity filtration and suction filtration using filter paper are techniques with which you should already be familiar. A membrane filter is the method of choice for particulates that are too small to be retained by filter paper. **Figure 7.13** provides information about three types of membrane filters.

**Dialysis** is another example of a separation technique in which size is used to separate the analyte and the interferent. A dialysis membrane usually is made using cellulose and fashioned into tubing, bags, or cassettes. **Figure 7.14** shows an example of a commercially available dialysis cassette. The sample is injected into the dialysis membrane, which is sealed tightly by a gasket, and the unit is placed in a container filled with a solution with a composition different from the sample. If there is a difference in a species’ concentration on the membrane’s two sides, the resulting concentration gradient provides a driving force for its diffusion across the membrane. While small species freely pass through the membrane, larger species are unable to pass. Dialysis frequently is used to purify proteins, hormones, and enzymes. During kidney dialysis, metabolic waste products, such as urea, uric acid, and creatinine, are removed from blood by passing it over a dialysis membrane.

**Size-exclusion chromatography** is a third example of a separation technique that uses size as a means to effect a separation. In this technique a column is packed with small, approximately 10-µm, porous polymer beads of cross-linked dextrin or polyacrylamide. The pore size of the particles is controlled by the degree of cross-linking, with more cross-linking producing smaller pore sizes. The sample is placed into a stream of solvent that is

<table>
<thead>
<tr>
<th>Basis of Separation</th>
<th>Separation Technique</th>
</tr>
</thead>
<tbody>
<tr>
<td>size</td>
<td>filtration</td>
</tr>
<tr>
<td></td>
<td>dialysis</td>
</tr>
<tr>
<td></td>
<td>size-exclusion chromatography</td>
</tr>
<tr>
<td>mass or density</td>
<td>centrifugation</td>
</tr>
<tr>
<td>complex formation</td>
<td>masking</td>
</tr>
<tr>
<td>change in physical state</td>
<td>distillation</td>
</tr>
<tr>
<td></td>
<td>sublimation</td>
</tr>
<tr>
<td></td>
<td>recrystallization</td>
</tr>
<tr>
<td>change in chemical state</td>
<td>precipitation</td>
</tr>
<tr>
<td></td>
<td>electrodeposition</td>
</tr>
<tr>
<td></td>
<td>volatilization</td>
</tr>
<tr>
<td>partitioning between phases</td>
<td>extraction</td>
</tr>
<tr>
<td></td>
<td>chromatography</td>
</tr>
</tbody>
</table>

**Table 7.4  Classification of Separation Techniques**

For applications of gravity filtration and suction filtration in gravimetric methods of analysis, see Chapter 8.
pumped through the column at a fixed flow rate. Those species too large
to enter the pores pass through the column at the same rate as the solvent.
Species that enter into the pores take longer to pass through the column,
with smaller species requiring more time to pass through the column. Size-
exclusion chromatography is widely used in the analysis of polymers, and
in biochemistry, where it is used for the separation of proteins.

7F.2 Separations Based on Mass or Density

If the analyte and the interferent have different masses or densities, then a
separation using CENTRIFUGATION may be possible. The sample is placed in
a centrifuge tube and spun at a high angular velocity, measured in revolu-
tions per minute (rpm). The sample’s constituents experience a centrifugal
force that pulls them toward the bottom of the centrifuge tube. Those spe-
cies that experience the greatest centrifugal force have the fastest sedimenta-
tion rate and are the first to reach the bottom of the centrifuge tube. If two
species have the same density, their separation is based on a difference in
mass, with the heavier species having the greater sedimentation rate. If the
species are of equal mass, then the species with the larger density has the
greatest sedimentation rate.
Centrifugation is an important separation technique in biochemistry. Table 7.5, for example, lists conditions for separating selected cellular components. We can separate lysosomes from other cellular components by several differential centrifugations, in which we divide the sample into a solid residue and a supernatant solution. After destroying the cells, the solution is centrifuged for 20 minutes at $15\,000 \times g$ (a centrifugal force that is 15,000 times the earth’s gravitational force), leaving a solid residue of cell membranes and mitochondria. The supernatant, which contains the lysosomes, is isolated by decanting it from the residue and then centrifuged for 30 minutes at $30\,000 \times g$, leaving a solid residue of lysosomes. Figure 7.15 shows a typical centrifuge capable of producing the centrifugal forces needed for biochemical separations.

An alternative approach to differential centrifugation is a density gradient centrifugation. To prepare a sucrose density gradient, for example, a solution with a smaller concentration of sucrose—and, thus, of lower density—is gently layered upon a solution with a higher concentration of sucrose. Repeating this process several times, fills the centrifuge tube with a multi-layer density gradient. The sample is placed on top of the density

**Table 7.5  Conditions for Separating Selected Cellular Components by Centrifugation**

<table>
<thead>
<tr>
<th>Components</th>
<th>Centrifugal Force ($\times g$)</th>
<th>Time (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>eukaryotic cells</td>
<td>1000</td>
<td>5</td>
</tr>
<tr>
<td>cell membranes, nuclei</td>
<td>4000</td>
<td>10</td>
</tr>
<tr>
<td>mitochondria, bacterial cells</td>
<td>15,000</td>
<td>20</td>
</tr>
<tr>
<td>lysosomes, bacterial membranes</td>
<td>30,000</td>
<td>30</td>
</tr>
<tr>
<td>ribosomes</td>
<td>100,000</td>
<td>180</td>
</tr>
</tbody>
</table>

Source: Adapted from Zubay, G. *Biochemistry*, 2nd ed. Macmillan: New York, 1988, p.120.

**Figure 7.14** Example of a dialysis cassette. The dialysis membrane in this unit has a molecular weight cut-off of 10,000 g/mol. Two sheets of the membrane are separated by a gasket and held in place by the plastic frame. Four ports, one of which is labeled, provide a means for injecting the sample between the dialysis membranes. The cassette is inverted and submerged in a beaker that contains the external solution. A foam buoy, used as a stand in the photo, serves as a float so that the unit remains suspended in the external solution. The external solution is stirred using a stir bar, and usually replaced several times during dialysis. When dialysis is complete, the solution in the cassette is removed through an injection port.

**Figure 7.15** Bench-top centrifuge capable of reaching speeds up to 14,000 rpm and centrifugal forces of $20,800 \times g$. This particular centrifuge is refrigerated, allowing samples to be cooled to temperatures as low as $-4 \, ^\circ C$. 
gradient and centrifuged using a force greater than $150,000 \times g$. During centrifugation, each of the sample’s components moves through the gradient until it reaches a position where its density matches the surrounding sucrose solution. Each component is isolated as a separate band positioned where its density is equal to that of the local density within the gradient. Figure 7.16 provides an example of a typical sucrose density centrifugation for separating plant thylakoid membranes.

**Figure 7.16** Example of a sucrose density gradient centrifugation of thylakoid membranes from wild type (WT) and *lut2* plants. The thylakoid membranes were extracted from the plant’s leaves and separated by centrifuging in a 0.1–1 M sucrose gradient for 22 h at 280,000 $\times g$ and at 4°C. Six bands and their chlorophyll contents are shown. Adapted from Dall’Osto, L.; Lico, C.; Alric, J.; Giuliano, G.; Havaux, M.; Bassi, R. *BMC Plant Biology* 2006, 6:32.

Technically, masking is not a separation technique because we do not physically separate the analyte and the interferent. We do, however, chemically isolate the interferent from the analyte, resulting in a pseudo-separation.

### 7F.3 Separations Based on Complexation Reactions (Masking)

One widely used technique for preventing an interference is to bind the interferent in a strong, soluble complex that prevents it from interfering in the analyte’s determination. This process is known as **masking**. As shown in Table 7.6, a wide variety of ions and molecules are useful masking agents, and, as a result, selectivity is usually not a problem.

**Example 7.12**

Using Table 7.6, suggest a masking agent for the analysis of aluminum in the presence of iron.

**Solution**

A suitable masking agent must form a complex with the interferent, but not with the analyte. Oxalate, for example, is not a suitable masking agent because it binds both Al and Fe. Thioglycolic acid, on the other hand, is a selective masking agent for Fe in the presence of Al. Other acceptable
masking agents are cyanide \((\text{CN}^-)\), thiocyanate \((\text{SCN}^-)\), and thiosulfate \((\text{S}_2\text{O}_3^{2-})\).

### Practice Exercise 7.6

Using Table 7.6, suggest a masking agent for the analysis of Fe in the presence of Al.

Click [here](#) to review your answer to this exercise.

As shown in Example 7.13, we can judge a masking agent’s effectiveness by considering the relevant equilibrium constants.

### Example 7.13

Show that \(\text{CN}^-\) is an appropriate masking agent for \(\text{Ni}^{2+}\) in a method where nickel’s complexation with EDTA is an interference.

**Solution**

The relevant reactions and formation constants are

\[
\text{Ni}^{2+} (aq) + Y^4- (aq) = \text{Ni}Y^2- (aq) \quad K_1 = 4.2 \times 10^{18} \\
\text{Ni}^{2+} (aq) + 4\text{CN}^- (aq) = \text{Ni} (\text{CN})_4^{2-} (aq) \quad \beta_4 = 1.7 \times 10^{10}
\]

where \(Y^4-\) is an abbreviation for EDTA. Cyanide is an appropriate masking agent because the formation constant for \(\text{Ni} (\text{CN})_4^{2-}\) is greater than that for the Ni–EDTA complex. In fact, the equilibrium constant for the reaction in which EDTA displaces the masking agent

\[
\text{Ni} (\text{CN})_4^{2-} (aq) + Y^4- (aq) = \text{Ni}Y^2- (aq) + 4\text{CN}^- (aq) \\
K = \frac{K_1}{\beta_4} = \frac{4.2 \times 10^{18}}{1.7 \times 10^{10}} = 2.5 \times 10^{-12}
\]

is sufficiently small that \(\text{Ni} (\text{CN})_4^{2-}\) is relatively inert in the presence of EDTA.

### Table 7.6  Selected Inorganic and Organic Masking Agents for Metal Ions

<table>
<thead>
<tr>
<th>Masking Agent</th>
<th>Elements Whose Ions Are Masked</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\text{CN}^-)</td>
<td>Ag, Au, Cd, Co, Cu, Fe, Hg, Mn, Ni, Pd, Pt, Zn</td>
</tr>
<tr>
<td>(\text{SCN}^-)</td>
<td>Ag, Cd, Co, Cu, Fe, Ni, Pd, Pt, Zn</td>
</tr>
<tr>
<td>(\text{NH}_3)</td>
<td>Ag, Co, Ni, Cu, Zn</td>
</tr>
<tr>
<td>(\text{F}^-)</td>
<td>Al, Co, Cr, Mg, Mn, Sn, Zn</td>
</tr>
<tr>
<td>(\text{S}_2\text{O}_3^{2-})</td>
<td>Au, Ce, Co, Cu, Fe, Hg, Mn, Pb, Pd, Pt, Sb, Sn, Zn</td>
</tr>
<tr>
<td>tartrate</td>
<td>Al, Ba, Bi, Ca, Ce, Co, Cr, Cu, Fe, Hg, Mn, Pb, Pd, Pt, Sb, Sn, Zn</td>
</tr>
<tr>
<td>oxalate</td>
<td>Al, Fe, Mg, Mn</td>
</tr>
<tr>
<td>thioglycolic acid</td>
<td>Cu, Fe, Sn</td>
</tr>
</tbody>
</table>

Practice Exercise 7.7

Use the formation constants in Appendix 12 to show that 1,10-phenanthroline is a suitable masking agent for Fe$^{2+}$ in the presence of Fe$^{3+}$. Use a ladder diagram to define any limitations on using 1,10-phenanthroline as a masking agent. See Chapter 6 for a review of ladder diagrams.

Click here to review your answer to this exercise.

7F.4 Separations Based on a Change of State

Because an analyte and its interferent are usually in the same phase, we can achieve a separation if one of them undergoes a change in its physical state or its chemical state.

Changes in Physical State

When the analyte and the interferent are miscible liquids, separation by distillation is possible if their boiling points are significantly different. Figure 7.17 shows the progress of a distillation as a plot of temperature versus the composition of mixture’s vapor-phase and liquid-phase. The initial liquid mixture (point A), contains more interferent than analyte. When this solution is brought to its boiling point, the vapor phase in equilibrium with the liquid phase is enriched in analyte (point B). The horizontal line that connects points A and B represents this vaporization equilibrium. Condensing the vapor phase at point B, by lowering the temperature, creates a new liquid phase with a composition identical to that in the vapor phase (point C). The vertical line that connects points B and C represents this condensation equilibrium. The liquid phase at point C has a lower boiling point than the original mixture, and is in equilibrium with the vapor phase at point D. This process of repeated vaporization and condensation gradually separates the analyte and the interferent.

Figure 7.17 Boiling point versus composition diagram for a near-ideal solution consisting of a low-boiling analyte and a high-boiling interferent. The horizontal lines represent vaporization equilibria and the vertical lines represent condensation equilibria. See the text for additional details.
Two experimental set-ups for distillations are shown in Figure 7.18. The simple distillation apparatus shown in Figure 7.18a is useful only for separating a volatile analyte (or interferent) from a non-volatile interferent (or analyte), or for separating an analyte and an interferent whose boiling points differ by more than 150 °C. A more efficient separation is achieved using the fractional distillation apparatus in Figure 7.18b. Packing the fractionating column with a high surface area material, such as a steel sponge or glass beads, provides more opportunity for the repeated process of vaporization and condensation necessary to effect a complete separation.

When the sample is a solid, sublimation may provide a useful separation of the analyte and the interferent. The sample is heated at a temperature and pressure below the analyte's triple point, allowing it to vaporize without passing through a liquid state. Condensing the vapor recovers the purified analyte (Figure 7.19). A useful analytical example of sublimation is the isolation of amino acids from fossil mollusk shells and deep-sea sediments.\(^\text{13}\)

Recrystallization is another method for purifying a solid. A solvent is chosen in which the analyte’s solubility is significant when the solvent is hot and minimal when the solvent is cold. The interferents must be less soluble in the hot solvent than the analyte or present in much smaller amounts. After heating a portion of the solvent in an Erlenmeyer flask, small amounts of sample are added until undissolved sample is visible. Additional hot [13] Glavin, D. P.; Bada, J. L. *Anal. Chem.* 1998, 70, 3119–3122.
solvent is added until the sample redissolves, or until only insoluble impurities remain. This process of adding sample and solvent is repeated until the entire sample is added to the Erlenmeyer flask. Any insoluble impurities are removed by filtering the hot solution. The solution is allowed to cool slowly, which promotes the growth of large, pure crystals, and then cooled in an ice bath to minimize solubility losses. The purified sample is isolated by filtration and rinsed to remove any soluble impurities. Finally, the sample is dried to remove any remaining traces of the solvent. Further purification, if necessary, is accomplished by additional recrystallizations.

**Changes in Chemical State**

Distillation, sublimation, and recrystallization use a change in physical state to effect a separation. Chemical reactivity also is a useful tool for separating analytes and interferents. For example, we can separate SiO$_2$ from a sample by reacting it with HF to form SiF$_4$. Because SiF$_4$ is volatile, it is easy to remove by evaporation. If we wish to collect the reaction’s volatile product, then a distillation is possible. For example, we can isolate the NH$_4^+$ in a sample by making the solution basic and converting it to NH$_3$. The ammonia is then removed by distillation. Table 7.7 provides additional examples of this approach for isolating inorganic ions.

Another reaction for separating analytes and interferents is precipitation. Two important examples of using a precipitation reaction in a separation are the pH-dependent solubility of metal oxides and hydroxides, and the pH-dependent solubility of metal sulfides.

Separations based on the pH-dependent solubility of oxides and hydroxides usually use a strong acid, a strong base, or an NH$_3$/NH$_4$Cl buffer to adjust the pH. Most metal oxides and hydroxides are soluble in hot concentrated HNO$_3$, although a few oxides, such as WO$_3$, SiO$_2$, and SnO$_2$ remain insoluble even under these harsh conditions. To determine the amount of Cu in brass, for example, we can avoid an interference from Sn by dissolving the sample with a strong acid and filtering to remove the solid residue of SnO$_2$.

Most metals form a hydroxide precipitate in the presence of concentrated NaOH. Those metals that form amphoteric hydroxides, however, do not precipitate because they react to form higher-order hydroxo-complexes.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Treatment</th>
<th>Isolated Species</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO$_3^{2-}$</td>
<td>CO$_3^{2-}(aq) + 2H_3O^+(aq) \rightarrow CO_2(g) + 3H_2O(l)$</td>
<td>CO$_2$</td>
</tr>
<tr>
<td>NH$_4^+$</td>
<td>NH$_4^+(aq) + OH^- (aq) \rightarrow NH_3(aq) + H_2O(l)$</td>
<td>NH$_3$</td>
</tr>
<tr>
<td>SO$_3^{2-}$</td>
<td>SO$_3^{2-}(aq) + 2H_3O^+(aq) \rightarrow SO_2(g) + 3H_2O(l)$</td>
<td>SO$_2$</td>
</tr>
<tr>
<td>S$^{2-}$</td>
<td>S$^{2-}(aq) + 2H_3O^+(aq) \rightarrow H_2S(g) + 2H_2O(l)$</td>
<td>H$_2$S</td>
</tr>
</tbody>
</table>
For example, Zn\(^{2+}\) and Al\(^{3+}\) do not precipitate in concentrated NaOH because they form the soluble complexes Zn(OH)\(^3\) and Al(OH)\(^4\). The solubility of Al\(^{3+}\) in concentrated NaOH allows us to isolate aluminum from impure samples of bauxite, an ore of Al\(_2\)O\(_3\). After crushing the ore, we place it in a solution of concentrated NaOH, dissolving the Al\(_2\)O\(_3\) and forming Al(OH)\(^4\). Other oxides in the ore, such as Fe\(_2\)O\(_3\) and SiO\(_2\), remain insoluble. After filtering, we recover the aluminum as a precipitate of Al(OH)\(_3\) by neutralizing some of the OH\(^-\) with acid.

The pH of an NH\(_3\)/NH\(_4\)Cl buffer (pK\(_a\) = 9.26) is sufficient to precipitate most metals as the hydroxide. The alkaline earths and alkaline metals, however, do not precipitate at this pH. In addition, metal ions that form soluble complexes with NH\(_3\), such as Cu\(^{2+}\), Zn\(^{2+}\), Ni\(^{2+}\), and Co\(^{2+}\) also do not precipitate under these conditions.

The use of S\(^2-\) as a precipitating reagent is one of the earliest examples of a separation technique. In Fresenius’s 1881 text *A System of Instruction in Quantitative Chemical Analysis*, sulfide frequently is used to separate metal ions from the remainder of the sample’s matrix.\(^{14}\) Sulfide is a useful reagent for separating metal ions for two reasons: (1) most metal ions, except for the alkaline earths and alkaline metals, form insoluble sulfides; and (2) these metal sulfides show a substantial variation in solubility. Because the concentration of S\(^2-\) is pH-dependent, we can control which metal ions precipitate by adjusting the pH. For example, in Fresenius’s gravimetric procedure for the determination of Ni in ore samples (see Figure 1.1 in Chapter 1 for a schematic diagram of this procedure), sulfide is used three times to separate Co\(^{2+}\) and Ni\(^{2+}\) from Cu\(^{2+}\) and, to a lesser extent, from Pb\(^{2+}\).

### 7F.5 Separations Based on a Partitioning Between Phases

The most important group of separation techniques uses a selective partitioning of the analyte or interferent between two immiscible phases. If we bring a phase that contains the solute, S, into contact with a second phase, the solute will partition itself between the two phases, as shown by the following equilibrium reaction.

\[
S_{\text{phase 1}} = S_{\text{phase 2}}
\]  

7.20

The equilibrium constant for reaction 7.20

\[
K_D = \frac{[S_{\text{phase 2}}]}{[S_{\text{phase 1}}]}
\]

is called the distribution constant or the **partition coefficient**. If \(K_D\) is sufficiently large, then the solute moves from phase 1 to phase 2. The solute will remain in phase 1 if the partition coefficient is sufficiently small. When we bring a phase that contains two solutes into contact with a second phase, a separation of the solutes is possible if \(K_D\) is favorable for only one of the solutes. The physical states of the phases are identified when we describe the

separation process, with the phase that contains the sample listed first. For example, if the sample is in a liquid phase and the second phase is a solid, then the separation involves liquid–solid partitioning.

**Extraction Between Two Phases**

We call the process of moving a species from one phase to another phase an [*extraction*](#). Simple extractions are particularly useful for separations where only one component has a favorable partition coefficient. Several important separation techniques are based on a simple extraction, including liquid–liquid, liquid–solid, solid–liquid, and gas–solid extractions.

**Liquid–Liquid Extractions**

A liquid–liquid extraction usually is accomplished using a separatory funnel (Figure 7.20). After placing the two liquids in the separatory funnel, we shake the funnel to increase the surface area between the phases. When the extraction is complete, we allow the liquids to separate. The stopcock at the bottom of the separatory funnel allows us to remove the two phases. We also can carry out a liquid–liquid extraction without a separatory funnel by adding the extracting solvent to the sample’s container. Pesticides in water, for example, are preserved in the field by extracting them into a small volume of hexane. A liquid–liquid microextraction, in which the extracting phase is a 1-µL drop suspended from a microsyringe (Figure 7.21), also has been described. Because of its importance, a more thorough discussion of liquid–liquid extractions is in Section 7G.

**Solid Phase Extractions**

In a solid phase extraction of a liquid sample, we pass the sample through a cartridge that contains a solid adsorbent, several examples of which are shown in Figure 7.22. The choice of adsorbent is determined by the species we wish to separate. Table 7.8 provides several representative examples of solid adsorbents and their applications.

As an example, let’s examine a procedure for isolating the sedatives secobarbital and phenobarbital from serum samples using a C-18 solid adsorbent. Before adding the sample, the solid phase cartridge is rinsed with 6 mL each of methanol and water. Next, a 500-µL sample of serum is pulled through the cartridge, with the sedatives and matrix interferents retained following a liquid–solid extraction (Figure 7.23a). Washing the cartridge with distilled water removes any interferents (Figure 7.23b). Finally, we elute the sedatives using 500 µL of acetone (Figure 7.23c). In comparison to a liquid–liquid extraction, a solid phase extraction has the advantage of being easier, faster, and requires less solvent.

---

**Figure 7.20** Example of a liquid–liquid extraction using a separatory funnel. (a) Before the extraction, 100% of the analyte is in phase 1. (b) After the extraction, most of the analyte is in phase 2, although some analyte remains in phase 1.

Although one liquid–liquid extraction can result in the complete transfer of analyte, a single extraction usually is not sufficient. See Section 7G for a discussion of extraction efficiency and multiple extractions.

**Figure 7.21** Schematic of a liquid–liquid microextraction showing a syringe needle with a µL drop of the extracting solvent.

---

Continuous Extractions

An extraction is possible even if the analyte has an unfavorable partition coefficient, provided that the sample’s other components have significantly smaller partition coefficients. Because the analyte’s partition coefficient is unfavorable, a single extraction will not recover all the analyte. Instead we continuously pass the extracting phase through the sample until we achieve a quantitative extraction.

A continuous extraction of a solid sample is carried out using a Soxhlet extractor (Figure 7.24). The extracting solvent is placed in the lower reservoir and heated to its boiling point. Solvent in the vapor phase moves upward through the tube on the far right side of the apparatus, reaching the condenser where it condenses back to the liquid state. The solvent then

![Solid phase extraction cartridges](image)

**Figure 7.22** Selection of solid phase extraction cartridges for liquid samples. The solid adsorbent is the white or black material in each cartridge. From left-to-right, the absorbent materials are octadecylsilane, carbon, octadecylsilane, polyamide resin, and diol; see Table 7.8 for additional details. The size of the cartridges dictates the volume of sample used; from left-to-right, these cartridges use samples of 1 mL, 3 mL, 6 mL, 3 mL, and 1 mL.

<table>
<thead>
<tr>
<th>Adsorbent</th>
<th>Structure</th>
<th>Properties and Uses</th>
</tr>
</thead>
</table>
| silica          | ![silica](structure) | • retains low to moderate polarity species from organic matrices  
                    • fat soluble vitamins, steroids                                           |
| aminopropyl     | ![aminopropyl](structure) | • retains polar compounds  
                    • carbohydrates, organic acids                                              |
| cyanopropyl     | ![cyanopropyl](structure) | • retains wide variety of species from aqueous and organic matrices  
                    • pesticides, hydrophobic peptides                                           |
| diol            | ![diol](structure) | • retains wide variety of species from aqueous and organic matrices  
                    • proteins, peptides, fungicides                                              |
| octadecyl (C-18)| ![octadecyl](structure) | • retains hydrophobic species from aqueous matrices  
                    • caffeine, sedatives, polyaromatic hydrocarbons, carbohydrates, pesticides |
| octyl (C-8)     | ![octyl](structure) | • similar to C-18                                                                  |
passes through the sample, which is held in a porous cellulose filter thimble, collecting in the upper reservoir. When the solvent in the upper reservoir reaches the return tube’s upper bend, the solvent and extracted analyte are siphoned back to the lower reservoir. Over time the analyte’s concentration in the lower reservoir increases.

Microwave-assisted extractions have replaced Soxhlet extractions in some applications. The process is the same as that described earlier for a microwave digestion. After placing the sample and the solvent in a sealed digestion vessel, a microwave oven is used to heat the mixture. Using a sealed digestion vessel allows the extraction to take place at a higher temperature and pressure, reducing the amount of time needed for a quantitative extraction. In a Soxhlet extraction the temperature is limited by the solvent’s boiling point at atmospheric pressure. When acetone is the solvent, for example, a Soxhlet extraction is limited to 56°C, but a microwave extraction can reach 150°C.

Two other continuous extractions deserve mention. Volatile organic compounds (VOCs) can be quantitatively removed from a liquid sample by a liquid–gas extraction. As shown in Figure 7.25, an inert purging gas, such as He, is passed through the sample. The purge gas removes the VOCs, which are swept to a primary trap where they collect on a solid absorbent. When the extraction is complete, the VOCs are removed from the primary trap by rapidly heating the tube while flushing with He. This technique is known as a PURGE-AND-TRAP. Because the analyte’s recovery may not be reproducible, an internal standard is required for quantitative work.

---

Figure 7.23 Steps in a typical solid phase extraction. After preconditioning the solid phase cartridge with solvent, (a) the sample is added to the cartridge, (b) the sample is washed to remove interferents, and (c) the analytes are eluted.

Figure 7.24 Soxhlet extractor. See text for details.

Continuous extractions also can be accomplished using supercritical fluids. If we heat a substance above its critical temperature and pressure it forms a supercritical fluid whose properties are between those of a gas and a liquid. A supercritical fluid is a better solvent than a gas, which makes it a better reagent for extractions. In addition, a supercritical fluid’s viscosity is significantly less than that of a liquid, which makes it easier to push it through a particulate sample. One example of a supercritical fluid extraction is the determination of total petroleum hydrocarbons (TPHs) in soils, sediments, and sludges using supercritical CO$_2$. An approximately 3-g sample is placed in a 10-mL stainless steel cartridge and supercritical CO$_2$ at a pressure of 340 atm and a temperature of 80°C is passed through the cartridge for 30 minutes at flow rate of 1–2 mL/min. To collect the TPHs, the effluent from the cartridge is passed through 3 mL of tetrachloroethylene at room temperature. At this temperature the CO$_2$ reverts to the gas phase and is released to the atmosphere.

**CHROMATOGRAPHIC SEPARATIONS**

In an extraction, the sample originally is in one phase and we extract the analyte or the interferent into a second phase. We also can separate the analyte and interferents by continuously passing one sample-free phase, called the mobile phase, over a second sample-free phase that remains fixed or stationary. The sample is injected into the mobile phase and the sample’s components partition themselves between the mobile phase and the stationary phase. Those components with larger partition coefficients are more likely to move into the stationary phase and take longer time to pass through the system. This is the basis of all chromatographic separations.

---

Chromatography provides both a separation of analytes and interferents, and a means for performing a qualitative or quantitative analysis for the analyte. For this reason a more thorough treatment of chromatography is found in Chapter 12.

7G Liquid–Liquid Extractions

A liquid–liquid extraction is an important separation technique for environmental, clinical, and industrial laboratories. A standard environmental analytical method illustrates the importance of liquid–liquid extractions. Municipal water departments routinely monitor public water supplies for trihalomethanes (CHCl$_3$, CHBrCl$_2$, CHBr$_2$Cl, and CHBr$_3$) because they are known or suspected carcinogens. Before their analysis by gas chromatography, trihalomethanes are separated from their aqueous matrix using a liquid–liquid extraction with pentane.$^{20}$

In a simple liquid–liquid extraction the solute partitions itself between two immiscible phases. One phase usually is an aqueous solvent and the other phase is an organic solvent, such as the pentane used to extract trihalomethanes from water. Because the phases are immiscible they form two layers, with the denser phase on the bottom. The solute initially is present in one of the two phases; after the extraction it is present in both phases. **Extraction efficiency**—that is, the percentage of solute that moves from one phase to the other—is determined by the equilibrium constant for the solute’s partitioning between the phases and any other side reactions that involve the solute. Examples of other reactions that affect extraction efficiency include acid–base reactions and complexation reactions.

7G.1 Partition Coefficients and Distribution Ratios

As we learned earlier in this chapter, a solute’s partitioning between two phases is described by a partition coefficient, $K_D$. If we extract a solute from an aqueous phase into an organic phase

\[
S_{\text{org}} = S_{\text{aq}}
\]

then the partition coefficient is

\[
K_D = \frac{[S_{\text{org}}]}{[S_{\text{aq}}]}
\]

A large value for $K_D$ indicates that extraction of solute into the organic phase is favorable.

To evaluate an extraction’s efficiency we must consider the solute’s total concentration in each phase, which we define as a **distribution ratio**, $D$.

\[
D = \frac{[S_{\text{org}}]_{\text{total}}}{[S_{\text{aq}}]_{\text{total}}}
\]

---

The partition coefficient and the distribution ratio are identical if the solute has only one chemical form in each phase; however, if the solute exists in more than one chemical form in either phase, then $K_D$ and $D$ usually have different values. For example, if the solute exists in two forms in the aqueous phase, $A$ and $B$, only one of which, $A$, partitions between the two phases, then

$$D = \frac{[S_{\text{org}}]_A}{[S_{\text{org}}]_A + [S_{\text{aq}}]_B} \leq K_D = \frac{[S_{\text{org}}]_A}{[S_{\text{aq}}]_A}$$

This distinction between $K_D$ and $D$ is important. The partition coefficient is a thermodynamic equilibrium constant and has a fixed value for the solute’s partitioning between the two phases. The distribution ratio’s value, however, changes with solution conditions if the relative amounts of $A$ and $B$ change. If we know the solute’s equilibrium reactions within each phase and between the two phases, we can derive an algebraic relationship between $K_D$ and $D$.

### 7G.2 Liquid–Liquid Extraction With No Secondary Reactions

In a simple liquid–liquid extraction, the only reaction that affects the extraction efficiency is the solute’s partitioning between the two phases (Figure 7.26). In this case the distribution ratio and the partition coefficient are equal.

$$D = \frac{[S_{\text{org}}]_{\text{total}}}{[S_{\text{aq}}]_{\text{total}}} = K_D = \frac{[S_{\text{org}}]}{[S_{\text{aq}}]} \quad 7.21$$

Let’s assume the solute initially is present in the aqueous phase and that we wish to extract it into the organic phase. A conservation of mass requires that the moles of solute initially present in the aqueous phase equal the combined moles of solute in the aqueous phase and the organic phase after the extraction.

$$(\text{mol} \ S_{\text{aq}})_0 = (\text{mol} \ S_{\text{aq}})_1 + (\text{mol} \ S_{\text{org}})_1 \quad 7.22$$

where the subscripts indicate the extraction number. After the extraction, the solute’s concentration in the aqueous phase is

$$[S_{\text{aq}}]_1 = \frac{(\text{mol} \ S_{\text{aq}})_1}{V_{\text{aq}}} \quad 7.23$$

and its concentration in the organic phase is

$$[S_{\text{org}}]_1 = \frac{(\text{mol} \ S_{\text{org}})_1}{V_{\text{org}}} \quad 7.24$$

where $V_{\text{aq}}$ and $V_{\text{org}}$ are the volumes of the aqueous phase and the organic phase. Solving equation 7.22 for $(\text{mol} \ S_{\text{org}})_1$ and substituting into equation 7.24 leave us with

$$[S_{\text{org}}]_1 = \frac{(\text{mol} \ S_{\text{aq}})_0 - (\text{mol} \ S_{\text{aq}})_1}{V_{\text{org}}} \quad 7.25$$

Substituting equation 7.23 and equation 7.25 into equation 7.21 gives

![Figure 7.26 Scheme for a simple liquid–liquid extraction in which the solute’s partitioning depends only on the $K_D$ equilibrium.](image-url)
\[
D = \frac{(\text{mol } S_{aq})_0 - (\text{mol } S_{aq})_1}{V_{aoq}} = \frac{(\text{mol } S_{aq})_0 \times V_{aq} - (\text{mol } S_{aq})_1 \times V_{aq}}{(\text{mol } S_{aq})_1 \times V_{aq}}
\]

Rearranging and solving for the fraction of solute that remains in the aqueous phase after one extraction, \((q_{aq})_1\), gives

\[
(q_{aq})_1 = \frac{(\text{mol } S_{aq})_1}{(\text{mol } S_{aq})_0} = \frac{V_{aq}}{DV_{org} + V_{aq}}
\]

The fraction present in the organic phase after one extraction, \((q_{org})_1\), is

\[
(q_{org})_1 = \frac{(\text{mol } S_{org})_1}{(\text{mol } S_{org})_0} = 1 - (q_{aq})_1 = \frac{DV_{org}}{DV_{org} + V_{aq}}
\]

Example 7.14 shows how we can use equation 7.26 to calculate the efficiency of a simple liquid-liquid extraction.

**Example 7.14**

A solute has a \(K_D\) between water and chloroform of 5.00. Suppose we extract a 50.00-mL sample of a 0.050 M aqueous solution of the solute using 15.00 mL of chloroform. (a) What is the separation’s extraction efficiency? (b) What volume of chloroform do we need if we wish to extract 99.9% of the solute?

**Solution**

For a simple liquid–liquid extraction the distribution ratio, \(D\), and the partition coefficient, \(K_D\), are identical.

(a) The fraction of solute that remains in the aqueous phase after the extraction is given by equation 7.26.

\[
(q_{aq})_1 = \frac{V_{aq}}{DV_{org} + V_{aq}} = \frac{50.00 \text{ mL}}{(5.00)(15.00 \text{ mL}) + 50.00 \text{ mL}} = 0.400
\]

The fraction of solute in the organic phase is \(1 - 0.400\), or 0.600. Extraction efficiency is the percentage of solute that moves into the extracting phase; thus, the extraction efficiency is 60.0%.

(b) To extract 99.9% of the solute \((q_{aq})_1\) must be 0.001. Solving equation 7.26 for \(V_{org}\), and making appropriate substitutions for \((q_{aq})_1\) and \(V_{aq}\) gives

\[
V_{org} = \frac{V_{aq} - (q_{aq})_1 V_{aq}}{(q_{aq})_1 D} = \frac{50.00 \text{ mL} - (0.001)(50.00 \text{ mL})}{(0.001)(5.00 \text{ mL})} = 9990 \text{ mL}
\]

In Example 7.14, a single extraction provides an extraction efficiency of only 60%. If we carry out a second extraction, the fraction of solute remaining in the aqueous phase, \((q_{aq})_2\), is
\[
(q_{aq})_2 = \frac{(mol \ S_{aq})_2}{(mol \ S_{aq})_1} = \frac{V_{aq}}{DV_{org} + V_{aq}}
\]

If \(V_{aq}\) and \(V_{org}\) are the same for both extractions, then the cumulative fraction of solute that remains in the aqueous layer after two extractions, \((Q_{aq})_2\), is the product of \((q_{aq})_1\) and \((q_{aq})_2\), or
\[
(Q_{aq})_2 = \left(\frac{mol \ S_{aq}}{mol \ S_{aq}}\right)_0 = (q_{aq})_1 \times (q_{aq})_2 = \left(\frac{V_{aq}}{DV_{org} + V_{aq}}\right)^2
\]

In general, for a series of \(n\) identical extractions, the fraction of analyte that remains in the aqueous phase after the last extraction is
\[
(Q_{aq})_n = \left(\frac{V_{aq}}{DV_{org} + V_{aq}}\right)^n
\]

### Example 7.15

For the extraction described in Example 7.14, determine (a) the extraction efficiency for two identical extractions and for three identical extractions; and (b) the number of extractions required to ensure that we extract 99.9% of the solute.

**Solution**

(a) The fraction of solute remaining in the aqueous phase after two extractions and three extractions is
\[
(Q_{aq})_2 = \left(\frac{50.00 \ mL}{(5.00)(15.00 \ mL) + 50.00 \ mL}\right)^2 = 0.160
\]
\[
(Q_{aq})_3 = \left(\frac{50.00 \ mL}{(5.00)(15.00 \ mL) + 50.00 \ mL}\right)^3 = 0.0640
\]

The extraction efficiencies are 84.0% for two extractions and 93.6% for three extractions.

(b) To determine the minimum number of extractions for an efficiency of 99.9%, we set \((Q_{aq})_n\) to 0.001 and solve for \(n\) using equation 7.27.
\[
0.001 = \left(\frac{50.00 \ mL}{(5.00)(15.00 \ mL) + 50.00 \ mL}\right)^n = (0.400)^n
\]

Taking the log of both sides and solving for \(n\)
\[
\log(0.001) = n \log(0.400) \quad \quad n = 7.54
\]

we find that a minimum of eight extractions is necessary.

The last two examples provide us with an important observation—for any extraction efficiency, we need less solvent if we complete several extractions using smaller portions of solvent instead of one extraction using a larger volume of solvent. For the conditions in Example 7.14 and Example 7.15, an extraction efficiency of 99.9% requires one extraction with 9990 mL of chloroform, or 120 mL when using eight 15-mL portions of chloroform. Although extraction efficiency increases dramatically with the
first few multiple, the effect diminishes quickly as we increase the number of extractions (Figure 7.27). In most cases there is little improvement in extraction efficiency after five or six extractions. For the conditions in Example 7.15, we reach an extraction efficiency of 99% after five extractions and need three additional extractions to obtain the extra 0.9% increase in extraction efficiency.

**Practice Exercise 7.8**

To plan a liquid–liquid extraction we need to know the solute’s distribution ratio between the two phases. One approach is to carry out the extraction on a solution that contains a known amount of solute. After the extraction, we isolate the organic phase and allow it to evaporate, leaving behind the solute. In one such experiment, 1.235 g of a solute with a molar mass of 117.3 g/mol is dissolved in 10.00 mL of water. After extracting with 5.00 mL of toluene, 0.889 g of the solute is recovered in the organic phase. (a) What is the solute’s distribution ratio between water and toluene? (b) If we extract 20.00 mL of an aqueous solution that contains the solute using 10.00 mL of toluene, what is the extraction efficiency? (c) How many extractions will we need to recover 99.9% of the solute?

Click [here](#) to review your answer to this exercise.

### 7G.3 Liquid–Liquid Extractions Involving Acid–Base Equilibria

As we see in equation 7.21, in a simple liquid–liquid extraction the distribution ratio and the partition coefficient are identical. As a result, the distribution ratio does not depend on the composition of the aqueous phase or the organic phase. A change in the pH of the aqueous phase, for example, will not affect the solute’s extraction efficiency when \( K_D \) and \( D \) have the same value.

If the solute participates in one or more additional equilibrium reactions within a phase, then the distribution ratio and the partition coefficient may not be the same. For example, Figure 7.28 shows the equilibrium reactions that affect the extraction of the weak acid, HA, by an organic phase in which ionic species are not soluble. In this case the partition coefficient and the distribution ratio are

\[
K_D = \frac{[HA_{org}]}{[HA_{aq}]} \tag{7.28}
\]

\[
D = \frac{[HA_{org}]_{total}}{[HA_{aq}]_{total}} = \frac{[HA_{org}]}{[HA_{aq}] + [A_{aq}]} \tag{7.29}
\]

Because the position of an acid–base equilibrium depends on pH, the distribution ratio, \( D \), is pH-dependent. To derive an equation for \( D \) that shows this dependence, we begin with the acid dissociation constant for HA.
K_a H_2O \rightarrow H_3O^+ + A^-

**Figure 7.28** Scheme for the liquid–liquid extraction of a weak acid, HA. Although the weak acid is soluble in both phases, its conjugate weak base, $A^-$, is soluble in the aqueous phase only. The $K_a$ reaction for HA, which is called a **SECONDARY EQUILIBRIUM REACTION**, affects weak acid’s extraction efficiency because it determines the relative abundance of HA in solution.

$$K_i = \frac{[H_3O^+]_a [A^-]_a}{[HA]_a} \tag{7.30}$$

Solving equation 7.30 for the concentration of $A^-$ in the aqueous phase

$$[A^-]_a = \frac{K_i [HA]_a}{[H_3O^+]_a}$$

and substituting into equation 7.29 gives

$$D = \frac{[HA]_a}{[HA]_a + \frac{K_i [HA]_a}{[H_3O^+]_a}}$$

Factoring $[HA]_a$ from the denominator, replacing $[HA]_o/[HA]_a$ with $K_D$ (equation 7.28), and simplifying leaves us with the following relationship between the distribution ratio, $D$, and the pH of the aqueous solution.

$$D = \frac{K_D [H_3O^+]_a}{[H_3O^+]_a + K_i} \tag{7.31}$$

**Example 7.16**

An acidic solute, HA, has a $K_a$ of $1.00 \times 10^{-5}$ and a $K_D$ between water and hexane of 3.00. Calculate the extraction efficiency if we extract a 50.00 mL sample of a 0.025 M aqueous solution of HA, buffered to a pH of 3.00, with 50.00 mL of hexane. Repeat for pH levels of 5.00 and 7.00.

**Solution**

When the pH is 3.00, $[H_3O^+]_a$ is $1.0 \times 10^{-3}$ and the distribution ratio is

$$D = \frac{(3.00) (1.0 \times 10^{-3})}{1.0 \times 10^{-3} + 1.00 \times 10^{-5}} = 2.97$$

The fraction of solute that remains in the aqueous phase is

$$(Q_{aq})_1 = \frac{50.00 \text{ mL}}{(2.97)(50.00 \text{ mL}) + 50.00 \text{ mL}} = 0.252$$
The extraction efficiency, therefore, is almost 75%. The same calculation at a pH of 5.00 gives the extraction efficiency as 60%. At a pH of 7.00 the extraction efficiency is just 3%.

The extraction efficiency in Example 7.16 is greater at more acidic pH levels because HA is the solute’s predominate form in the aqueous phase. At a more basic pH, where A\(^-\) is the solute’s predominate form, the extraction efficiency is smaller. A graph of extraction efficiency versus pH is shown in Figure 7.29. Note that extraction efficiency essentially is independent of pH for pH levels more acidic than the HA’s pK\(_a\), and that it is essentially zero for pH levels more basic than HA’s pK\(_a\). The greatest change in extraction efficiency occurs at pH levels where both HA and A\(^-\) are predominate species. The ladder diagram for HA along the graph’s x-axis helps illustrate this effect.

**Practice Exercise 7.9**

The liquid–liquid extraction of the weak base B is governed by the following equilibrium reactions:

\[
B(aq) = B(org) \quad K_D = 5.00
\]

\[
B(aq) + H_2O(l) = OH^-(aq) + HB^+(aq) \quad K_a = 1.0 \times 10^{-4}
\]

Derive an equation for the distribution ratio, \(D\), and calculate the extraction efficiency if 25.0 mL of a 0.025 M solution of B, buffered to a pH of 9.00, is extracted with 50.0 mL of the organic solvent.

Click here to review your answer to this exercise.

7G.4 Liquid–Liquid Extraction of a Metal–Ligand Complex

One important application of a liquid–liquid extraction is the selective extraction of metal ions using an organic ligand. Unfortunately, many organic ligands are not very soluble in water or undergo hydrolysis or oxidation reactions in aqueous solutions. For these reasons the ligand is added to the organic solvent instead of the aqueous phase. Figure 7.30 shows the relevant equilibrium reactions (and equilibrium constants) for the extraction of M\(^{n+}\) by the ligand HL, including the ligand’s extraction into the aqueous phase (\(K_{D,HL}\)), the ligand’s acid dissociation reaction (\(K_a\)), the formation of the metal–ligand complex (\(\beta_n\)), and the complex’s extraction into the organic phase (\(K_{D,c}\)).

If the ligand’s concentration is much greater than the metal ion’s concentration, then the distribution ratio is

\[
D = \frac{\beta_n K_{D,c} (C_{HL})^n}{(K_{D,HL})^n [H_3O^+]^n + \beta_n (K_a)^n (C_{HL})^n}
\]

where \(C_{HL}\) is the ligand’s initial concentration in the organic phase. As shown in Example 7.17, the extraction efficiency for metal ions shows a marked pH dependency.

Problem 31 in the end-of-chapter problems asks you to derive equation 7.32.
Example 7.17

A liquid–liquid extraction of the divalent metal ion, M$^{2+}$, uses the scheme outlined in Figure 7.30. The partition coefficients for the ligand, $K_{D,HL}$, and for the metal–ligand complex, $K_{D,c}$, are $1.0 \times 10^4$ and $7.0 \times 10^4$, respectively. The ligand’s acid dissociation constant, $K_a$, is $5.0 \times 10^{-5}$, and the formation constant for the metal–ligand complex, $\beta_2$, is $2.5 \times 10^{16}$. What is the extraction efficiency if we extract 100.0 mL of a $1.0 \times 10^{-6}$ M aqueous solution of M$^{2+}$, buffered to a pH of 1.00, with 10.00 mL of an organic solvent that is 0.1 mM in the chelating agent? Repeat the calculation at a pH of 3.00.

SOLUTION

When the pH is 1.00 the distribution ratio is

$$D = \frac{(2.5 \times 10^6)(7.0 \times 10^3)(5.0 \times 10^{-5})^2(1.0 \times 10^{-4})^2}{(1.0 \times 10^{-4})^2(0.10)^2 + (2.5 \times 10^6)(5.0 \times 10^{-5})^2(1.0 \times 10^{-4})^2}$$

or a $D$ of 0.0438. The fraction of metal ion that remains in the aqueous phase is

$$(Q_{aq}) = \frac{100.0 \text{ mL}}{0.0438(10.00 \text{ mL}) + 100.0 \text{ mL}} = 0.996$$

At a pH of 1.00, we extract only 0.40% of the metal into the organic phase. Changing the pH to 3.00, however, increases the extraction efficiency to 97.8%. Figure 7.31 shows how the pH of the aqueous phase affects the extraction efficiency for M$^{2+}$.

**Figure 7.30** Scheme for the liquid–liquid extraction of a metal ion, M$^{n+}$, by the ligand L$^-$. The ligand initially is present in the organic phase as HL. Four equilibrium reactions are needed to explain the extraction efficiency.

**Figure 7.31** Plot of extraction efficiency versus pH for the extraction of the metal ion, M$^{2+}$, in Example 7.17.
One advantage of using a ligand to extract a metal ion is the high degree of selectivity that it brings to a liquid–liquid extraction. As seen in Figure 7.31, a divalent metal ion’s extraction efficiency increases from approximately 0% to 100% over a range of 2 pH units. Because a ligand’s ability to form a metal–ligand complex varies substantially from metal ion to metal ion, significant selectivity is possible if we carefully control the pH. Table 7.9 shows the minimum pH for extracting 99% of a metal ion from an aqueous solution using an equal volume of 4 mM dithizone in CCl$_4$.

**Example 7.18**

Using Table 7.9, explain how we can separate the metal ions in an aqueous mixture of Cu$^{2+}$, Cd$^{2+}$, and Ni$^{2+}$ by extracting with an equal volume of dithizone in CCl$_4$.

**SOLUTION**

From Table 7.9, a quantitative separation of Cu$^{2+}$ from Cd$^{2+}$ and from Ni$^{2+}$ is possible if we acidify the aqueous phase to a pH of less than 1. This pH is greater than the minimum pH for extracting Cu$^{2+}$ and significantly less than the minimum pH for extracting either Cd$^{2+}$ or Ni$^{2+}$. After the extraction of Cu$^{2+}$ is complete, we shift the pH of the aqueous phase to 4.0, which allows us to extract Cd$^{2+}$ while leaving Ni$^{2+}$ in the aqueous phase.

**Table 7.9 Minimum pH for Extracting 99% of an Aqueous Metal Ion Using 4.0 mM Dithizone in CCl$_4$ ($V_{aq} = V_{org}$)**

<table>
<thead>
<tr>
<th>Metal Ion</th>
<th>Minimum pH</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hg$^{2+}$</td>
<td>-8.7</td>
</tr>
<tr>
<td>Ag$^+$</td>
<td>-1.7</td>
</tr>
<tr>
<td>Cu$^{2+}$</td>
<td>-0.8</td>
</tr>
<tr>
<td>Bi$^{3+}$</td>
<td>0.9</td>
</tr>
<tr>
<td>Zn$^{2+}$</td>
<td>2.3</td>
</tr>
<tr>
<td>Cd$^{2+}$</td>
<td>3.6</td>
</tr>
<tr>
<td>Co$^{2+}$</td>
<td>3.6</td>
</tr>
<tr>
<td>Pb$^{2+}$</td>
<td>4.1</td>
</tr>
<tr>
<td>Ni$^{2+}$</td>
<td>6.0</td>
</tr>
<tr>
<td>Tl$^+$</td>
<td>8.7</td>
</tr>
</tbody>
</table>

**7H Separation Versus Preconcentration**

Two common analytical problems are matrix components that interfere with an analyte’s analysis and an analyte with a concentration that is too small to analyze accurately. As we have learned in this chapter, we can use a separation to solve the first problem. Interestingly, we often can use a separation to solve the second problem as well. For a separation in which we recover the analyte in a new phase, it may be possible to increase the analyte’s concentration if we can extract the analyte from a larger volume into a smaller volume. This step in an analytical procedure is known as a **PRECONCENTRATION**.

An example from the analysis of water samples illustrates how we can simultaneously accomplish a separation and a preconcentration. In the gas chromatographic analysis for organophosphorous pesticides in environmental waters, the analytes in a 1000-mL sample are separated from their aqueous matrix by a solid-phase extraction that uses 15 mL of ethyl acetate.\(^{21}\) After the extraction, the analytes in the ethyl acetate have a concentration that is 67 times greater than that in the original sample (assuming the extraction is 100% efficient).

\[\frac{1000 \text{ mL}}{15 \text{ mL}} \approx 67 \times\]

**7I Key Terms**

- centrifugation
- composite sample
- coning and quartering
- convenience sampling
density gradient
centrifugation
dialysis
distillation
distribution ratio
extraction
equilibrium
filtrate
filtration
gross sample
heterogeneous
grab sample
judgmental sampling
homogeneous
laboratory sample
masking
Nyquist theorem
partition coefficient
preconcentration
purge-and-trap
random sampling
recovery
recrystallization
reentrate
sampling plan
secondary equilibrium
selectivity coefficient
separation factor
reaction
Soxhlet extractor
stratified sampling
size exclusion
subsamples
stratification of density
sublimation
supercritical fluid
systematic–judgmental
sampling
systematic sampling
target population

**7J Chapter Summary**

An analysis requires a sample and how we acquire that sample is critical. The samples we collect must accurately represent their target population, and our sampling plan must provide a sufficient number of samples of ap-

---

propriate size so that uncertainty in sampling does not limit the precision of our analysis.

A complete sampling plan requires several considerations, including the type of sample to collect (random, judgmental, systematic, systematic–judgmental, stratified, or convenience); whether to collect grab samples, composite samples, or in situ samples; whether the population is homogeneous or heterogeneous; the appropriate size for each sample; and the number of samples to collect.

Removing a sample from its population may induce a change in its composition due to a chemical or physical process. For this reason, we collect samples in inert containers and we often preserve them at the time of collection.

When an analytical method’s selectivity is insufficient, we may need to separate the analyte from potential interferents. Such separations take advantage of physical properties—such as size, mass or density—or chemical properties. Important examples of chemical separations include masking, distillation, and extractions.

7K Problems

1. Because of the risk of lead poisoning, the exposure of children to lead-based paint is a significant public health concern. The first step in the quantitative analysis of lead in dried paint chips is to dissolve the sample. Corl evaluated several dissolution techniques.\textsuperscript{22} Samples of paint were collected and then pulverized using a Pyrex mortar and pestle. Replicate portions of the powdered paint were taken for analysis. The following table shows results for a paint sample and for a standard reference material. Both samples and standards were digested with HNO\textsubscript{3} on a hot plate.

<table>
<thead>
<tr>
<th>Replicate</th>
<th>% w/w Pb in Sample</th>
<th>% w/w Pb in Standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>5.09</td>
<td>11.48</td>
</tr>
<tr>
<td>2</td>
<td>6.29</td>
<td>11.62</td>
</tr>
<tr>
<td>3</td>
<td>6.64</td>
<td>11.47</td>
</tr>
<tr>
<td>4</td>
<td>4.63</td>
<td>11.86</td>
</tr>
</tbody>
</table>

(a) Determine the overall variance, the variance due to the method and the variance due to sampling. (b) What percentage of the overall variance is due to sampling? (c) How might you decrease the variance due to sampling?

2. To analyze a shipment of 100 barrels of an organic solvent, you plan to collect a single sample from each of 10 barrels selected at random.

\textsuperscript{22} Corl, W. E. Spectroscopy 1991, 6(8), 40–43.
From which barrels should you collect samples if the first barrel is given by the twelfth entry in the random number table in Appendix 14, with subsequent barrels given by every third entry? Assume that entries in the random number table are arranged by rows.

3. The concentration of dissolved $O_2$ in a lake shows a daily cycle from the effect of photosynthesis, and a yearly cycle due to seasonal changes in temperature. Suggest an appropriate systematic sampling plan to monitor the daily change in dissolved $O_2$. Suggest an appropriate systematic sampling plan for monitoring the yearly change in dissolved $O_2$.

4. The data in the following table were collected during a preliminary study of the pH of an industrial wastewater stream.

<table>
<thead>
<tr>
<th>Time (hr)</th>
<th>pH</th>
<th>Time (hr)</th>
<th>pH</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5</td>
<td>4.4</td>
<td>9.0</td>
<td>5.7</td>
</tr>
<tr>
<td>1.0</td>
<td>4.8</td>
<td>9.5</td>
<td>5.5</td>
</tr>
<tr>
<td>1.5</td>
<td>5.2</td>
<td>10.0</td>
<td>6.5</td>
</tr>
<tr>
<td>2.0</td>
<td>5.2</td>
<td>10.5</td>
<td>6.0</td>
</tr>
<tr>
<td>2.5</td>
<td>5.6</td>
<td>11.0</td>
<td>5.8</td>
</tr>
<tr>
<td>3.0</td>
<td>5.4</td>
<td>11.5</td>
<td>6.0</td>
</tr>
<tr>
<td>3.5</td>
<td>5.4</td>
<td>12.0</td>
<td>5.6</td>
</tr>
<tr>
<td>4.0</td>
<td>4.4</td>
<td>12.5</td>
<td>5.6</td>
</tr>
<tr>
<td>4.5</td>
<td>4.8</td>
<td>13.0</td>
<td>5.4</td>
</tr>
<tr>
<td>5.0</td>
<td>4.8</td>
<td>13.5</td>
<td>4.9</td>
</tr>
<tr>
<td>5.5</td>
<td>4.2</td>
<td>14.0</td>
<td>5.2</td>
</tr>
<tr>
<td>6.0</td>
<td>4.2</td>
<td>14.5</td>
<td>4.4</td>
</tr>
<tr>
<td>6.5</td>
<td>3.8</td>
<td>15.0</td>
<td>4.0</td>
</tr>
<tr>
<td>7.0</td>
<td>4.0</td>
<td>15.5</td>
<td>4.5</td>
</tr>
<tr>
<td>7.5</td>
<td>4.0</td>
<td>16.0</td>
<td>4.0</td>
</tr>
<tr>
<td>8.0</td>
<td>3.9</td>
<td>16.5</td>
<td>5.0</td>
</tr>
<tr>
<td>8.5</td>
<td>4.7</td>
<td>17.0</td>
<td>5.0</td>
</tr>
</tbody>
</table>

Prepare a figure showing how the pH changes as a function of time and suggest an appropriate sampling frequency for a long-term monitoring program.

5. You have been asked to monitor the daily fluctuations in atmospheric ozone in the downtown area of a city to determine if there is relationship between daily traffic patterns and ozone levels. (a) Which of the following sampling plans will you use and why: random, systematic, judgmental, systematic–judgmental, or stratified? (b) Do you plan to collect and analyze a series of grab samples, or will you form a single composite sample? (c) Will your answers to these questions change
if your goal is to determine if the average daily ozone level exceeds a threshold value? If yes, then what is your new sampling strategy?

6. The distinction between a homogeneous population and a heterogeneous population is important when we develop a sampling plan. (a) Define homogeneous and heterogeneous. (b) If you collect and analyze a single sample, can you determine if the population is homogeneous or is heterogeneous?

7. Beginning with equation 7.4, derive equation 7.5. Assume that the particles are spherical with a radius of $r$ and a density of $d$.

8. The sampling constant for the radioisotope $^{24}$Na in homogenized human liver is approximately 35 g. (a) What is the expected relative standard deviation for sampling if we analyze 1.0-g samples? (b) How many 1.0-g samples must we analyze to obtain a maximum sampling error of ±5% at the 95% confidence level?

9. Engels and Ingamells reported the following results for the % w/w K$_2$O in a mixture of amphibolite and orthoclase.

<p>| | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>0.247</td>
<td>0.300</td>
<td>0.236</td>
</tr>
<tr>
<td>0.247</td>
<td>0.275</td>
<td>0.212</td>
</tr>
<tr>
<td>0.258</td>
<td>0.311</td>
<td>0.304</td>
</tr>
<tr>
<td>0.258</td>
<td>0.330</td>
<td>0.187</td>
</tr>
</tbody>
</table>

Each of the 12 samples had a nominal mass of 0.1 g. Using this data, calculate the approximate value for $K_s$, and then, using this value for $K_s$, determine the nominal mass of sample needed to achieve a percent relative standard deviation of 2%.

10. The following data was reported for the determination of KH$_2$PO$_4$ in a mixture of KH$_2$PO$_4$ and NaCl.

<table>
<thead>
<tr>
<th>Nominal Mass (g)</th>
<th>Actual Mass (g)</th>
<th>% w/w KH$_2$PO$_4$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.10</td>
<td>0.1039</td>
<td>0.085</td>
</tr>
<tr>
<td>0.1015</td>
<td>0.1012</td>
<td>0.1010</td>
</tr>
<tr>
<td>0.1060</td>
<td>0.0997</td>
<td>0.507</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Nominal Mass (g)</th>
<th>Actual Mass (g)</th>
<th>% w/w KH₂PO₄</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.25</td>
<td>0.2515</td>
<td>0.847</td>
</tr>
<tr>
<td></td>
<td>0.2465</td>
<td>0.598</td>
</tr>
<tr>
<td></td>
<td>0.2770</td>
<td>0.431</td>
</tr>
<tr>
<td></td>
<td>0.2460</td>
<td>0.842</td>
</tr>
<tr>
<td></td>
<td>0.2485</td>
<td>0.964</td>
</tr>
<tr>
<td></td>
<td>0.2590</td>
<td>1.178</td>
</tr>
<tr>
<td>0.50</td>
<td>0.5084</td>
<td>1.009</td>
</tr>
<tr>
<td></td>
<td>0.4954</td>
<td>0.947</td>
</tr>
<tr>
<td></td>
<td>0.5286</td>
<td>0.618</td>
</tr>
<tr>
<td></td>
<td>0.5232</td>
<td>0.744</td>
</tr>
<tr>
<td></td>
<td>0.4965</td>
<td>0.572</td>
</tr>
<tr>
<td></td>
<td>0.4995</td>
<td>0.709</td>
</tr>
<tr>
<td>1.00</td>
<td>1.027</td>
<td>0.696</td>
</tr>
<tr>
<td></td>
<td>0.987</td>
<td>0.843</td>
</tr>
<tr>
<td></td>
<td>0.991</td>
<td>0.535</td>
</tr>
<tr>
<td></td>
<td>0.998</td>
<td>0.750</td>
</tr>
<tr>
<td></td>
<td>0.997</td>
<td>0.711</td>
</tr>
<tr>
<td></td>
<td>1.001</td>
<td>0.639</td>
</tr>
<tr>
<td>2.50</td>
<td>2.496</td>
<td>0.766</td>
</tr>
<tr>
<td></td>
<td>2.504</td>
<td>0.769</td>
</tr>
<tr>
<td></td>
<td>2.496</td>
<td>0.682</td>
</tr>
<tr>
<td></td>
<td>2.496</td>
<td>0.609</td>
</tr>
<tr>
<td></td>
<td>2.557</td>
<td>0.589</td>
</tr>
<tr>
<td></td>
<td>2.509</td>
<td>0.617</td>
</tr>
</tbody>
</table>

(a) Prepare a graph of % w/w KH₂PO₄ vs. the actual sample mass. Is this graph consistent with your understanding of the factors that affect sampling variance. (b) For each nominal mass, calculate the percent relative standard deviation, \( R_{\text{exp}} \), based on the data. The value of \( K_s \) for this analysis is estimated as 350. Use this value of \( K_s \) to determine the theoretical percent relative standard deviation, \( R_{\text{theo}} \), due to sampling. Considering these calculations, what is your conclusion about the importance of indeterminate sampling errors for this analysis? (c) For each nominal mass, convert \( R_{\text{theo}} \) to an absolute standard deviation. Plot points on your graph that correspond to ±1 absolute standard deviations about the overall average % w/w KH₂PO₄ for all samples. Draw smooth curves through these two sets of points. Does the sample appear homogeneous on the scale at which it is sampled?
11. In this problem you will collect and analyze data to simulate the sampling process. Obtain a pack of M&M’s (or other similar candy). Collect a sample of five candies and count the number that are red (or any other color of your choice). Report the result of your analysis as % red. Return the candies to the bag, mix thoroughly, and repeat the analysis for a total of 20 determinations. Calculate the mean and the standard deviation for your data. Remove all candies from the bag and determine the true % red for the population. Sampling in this exercise should follow binomial statistics. Calculate the expected mean value and the expected standard deviation, and compare to your experimental results.

12. Determine the error ($\alpha = 0.05$) for the following situations. In each case assume that the variance for a single determination is 0.0025 and that the variance for collecting a single sample is 0.050. (a) Nine samples are collected, each analyzed once. (b) One sample is collected and analyzed nine times. (c) Five samples are collected, each analyzed twice.

13. Which of the sampling schemes in problem 12 is best if you wish to limit the overall error to less than $\pm 0.30$ and the cost to collect a single sample is $1$ and the cost to analyze a single sample is $10$? Which is the best sampling scheme if the cost to collect a single sample is $7$ and the cost to analyze a single sample is $3$?

14. Maw, Witry, and Emond evaluated a microwave digestion method for Hg against the standard open-vessel digestion method. The standard method requires a 2-hr digestion and is operator-intensive. The microwave digestion is complete in approximately 0.5 hr and requires little monitoring by the operator. Samples of baghouse dust from air-pollution-control equipment were collected from a hazardous waste incinerator and digested in triplicate before determining the concentration of Hg in ppm. Results are summarized in the following two tables.

<table>
<thead>
<tr>
<th>ppm Hg Following Microwave Digestion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample</td>
</tr>
<tr>
<td>--------</td>
</tr>
<tr>
<td>1</td>
</tr>
<tr>
<td>2</td>
</tr>
<tr>
<td>3</td>
</tr>
<tr>
<td>4</td>
</tr>
<tr>
<td>5</td>
</tr>
<tr>
<td>6</td>
</tr>
<tr>
<td>7</td>
</tr>
<tr>
<td>8</td>
</tr>
</tbody>
</table>

Chapter 7 Collecting and Preparing Samples

<table>
<thead>
<tr>
<th>Sample</th>
<th>Replicate 1</th>
<th>Replicate 2</th>
<th>Replicate 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>5.50</td>
<td>5.54</td>
<td>5.40</td>
</tr>
<tr>
<td>2</td>
<td>13.1</td>
<td>12.8</td>
<td>13.0</td>
</tr>
<tr>
<td>3</td>
<td>5.39</td>
<td>5.12</td>
<td>5.36</td>
</tr>
<tr>
<td>4</td>
<td>6.59</td>
<td>6.52</td>
<td>7.20</td>
</tr>
<tr>
<td>5</td>
<td>6.20</td>
<td>6.03</td>
<td>5.77</td>
</tr>
<tr>
<td>6</td>
<td>6.25</td>
<td>5.65</td>
<td>5.61</td>
</tr>
<tr>
<td>7</td>
<td>15.0</td>
<td>13.9</td>
<td>14.0</td>
</tr>
<tr>
<td>8</td>
<td>20.4</td>
<td>16.1</td>
<td>20.0</td>
</tr>
</tbody>
</table>

Does the microwave digestion method yields acceptable results when compared to the standard digestion method?

15. Simpson, Apte, and Batley investigated methods for preserving water samples collected from anoxic (O$_2$-poor) environments that have high concentrations of dissolved sulfide. They found that preserving water samples with HNO$_3$ (a common method for preserving aerobic samples) gave significant negative determinate errors when analyzing for Cu$^{2+}$. Preserving samples by first adding H$_2$O$_2$ and then adding HNO$_3$ eliminated the determinate error. Explain their observations.

16. In a particular analysis the selectivity coefficient, $K_{A,I}$, is 0.816. When a standard sample with an analyte-to-interferent ratio of 5:1 is carried through the analysis, the error when determining the analyte is +6.3%.

(a) Determine the apparent recovery for the analyte if $R_I = 0$. (b) Determine the apparent recovery for the interferent if $R_A = 0$.

17. The amount of Co in an ore is determined using a procedure for which Fe in an interferent. To evaluate the procedure’s accuracy, a standard sample of ore known to have a Co/Fe ratio of 10.2 is analyzed. When pure samples of Co and Fe are taken through the procedure the following calibration relationships are obtained:

\[ S_{Co} = 0.786 \times m_{Co} \quad \text{and} \quad S_{Fe} = 0.699 \times m_{Fe} \]

where $S$ is the signal and $m$ is the mass of Co or Fe. When 278.3 mg of Co are taken through the separation step, 275.9 mg are recovered. Only 3.6 mg of Fe are recovered when a 184.9 mg sample of Fe is carried through the separation step. Calculate (a) the recoveries for Co and Fe; (b) the separation factor; (c) the selectivity ratio; (d) the error if no attempt is made to separate the Co and Fe; (e) the error if the separation step is carried out; and (f) the maximum possible recovery for Fe if the recovery for Co is 1.00 and the maximum allowed error is 0.05%.

---

18. The amount of calcium in a sample of urine is determined by a method for which magnesium is an interferent. The selectivity coefficient, $K_{Ca,Mg}$, for the method is 0.843. When a sample with a Mg/Ca ratio of 0.50 is carried through the procedure, an error of $-3.7\%$ is obtained. The error is $+5.5\%$ when using a sample with a Mg/Ca ratio of 2.0. (a) Determine the recoveries for Ca and Mg. (b) What is the expected error for a urine sample in which the Mg/Ca ratio is 10.0?

19. Using the formation constants in Appendix 12, show that $F^-$ is an effective masking agent for preventing a reaction between $Al^{3+}$ and EDTA. Assume that the only significant forms of fluoride and EDTA are $F^-$ and $Y^{4-}$.

20. Cyanide is frequently used as a masking agent for metal ions. Its effectiveness as a masking agent is better in more basic solutions. Explain the reason for this dependence on pH.

21. Explain how we can separate an aqueous sample that contains $Cu^{2+}$, $Sn^{4+}$, $Pb^{2+}$, and $Zn^{2+}$ into its component parts by adjusting the pH of the solution.

22. A solute, S, has a distribution ratio between water and ether of 7.5. Calculate the extraction efficiency if we extract a 50.0-mL aqueous sample of S using 50.0 mL of ether as (a) a single portion of 50.0 mL; (b) two portions, each of 25.0 mL; (c) four portions, each of 12.5 mL; and (d) five portions, each of 10.0 mL. Assume the solute is not involved in any secondary equilibria.

23. What volume of ether is needed to extract 99.9% of the solute in problem 23 when using (a) 1 extraction; (b) 2 extractions; (c) four extractions; and (d) five extractions.

24. What is the minimum distribution ratio if 99% of the solute in a 50.0-mL sample is extracted using a single 50.0-mL portion of an organic solvent? Repeat for the case where two 25.0-mL portions of the organic solvent are used.

25. A weak acid, HA, with a $K_a$ of $1.0 \times 10^{-5}$ has a partition coefficient, $K_D$, of $1.2 \times 10^3$ between water and an organic solvent. What restriction on the sample’s pH is necessary to ensure that 99.9% of the weak acid in a 50.0-mL sample is extracted using a single 50.0-mL portion of the organic solvent?

26. For problem 25, how many extractions are needed if the sample’s pH cannot be decreased below 7.0?
27. A weak base, B, with a $K_b$ of $1.0 \times 10^{-3}$ has a partition coefficient, $K_D$, of $5.0 \times 10^2$ between water and an organic solvent. What restriction on the sample’s pH is necessary to ensure that 99.9% of the weak base in a 50.0-mL sample is extracted when using two 25.0-mL portions of the organic solvent?

28. A sample contains a weak acid analyte, HA, and a weak acid interferent, HB. The acid dissociation constants and the partition coefficients for the weak acids are $K_{a,HA} = 1.0 \times 10^{-3}$, $K_{a,HB} = 1.0 \times 10^{-7}$, $K_{D,HA} = K_{D,HB} = 5.0 \times 10^2$. (a) Calculate the extraction efficiency for HA and HB when a 50.0-mL sample, buffered to a pH of 7.0, is extracted using 50.0 mL of the organic solvent. (b) Which phase is enriched in the analyte? (c) What are the recoveries for the analyte and the interferent in this phase? (d) What is the separation factor? (e) A quantitative analysis is conducted on the phase enriched in analyte. What is the expected relative error if the selectivity coefficient, $K_{HA,HB}$, is 0.500 and the initial ratio of HB/HA is 10.0?

29. The relevant equilibria for the extraction of I$_2$ from an aqueous solution of KI into an organic phase are shown in Figure 7.32. (a) Is the extraction efficiency for I$_2$ better at higher or at a lower concentrations of I$^-$. (b) Derive an expression for the distribution ratio for this extraction.

30. The relevant equilibria for the extraction of the metal-ligand complex ML$_2$ from an aqueous solution into an organic phase are shown in Figure 7.33. (a) Derive an expression for the distribution ratio for this extraction. (b) Calculate the extraction efficiency when a 50.0-mL aqueous sample that is 0.15 mM in M$^{2+}$ and 0.12 M in L$^-$ is extracted using 25.0 mL of the organic phase. Assume that $K_D$ is 10.3 and that $\beta_2$ is 560.

31. Derive equation 7.32 for the extraction scheme outlined in Figure 7.30.

32. The following information is available for the extraction of Cu$^{2+}$ by CCl$_4$ and dithizone: $K_{D,c} = 7 \times 10^4$; $\beta_2 = 5 \times 10^{22}$; $K_{a,HL} = 3 \times 10^{-5}$; $K_{D,HL} = 1.1 \times 10^4$; and $n = 2$. What is the extraction efficiency if a 100.0-mL sample of an aqueous solution that is $1.0 \times 10^{-7}$ M Cu$^{2+}$ and 1 M in HCl is extracted using 10.0 mL of CCl$_4$ containing $4.0 \times 10^{-4}$ M dithizone (HL)?

33. Cupferron is a ligand whose strong affinity for metal ions makes it useful as a chelating agent in liquid–liquid extractions. The following table provides pH-dependent distribution ratios for the extraction of Hg$^{2+}$, Pb$^{2+}$, and Zn$^{2+}$ from an aqueous solution to an organic solvent.
(a) Suppose you have a 50.0-mL sample of an aqueous solution that contains Hg\(^{2+}\), Pb\(^{2+}\), and Zn\(^{2+}\). Describe how you can separate these metal ions. (b) Under the conditions for your extraction of Hg\(^{2+}\), what percent of the Hg\(^{2+}\) remains in the aqueous phase after three 50.0-mL extractions with the organic solvent? (c) Under the conditions for your extraction of Pb\(^{2+}\), what is the minimum volume of organic solvent needed to extract 99.5\% of the Pb\(^{2+}\) in a single extraction? (d) Under the conditions for your extraction of Zn\(^{2+}\), how many extractions are needed to remove 99.5\% of the Zn\(^{2+}\) if each extraction uses 25.0 mL of organic solvent?

### 7L Solutions to Practice Exercises

**Practice Exercise 7.1**

To reduce the overall variance by improving the method’s standard deviation requires that

\[ s^2 = 5.00 \text{ ppm}^2 = s_{samp}^2 + s_{meth}^2 = (2.1 \text{ ppm})^2 + s_{meth}^2 \]

Solving for \( s_{meth} \) gives its value as 0.768 ppm. Relative to its original value of 1.1 ppm, this is a reduction of \( 3.0 \times 10^1 \% \). To reduce the overall variance by improving the standard deviation for sampling requires that

\[ s^2 = 5.00 \text{ ppm}^2 = s_{samp}^2 + s_{meth}^2 = s_{samp}^2 + (1.1 \text{ ppm})^2 \]

Solving for \( s_{samp} \) gives its value as 1.95 ppm. Relative to its original value of 2.1 ppm, this is reduction of 7.1%.

Click [here](#) to return to the chapter.

**Practice Exercise 7.2**

The analytical method’s standard deviation is \( 1.96 \times 10^{-3} \text{ g/cm}^3 \) as this is the standard deviation for the analysis of a single sample of the polymer. The sampling variance is
Converting the variance to a standard deviation gives $s_{\text{meth}}$ as $3.64 \times 10^{-2} \text{g/cm}^3$.

Click here to return to the chapter.

### Practice Exercise 7.3

To determine the sampling constant, $K_s$, we need to know the average mass of the samples and the percent relative standard deviation for the concentration of olaquindox in the feed. The average mass for the five samples is 0.95792 g. The average concentration of olaquindox in the samples is 23.14 mg/kg with a standard deviation of 2.200 mg/kg. The percent relative standard deviation, $R$, is

$$ R = \frac{s_{\text{amp}}}{\bar{X}} \times 100 = \frac{2.200 \text{ mg/kg}}{23.14 \text{ mg/kg}} \times 100 = 9.507 \approx 9.51 $$

Solving for $K_s$ gives its value as

$$ K_s = mR^2 = (0.95792 \text{ g})(9.507)^2 = 86.58 \text{ g} \approx 86.6 \text{ g} $$

To obtain a percent relative standard deviation of 5.0%, individual samples need to have a mass of at least

$$ m = \frac{K_s}{R^2} = \frac{86.58 \text{ g}}{(5.0)^2} = 3.5 \text{ g} $$

To reduce the sample's mass from 3.5 g to 1 g, we must change the mass by a factor of

$$ \frac{3.5 \text{ g}}{1 \text{ g}} = 3.5 \times $$

If we assume that the sample's particles are spherical, then we must reduce a particle's radius by a factor of

$$ r^3 = 3.5 \times $$

$$ r = 1.5 \times $$

Click here to return to the chapter.

### Practice Exercise 7.4

Because the value of $t$ depends on the number of samples—a result we have yet to calculate—we begin by letting $n_{\text{amp}} = \infty$ and using $t(0.05, \infty)$ for the value of $t$. From Appendix 4, the value for $t(0.05, \infty)$ is 1.960. Our first estimate for $n_{\text{amp}}$ is

$$ n_{\text{amp}} = \frac{t^2 s_{\text{amp}}}{\epsilon^2} = \frac{(1.96)^2(5.0)^2}{(2.5)^2} = 15.4 \approx 15 $$

Letting $n_{\text{amp}} = 15$, the value of $t(0.05, 14)$ from Appendix 4 is 2.145. Recalculating $n_{\text{amp}}$ gives
\[ n_{\text{amp}} = \frac{t^2 s^2_{\text{amp}}}{e^2} = \frac{(2.145)^2 (5.0)^2}{(2.5)^2} = 18.4 \approx 18 \]

Letting \( n_{\text{amp}} = 18 \), the value of \( t(0.05, 17) \) from Appendix 4 is 2.103. Recalculating \( n_{\text{amp}} \) gives
\[ n_{\text{amp}} = \frac{t^2 s^2_{\text{amp}}}{e^2} = \frac{(2.103)^2 (5.0)^2}{(2.5)^2} = 17.7 \approx 18 \]

Because two successive calculations give the same value for \( n_{\text{amp}} \), we need 18 samples to achieve a sampling error of \( \pm 2.5\% \) at the 95\% confidence interval.

Click [here](#) to return to the chapter.

**Practice Exercise 7.5**

If we collect a single sample (cost $20), then we can analyze that sample 13 times (cost $650) and stay within our budget. For this scenario, the percent relative error is
\[
e = t \sqrt{\frac{s^2_{\text{amp}}}{n_{\text{amp}}} + \frac{s^2_{\text{meth}}}{n_{\text{amp}} n_{\text{rep}}}} = 2.179 \sqrt{\frac{0.10}{1} + \frac{0.20}{1 \times 13}} = 0.74
\]
where \( t(0.05, 12) \) is 2.179. Because this percent relative error is larger than \( \pm 0.50\% \), this is not a suitable sampling strategy.

Next, we try two samples (cost $40), analyzing each six times (cost $600). For this scenario, the percent relative error is
\[
e = t \sqrt{\frac{s^2_{\text{amp}}}{n_{\text{amp}}} + \frac{s^2_{\text{meth}}}{n_{\text{amp}} n_{\text{rep}}}} = 2.2035 \sqrt{\frac{0.10}{2} + \frac{0.20}{2 \times 6}} = 0.57
\]
where \( t(0.05, 11) \) is 2.2035. Because this percent relative error is larger than \( \pm 0.50\% \), this also is not a suitable sampling strategy.

Next we try three samples (cost $60), analyzing each four times (cost $600). For this scenario, the percent relative error is
\[
e = t \sqrt{\frac{s^2_{\text{amp}}}{n_{\text{amp}}} + \frac{s^2_{\text{meth}}}{n_{\text{amp}} n_{\text{rep}}}} = 2.2035 \sqrt{\frac{0.10}{3} + \frac{0.20}{3 \times 4}} = 0.49
\]
where \( t(0.05, 11) \) is 2.2035. Because both the total cost ($660) and the percent relative error meet our requirements, this is a suitable sampling strategy.

There are other suitable sampling strategies that meet both goals. The strategy that requires the least expense is to collect eight samples, analyzing each once for a total cost of $560 and a percent relative error of \( \pm 0.46\% \). Collecting 10 samples and analyzing each one time, gives a percent relative error of \( \pm 0.39\% \) at a cost of $700.

Click [here](#) to return to the chapter.
Practice Exercise 7.6
The fluoride ion, F\(^-\), is a suitable masking agent as it binds with Al\(^{3+}\) to form the stable AlF\(_6\)\(^{3-}\) complex, leaving iron in solution.

Click here to return to the chapter.

Practice Exercise 7.7
The relevant reactions and equilibrium constants are
\[
\begin{align*}
\text{Fe}^{2+}(aq) + 3\text{phen}(aq) &= \text{Fe(phen)}^{3+}(aq) \\
\beta_3 &= 5 \times 10^{20}
\end{align*}
\]
\[
\begin{align*}
\text{Fe}^{3+}(aq) + 3\text{phen}(aq) &= \text{Fe(phen)}^{3+}(aq) \\
\beta_3 &= 6 \times 10^{13}
\end{align*}
\]
where \text{phen} is an abbreviation for 1,10-phenanthroline. Because \(\beta_3\) is larger for the complex with \text{Fe}^{2+} than it is for the complex with \text{Fe}^{3+}, 1,10-phenanthroline will bind \text{Fe}^{2+} before it binds \text{Fe}^{3+}. A ladder diagram for this system (Figure 7.34) suggests that an equilibrium \(p(\text{phen})\) between 5.6 and 5.9 will fully complex \text{Fe}^{2+} without any significant formation of the \text{Fe(phen)}^{3+} complex. Adding a stoichiometrically equivalent amount of 1,10-phenanthroline to a solution of \text{Fe}^{2+} is sufficient to mask \text{Fe}^{2+} in the presence of \text{Fe}^{3+}. A large excess of 1,10-phenanthroline, however, decreases \(p(\text{phen})\) and allows for the formation of both metal–ligand complexes.

Click here to return to the chapter.

Practice Exercise 7.8
(a) The solute’s distribution ratio between water and toluene is
\[
D = \frac{[S_{org}]}{[S_{aq}]} = \frac{0.889 \text{ g} \times \frac{1 \text{ mol}}{117.3 \text{ g}} \times \frac{1}{0.00500 \text{ L}}}{(1.235 \text{ g} - 0.889 \text{ g}) \times \frac{1 \text{ mol}}{117.3 \text{ g}} \times \frac{1}{0.01000 \text{ L}}} = 5.14
\]
(b) The fraction of solute remaining in the aqueous phase after one extraction is
\[
(q_{aq})_1 = \frac{V_{aq}}{DV_{org} + V_{aq}} = \frac{20.00 \text{ mL}}{(5.14)(10.00 \text{ mL}) + 20.00 \text{ mL}} = 0.280
\]
The extraction efficiency, therefore, is 72.0%.
(c) To extract 99.9% of the solute requires
\[
(Q_{aq})_n = 0.001 = \left(\frac{20.00 \text{ mL}}{(5.14)(10.00 \text{ mL}) + 20.00 \text{ mL}}\right)^n = (0.280)^n
\]
\[
\log(0.001) = n \log(0.280)
\]
\[
n = 5.4
\]
a minimum of six extractions.

Click here to return to the chapter.
Practice Exercise 7.9

Because the weak base exists in two forms, only one of which extracts into the organic phase, the partition coefficient, $K_D$, and the distribution ratio, $D$, are not identical.

$$K_D = \frac{[B_{org}]}{[B_{aq}]}$$

$$D = \frac{[B_{org}]}{[B_{aq}]_{total}} = \frac{[B_{org}]}{[B_{aq}] + [HB_{aq}]}$$

Using the $K_b$ expression for the weak base

$$K_b = \frac{[OH_{aq}][HB_{aq}^+]}{[B_{aq}]}$$

we solve for the concentration of $HB^{+}$ and substitute back into the equation for $D$, obtaining

$$D = \frac{[B_{org}]}{[B_{aq}] + \frac{K_b \times [B_{aq}]}{[OH_{aq}]}} = \frac{[B_{org}]}{[B_{aq}] \left(1 + \frac{K_b}{[OH_{aq}]}\right)} = \frac{K_D [OH_{aq}]}{[OH_{aq}^+] + K_b}$$

At a pH of 9.0, the $[OH^{-}]$ is $1 \times 10^{-5}$ M and the distribution ratio has a value of

$$D = \frac{K_D [OH_{aq}]}{[OH_{aq}^+] + K_b} = \frac{(5.00)(1.0 \times 10^{-5})}{1.0 \times 10^{-5} + 1.0 \times 10^{-4}} = 0.455$$

After one extraction, the fraction of $B$ remaining in the aqueous phase is

$$q_{aq(1)} = \frac{25.00 \text{ mL}}{(0.455)(50.00 \text{ mL}) + 25.00 \text{ mL}} = 0.524$$

The extraction efficiency, therefore, is 47.6%. At a pH of 9, most of the weak base is present as $HB^+$, which explains why the overall extraction efficiency is so poor.

Click [here](#) to return to the chapter.