An Introduction to Probability and Statistics (statistics 325)

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The entity of primary interest is a group

Statistical methods deal with properties of groups or aggregates. In many applications the entity of primary interest is an actual, physical group (population) of objects. These objects may be animate (*e.g.*, people or animals) or inanimate (*e.g.*, farm field plots, trees, or days). We will refer to the individual objects that comprise the group of interest as **units**. In certain contexts we may refer to the unit as a *population unit*, a sampling unit, an experimental unit, or a treatment unit.

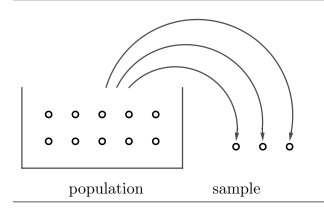
Information about a unit – variables

In order to obtain information about a group of units we first need to obtain information about each of the units in the group. A **variable** is a measurable characteristic of an individual unit. Since our goal is to learn something about the group, we are most interested in the **distribution of the variable**, *i.e.*, the way in which the possible values of the variable are distributed among the units in the group.

The population and the sample

The population is the collection of all of the units that we are interested in. The sample is the subset of the population that we will examine. (We will define a sample more precisely when we discuss random sampling.)

Figure 1.1. Population (box of balls) and sample. A ball represents a population unit. The balls removed from the box represent the sample.



2 1.1 Basic ideas

When the units are actual, physical objects we define the **population** as the collection of all of the units that we are interested in. In most applications it is unnecessary or undesirable to examine the entire population. Thus we define a **sample** as a subset or part of the population for which we have or will obtain data. The collection of observed values of one or more variables corresponding to the individual units in the sample constitute the **data**. Once the data are obtained we can use the distributions of the variables among the units in the sample to characterize the sample itself and to make inferences or generalizations about the entire population, *i.e.*, inferences about the distributions of these variables among the units in the population.

In some applications, such as experimental studies, the population is best viewed as a hypothetical population of values of one or more variables. For example, suppose that we are interested in the effects of an alternative diet on weight gain in some population of experimental animals. We might conduct an experiment by randomly assigning animals to two groups; feeding one group a standard diet and the other group the alternative diet; and then recording the weight gained by each animal over some fixed period of time. In this example we can envision two hypothetical populations of weight gains: The population of weight gains we would have observed if all of the animals were given the standard diet; and, the population of weight gains we would have observed if all of the animals were given the alternative diet.

Information about a group – parameters and statistics

Recall that a **variable** is a measurable characteristic of an individual unit. One way to characterize a group of units is to examine the values of the variable corresponding to all of the units in the group and determine one or more suitable summary values. For example, given a group of adults, we might compute the average age of the group or the proportion who have full-time jobs. A **parameter** is a numerical characteristic of the population. A **statistic** is a numerical characteristic of the sample. That is, a parameter is a number which characterizes a population and a statistic is a number which characterizes a sample.

An illustration is provided in Figure 1.2 for a population of 10 balls and sample of 3 balls. In this figure the characteristic of interest is the color of the ball and the color red (darker shade) is of particular interest. The parameter is the proportion of red balls in the population, 6/10, and the statistic is the proportion of red balls in the sample, 2/3.

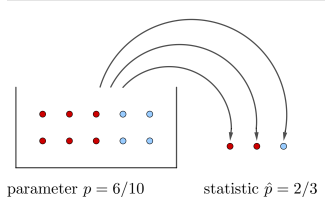


Figure 1.2. Parameter and statistic for a population of 10 balls and sample of 3 balls.

(population proportion red) (sample proportion red)

Example 1.1 NHANES The National Health and Nutrition Examination Survey (NHANES) is a program of studies designed to assess the health and nutritional status of adults and children in the United States. The survey is unique in that it combines interviews and physical examinations. We will use some data from the 2013–2014 NHANES to illustrate the basic ideas we are discussing. For now we will concentrate our attention on some body size measurements for the 5588 adults (age 20 and over) in the 2013–2014 NHANES.

For present purposes we will view this group of N = 5588 adults as the population. In the original context of the survey this is a sample. Thus, for our purposes:

1. A unit is an individual adult.

2. The population is the collection of N = 5588 adults about whom we have information.

3. The sample is a collection of n = 50 individuals (units) which I selected at random from the population of N = 5588 adults.

4. We will consider six variables: The sex of the person (male or female); The age of the person (years); The weight of the person (pounds); The height of the person (inches); The BMI (body mass index) of the person (kg/m^2) ; and, The waist circumference of the person (inches).

4 1.1 Basic ideas

Table 1.1 contains the values of the six variables for the n = 50 people in the sample. The values in a particular row correspond to an individual (one unit).

NULANES	2013-20	$14 \operatorname{sim}$	ble randoi	n sample o	n = 0	0.	
line			moight	hoimht	h ma i	ict	
line	sex	age	weight	height	bmi	waist	
1	male	48	285.34	70.1181	40.9	48.8976	
2	female	80	172.48	63.5433	30.1	41.6535	
3	male	48	186.78	69.8819	26.9	37.5197	
4	male	80	167.2	67.5591	25.8	37.0079	
5	female	20	156.2	67.7953	23.9	32.2835	
6	female	43	196.02	66.1811	31.5	46.8504	
7	male	54	200.86	64.8425	33.7	42.5197	
8	female	24	153.78	63.4252	26.9	40.7874	
9	female	25	122.32	64.1732	20.9	29.0945	
10	male	58	142.34	69.0157	21.1	34.252	
11	female	74	154.22	63.1496	27.2	37.8346	
12	female	74	173.8	61.378	32.5	45.3937	
13	male	78	154.22	67.4016	23.9	33.4646	
14	female	72	89.1	56.9291	19.4	29.1339	
15	male	48	178.2	72.2835	24	36.811	
16	male	64	231.66	62.874	41.3	51.7323	
17	male	41	164.34	68.0315	25	36.7717	
18	female	39	143.88	60.7874	27.4	33.7402	
19	male	49	305.14	72.1654	41.3	missing	
20	male	73	141.24	70.3937	20.1	37.4409	
21	female	67	203.94	62.0079	37.4	44.7244	
22	female	26	101.86	59.8031	20.1	29.0157	
23	female	73	150.7	65.7087	24.6	38.189	
24	male	60	199.32	72.1654	27	40.0394	
25	male	40	206.58	67.4409	32	43.2677	
26	male	27	181.94	67.3622	28.2	38.622	
27	male	62	199.98	67.3622	31.1	43.3858	
28	female	71	176.66	64.6063	29.8	42.9134	
29	female	80	166.1	62.3228	30.1	39.3701	
30	male	39	280.72	71.9291	38.2	52.0472	
31	female	48	127.6	61.4173	23.8	31.2598	
32	male	46	156.64	67.7953	24	missing	
33	female	80	147.4	61.3386	27.6	40.7087	
34	female	35	183.04	65.2362	30.3	41.2205	
35	female	57	140.36	62.2047	25.6	34.9213	
		CO	ontinued	below			

Table 1.1 NHANES 2013-2014 simple random sample of n = 50.

line	sex	age	weight	height	bmi	waist	
36	female	40	170.06	65.9055	27.6	35.748	
37	female	56	182.6	59.685	36.1	39.4882	
38	male	34	192.5	70.3937	27.4	37.9921	
39	male	75	163.9	64.7244	27.6	48.8189	
40	female	76	147.84	60.4724	28.5	36.6535	
41	male	57	254.32	70.7087	35.8	43.8189	
42	male	77	187.22	69.2126	27.5	40.1181	
43	male	45	179.3	70.7087	25.3	36.5354	
44	male	48	249.48	67.126	39	51.063	
45	female	47	214.06	64.6457	36.1	42.5984	
46	male	72	205.26	70.3543	29.2	42.8346	
47	male	80	127.38	66.4567	20.3	missing	
48	male	24	199.32	71.2205	27.7	40.6299	
49	male	80	162.8	66.7323	25.8	39.4094	
50	male	21	119.24	66.6929	18.9	26.3386	

Table 1.1 continuation of NHANES 2013-2014 simple random sample of n = 50.

Some parameters and statistics (population and sample means) for the numerical variables in this example are given in Table 1.2. With respect to the categorical variable sex of the person; there are 2919 females and 2669 males in the population and there are 22 females and 28 males in the sample. This gives the (parameter) population percentage female as 52.24% and the (statistic) sample percentage female as 44%.

variable	parameter population mean	statistic sample mean
age	49.15	54.70
weight	179.22	177.94
height	65.77	66.11
bmi	29.10	28.53
waist	39.05	39.47

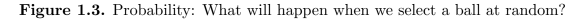
 Table 1.2 Population and sample means for the HANES example.

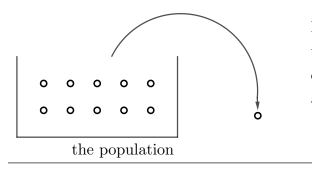
Which direction? Probability versus statistics

Probability theory is used to model the outcomes of random processes. The basic probability situation is illustrated in Figure 1.3. Here we know all there is to know about the

6 1.2 Sampling

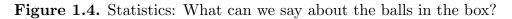
characteristics of the balls in the box and we want to make a statement about what will happen when we select a ball at random from the box and examine it. For example, for the box of Figure 1.2, where 60% of the balls in the box are red, if we select one ball at random, there is a 60% chance (probability) that it will be red.

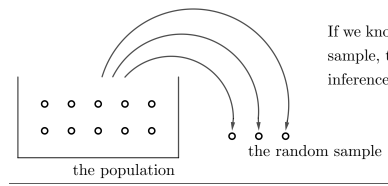




If we know all about the balls in the box, then we can assign probabilities to the outcomes we may observe when we select a ball at random.

Statistical theory is used to make inferences from a random sample to a population. The basic statistics situation is illustrated in Figure 1.4. Here we know all there is to know about the characteristics of the balls in the random sample and we want to make a statement about what we would find if we examined all of the balls in the box (the entire population).





If we know all about the balls in the random sample, then we can use statistics to make inferences about the balls in the box.

1.2 Sampling

toc

A sampling study is conducted by selecting a random sample of units from a population, observing the values of a variable for the units in the sample, and then making inferences or generalizations about the population. More specifically, the distribution of the values of the variable among the units in a random sample is used to make inferences about the distribution of the variable among the units in the population. The first consideration in planning or interpreting the results of a sampling study is the determination of exactly which units could be in the sample. The second consideration concerns the proper selection of the units which constitute the sample. We will discuss these considerations in more depth in the rest of this section.

Sampling is the process of obtaining a sample from a population. Our ultimate goal is to use the sample (which we will examine) to make inferences about the population (which we will not examine in its entirety). If the sample is selected from the population in an *appropriate fashion*, then we can use the information in the sample to make reliable and quantifiable inferences about the population. When the sample is obtained we will use the distribution of the variable among the units in the sample to make inferences about the distribution of the variable among the units in the population. If the distribution of the variable in the sample was exactly the same as the distribution of the variable in the population, then it would be easy to make inferences about the population; but, this is clearly too much to ask. Therefore we need to determine how to select a sample so that the sample is representative of the population.

The first step in deciding whether a method of choosing a sample will yield a representative sample requires a distinction between two populations. Before we obtain a sample we need to decide exactly which population we are interested in. The **target population** is the collection of all of the units that we want to make inferences about. We then need to determine which population our sample actually comes from. The **sampled population** is the collection of all of the units that *could* be in the sample. Notice that the sampled population is determined by the method used to select the sample.

Ideally the sampling method is chosen so that the sampled population is exactly the same as the target population and we can refer to this collection as the population. In practice, there may be some differences between the target population and the sampled population. When the sampled population is not identical to the target population we cannot be confident that the sample (which comes from the sampled population) will be representative of the target population. Furthermore, we cannot be confident that the statistic (which is based on a sample from the sampled population) will be suitable for inference about the parameter (which corresponds to the target population).

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If there is a difference between the sampled population and the target population, in the sense that the distribution of the variable in the sampled population is different from the distribution of the variable in the target population, then a sample (obtained from the sampled population) is said to be **biased** for making inferences about the target population. If we use a biased sample to make inferences about the target population, the resulting inferences will not be appropriate. For example, a statistic based on a biased sample, may provide a suitable estimate of the corresponding parameter in the sampled population; but, it may not provide a suitable estimate of the corresponding parameter in the target population. Therefore, if the sampled population is different from the target population, then we must modify our goals by redefining the target population or we must change the sampled population by modifying our sampling method, since we want these two populations to be the same so that our inferences will be valid for our target population. It may be possible to change the method of obtaining our sample so that all of the units in the target population could be in our sample and these two populations are the same. If it is not possible to change the sampling method, then we must change our goals by restricting our inferences to the sampled population. In any case, once a sampling method has been chosen, the sampled population is determined and we should restrict our inferences to this sampled population. In conclusion, when making inferences from a sample we must carefully consider the restrictions imposed by the sampling method, since statistical theory can only justify inferences about the sampled population.

Assuming that we have defined a method of selecting a sample so that the sampled population is the same as the target population, we next need to consider exactly how we should select the units that constitute the sample. Since we are assuming that the sampled and target populations are the same, we do not need to worry about the type of bias described above. However, we might introduce bias if we do not select the units for the sample in an appropriate fashion. The approach to sampling that we will adopt is called random sampling. The idea behind random sampling is to eliminate potential bias (intentional or unintentional) from the selection process by using *impersonal random chance* to select the sample. In addition to eliminating bias random sampling also provides the basis for theoretical justification and quantification of inferences based on the sample.

All of the sampling situations we consider can be viewed as being abstractly the same as the simple situation of selecting a sample of balls from a box of balls. This scenario was illustrated in Figure 1.1. The simplest type of random sample is called a simple random sample. A **simple random sample of size n** is a sample of n units selected from the population in such a way that every possible sample of n units has the same chance of being selected. This definition of a simple random sample can be refined to distinguish between two versions of simple random samples. If we require that the possible samples of n units are such that a particular unit can occur at most once in a sample, then we refer to the sample as being a **simple random sample of size n**, **selected without replacement**. On the other hand, if we allow a particular unit to occur more than once in the sample, then we refer to the sample as a **simple random** sample of size n, selected with replacement.

To obtain a simple random sample of size n from the balls in our box, we first mix the balls in the box and select one ball at random (so that each ball in the box has the same chance of being selected). We then determine the value of the variable for the selected ball giving us the value of the variable for one of the balls in our random sample. If we are sampling with replacement we return the ball to the box before the next draw. If we are sampling without replacement we do not return the ball to the box. We then mix the balls in the box and continue this process of selecting a ball from the box at random until n balls have been selected. These n balls (or the values of the variable for these balls) form the simple random sample of size n.

If the population from which we wish to select a random sample is not too large, then it is possible to envision actually labeling each unit in the population, placing these labels on a collection of balls, placing these labeled balls in a box, and selecting a simple random sample of these balls as described above. In fact, state lotteries, where a simple random sample of numbers is selected from a collection of allowable numbers (the units), are conducted in this way. If you have ever observed the complicated mechanisms used to select winning lottery numbers, you know that it is difficult to convince people that a method of "drawing balls from a box" yields a proper simple random sample. For most purposes it is best to use a computer or calculator to select a simple random sample. The computer will simulate the process of drawing balls at random from a box.

When we take a simple random sample, all of the possible samples have the same chance of being selected. There are situations where it is not appropriate for all of the possible samples to have the same chance of being selected. Suppose that there are two or more identifiable subsets of the population (subpopulations). If we obtain a simple random

10 1.2 Sampling

sample from the whole population, then it is possible for the resulting sample to come entirely from one of the subpopulations so that the sample does not contain any units from one or more of the subpopulations. If we know or suspect that the distribution of the variable of interest varies among the subpopulations, then a sample which does not contain any units from some of the subpopulations will not be representative of the whole population. Therefore, in a situation like this we should not use a simple random sample to make inferences about the whole population. Instead we should use a more complex kind of random sample. One possibility is to use a sampling method known as **stratified random sampling** which is described below in the context of a simple example.

Suppose we wish to estimate the proportion of all registered voters in the United States who favor a particular candidate in an upcoming presidential election. We might expect there to be differences in the proportion of registered voters who favor this candidate among the various states. For example, we might expect support for this candidate to be particularly strong in his or her home state. Because we are interested in the proportion of all registered voters in the United States who favor this candidate, we want to be sure that all of the states are represented fairly in our sample.

We can use the states to define **strata** (subpopulations), take a random sample from each state (stratum), and then combine these samples to get a sample that is representative of the entire country. This is an example of a stratified random sample. The simplest type of **stratified random sample** is obtained as described in the following three steps.

1. Divide the population into appropriate strata (subpopulations).

2. Obtain a simple random sample within each stratum.

3. Combine these simple random samples to get the stratified random sample from the whole population.

To obtain a representative sample in the opinion poll example, we would need to determine the number of registered voters in each state and select simple random samples of sizes that are proportional to the numbers of registered voters in the states.

1.3 Experimentation

An experimental study differs from a sampling study in that the units used in the experimental study are manipulated and the responses of the units to this experimental manipulation are recorded. For an experimental study the relevant population or populations are hypothetical populations of values of the variable defined by the experimental treatment(s) and corresponding to all of the units available for use in the experiment. That is, the relevant population(s) is the population(s) of values of the variable which would be observed if all of the available units were subjected to the experimental treatment(s). In the context of a comparative experiment we cannot properly quantify inferences unless the units are assigned to the treatments being compared using an appropriate method of random assignment. This random assignment of units to treatments is analogous to the random sampling of a sampling study.

In an **experimental study** we manipulate the units and observe their response to this manipulation. In the experimental context, a particular combination of experimental conditions is known as a **treatment**. The purpose of an experiment is to obtain information about how the units in the population would respond to a treatment; or, to compare the responses of the units to two or more treatments. The response of a unit to a particular treatment is determined by measuring the value of a suitable **response variable**.

The steps involved in conducting a simple experimental study based on a random sample are summarized below.

1. Obtain a random sample of units from the population of interest.

2. Subject the units in the sample to the experimental treatment of interest.

3. Obtain the data. That is, determine the values of the response variable for the units in the sample.

4. Use the data to make inferences about the how the units in the population would respond to the treatment. More specifically, use the distribution of the response variable in the sample to make inferences about the distribution of the response variable in the population from which the sample was taken. In this context it may be easiest to think of the population as the hypothetical population of values of the response variable which would result if all of the units in the population were subjected to the treatment.

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We will now discuss the basic ideas of experimentation in more detail in the context of a simple hypothetical experiment. Suppose that a new drug has been developed to reduce the blood pressure of hypertensive patients. The treatment of interest is the administration of the new drug to a hypertensive patient. The change in a patient's blood pressure will be used as the response variable. For this example the plan of the simple experiment described above is summarized in the steps below.

1. Obtain a random sample of n hypertensive patients.

- 2. Measure the blood pressure of each patient before the new drug is administered.
- 3. Administer the new drug to each of these patients.

4. After a suitable period of time, measure the blood pressure of each patient.

5. For each patient determine the change in his or her blood pressure by computing the difference between the patient's blood pressure before the drug was administered and the patient's blood pressure after the new drug was administered. This change or difference will serve as the response variable for assessing the effects of the new drug. In this example the relevant population is the hypothetical population of changes in blood pressure that we would observe if all of the hypertensive patients in the population from which the sample was selected had been subjected to this experiment.

Suppose that we actually conducted this experiment. Furthermore, suppose that the data indicate that the hypertensive patients' blood pressures tend to decrease after they are given the new drug, *i.e.*, suppose that the data indicate that most of the patients experienced a meaningful reduction in blood pressure. We can conclude that there is an association between the new drug and a reduction in blood pressure. This association is clear, since the patients (as a group) tended to experience a decrease in blood pressure after they received the new drug. Can we conclude that the new drug caused this decrease in blood pressure? The support for the contention that the new drug caused the decrease in blood pressure is not so clear. In addition to the new drug there may be other factors associated with the observed decrease in blood pressure. For example, the decrease in blood pressure might be explained, in whole or in part, as the physical manifestation of the psychological effect of receiving medication. In other words, we might observe a similar decrease in blood pressure if we administered a placebo to the patients instead of the new drug. It is also possible that some other aspects of the experimental protocol are

affecting the patients' blood pressures. The way that this experiment is being conducted does not allow us to separate out the effects of the many possible causes of the decrease in blood pressure. If we hope to establish a cause and effect relationship between taking the new drug and observing a decrease in blood pressure, then we need to use a comparative experiment.

In a randomized comparative experiment baseline data is obtained at the same time as the data concerning the treatment of interest. This is done by randomly dividing the available units (patients) into two or more groups and comparing the responses for these groups. In the drug example there is one treatment of interest, administer the new drug. Therefore, in this situation we only need two groups, a control group and a treatment group. The units (patients) in the **control group** do not receive the treatment (do not receive the new drug). The units (patients) in the **treatment group** do receive the treatment (do receive the drug). During the course of the experiment we need to keep all aspects of the experiment, other than the treatment itself, as similar as possible for all of the units in the study. The idea is that, if the only difference between the units in the control group and the units in the treatment group is that the units in the treatment group received the treatment, then any observed differences between the responses of the two groups must be caused by the treatment. In the drug example it would be a good idea to administer a placebo to the patients in the control group, so that they do not know that they did not receive the new drug. It would also be a good idea to "blind" the patients and the people administering the drug or placebo by not telling them which patients are receiving the placebo and which patients are receiving the new drug. The purpose of such blinding is to eliminate intentional or unintentional effects due to patient or administrative actions which might affect a patient's response. The steps for conducting such a randomized comparative experiment are given below.

1. Randomly divide the group of available patients into two groups: a group of n_1 patients to serve as the control group and a group of n_2 patients to serve as the treatment group. These two groups are random samples of sizes n_1 and n_2 from the group of available patients. The samples sizes n_1 and n_2 may be different.

2. Administer the placebo to the patients in the control group and administer the new drug to the patients in the treatment group.

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3. Obtain the data. That is, measure the response variable for each of the $n_1 + n_2$ patients in the experiment. For example, we could determine the change (difference) in each patient's blood pressure as measured before and after administration of the placebo or new drug.

4. Compare the responses of the patients in the treatment group to the responses of the patients in the control group and make inferences about the effects of the new drug versus the placebo.

In this example there are two hypothetical populations of changes in blood pressure. The hypothetical population of changes in blood pressure that we would observe if all of the available hypertensive patients were subjected to this experiment and given the placebo and the hypothetical population of changes in blood pressure that we would observe if all of the available hypertensive patients were subjected to this experiment and given the new drug. Notice that, strictly speaking, our inferences in this example only apply to the hypertensive patients who were available for assignment to the groups used in the experiment. If we want to make inferences about a larger population of hypertensive patients, then the group of available patients used in the study should form a random sample from this larger population.

The experiment described above is designed to compare the effects of the new drug to the effects of a placebo. Suppose that we wanted to compare the effects of the new drug to the effects of a standard drug. To make this comparison we could design the experiment with three groups: a control group, a treatment group for the new drug, and a treatment group for the standard drug. If our only goal is to compare the two drugs (treatments), then we could eliminate the placebo control group and run the experiment with the two treatment groups alone.

1.4 Problems

toc

The problem sets and solutions can be found on the book web page at https://www.ucs.louisiana.edu/~jcb0773/.

2 Descriptive Statistics

2.1 Tabular and graphical summary

Consider the problem of using data to learn something about the characteristics of the group of units which comprise the sample. Recall that the distribution of a variable is the way in which the possible values of the variable are distributed among the units in the group. A variable is chosen to measure some characteristic of the units in the group; therefore, the distribution of a variable contains all of the available information about the characteristic (as measured by that variable) for the group. Other variables, either alone or in conjunction with the primary variable, may also contain information about the characteristic of interest. A meaningful summary of the distribution of a variable provides an indication of the overall pattern of the distribution and serves to highlight possible unusual or particularly interesting aspects of the distribution.

Generally speaking, it is hard to tell much about the distribution of a variable by examining the data in raw form. Therefore, the first step in summarizing the distribution of a variable is to tabulate the frequencies with which the possible values of the variable appear in the sample. A **frequency distribution** is a table listing the possible values of the variable and their frequencies (counts of the number of times each value occurs). A frequency distribution provides a decomposition of the total number of observations (the sample size) into frequencies for each possible value. In general, especially when comparing two distributions based on different sample sizes, it is preferable to provide a decomposition in terms of relative frequencies. A **relative frequency distribution** is a table listing the possible values of the variable along with their relative frequencies (proportions). A relative frequency distribution provides a decomposition of the total relative frequencies for each possible values area to the variable along with their relative frequencies (proportions). A

Many aspects of the distribution of a variable are most easily communicated by a graphical representation of the distribution. The basic idea of a graphical representation of a distribution is to use area to represent relative frequency. The total area of the graphical representation is taken to be one (100%) and sections with area equal to the relative frequency (percentage) of occurrence of a value are used to represent each possible value of the variable.

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2.2 Types of variables

When discussing the distribution of a variable we need to consider the structure possessed by the possible values of the variable. This leads to the following classification of variables into four basic types.

A qualitative variable (categorical variable) classifies a unit into one of several possible categories. The possible values of a qualitative variable are names for these categories. We can distinguish between two types of qualitative variables. A qualitative variable is said to be **nominal** if there is no inherent ordering among its possible values. If there is an inherent ordering of the possible values of a qualitative variable, then it is said to be **ordinal**. For example the sex (female or male) of a college student is nominal while the classification (freshman, sophomore, junior, senior) is ordinal.

A quantitative variable (numerical variable) assigns a meaningful numerical value to a unit. Because the possible values of a quantitative variable are meaningful numerical quantities, they can be viewed as points on a number line. If the possible values of a quantitative variable correspond to isolated points on the number line, then there is a discrete jump between adjacent possible values and the variable is said to be a **discrete** quantitative variable. The most common example of a discrete quantitative variable is a count such as the number of babies in a litter of animals or the number of plants in a field plot. If there is a continuous transition from one value of the variable to the next, then the variable is said to be a **continuous** quantitative variable. For a continuous quantitative variable there is always another possible value between any two possible values, no matter how close together the values are. In practice all quantitative variables are discrete in the sense that the observed values are rounded to a reasonable number of decimal places. Thus the distinction between a continuous quantitative variable and a discrete quantitative variable is often more conceptual than real. If a value of the variable represents a measurement of the size of a unit, such as height, weight, or length, or the amount of some quantity, then it is reasonable to think of the possible values of the variable as forming a continuum of values on the number line and to view the variable as continuous.

We can also classify variables with respect to the roles they play in a statistical analysis. That is, we can distinguish between response variables and explanatory variables. A **response variable** is a variable that measures the response of a unit to natural or experimental stimuli. A response variable provides us with a measurement or observation that characterizes a unit with respect to a characteristic of primary interest. An **explanatory variable** is a variable that can be used to explain, in whole or in part, how a unit responds to natural or experimental stimuli. This terminology is clearest in the context of an experimental study. Consider an experiment where a unit is subjected to a treatment (some combination of conditions) and the response of the unit to the treatment is recorded. A variable that describes the treatment conditions is called an explanatory variable, since it may be used to explain the outcome of the experiment. A variable that measures the outcome of the experiment is called a response variable, since it measures the response of the unit to the treatment. An explanatory variable may also be used to subdivide a group so that the distributions of a response variable can be compared among subgroups. **2.3** Qualitative data **toc**

Bar graphs are used to summarize the distribution of a qualitative variable. A **bar graph** consists of a collection of bars (rectangles) such that the combined area of all the bars is one (100%) and the area of a particular bar is the relative frequency of the corresponding value of the variable. Two other common forms for such a graphical representation are segmented bar graphs and pie graphs. A **segmented bar graph** consists of a single bar of area one (100%) that is divided into segments with a segment of the appropriate area for each observed value of the variable. A segmented bar graph can be obtained by joining the separate bars of a bar graph. If the bar of the segmented bar graph is replaced by a disk, the result is a pie graph or pie chart. In a **pie graph** or pie chart the interior of a disk (the pie) is used to represent the total area of one (100%); and the pie is divided into slices of the appropriate area or relative frequency, with one slice for each observed value of the variable.

Example 2.1 Immigrants to the United States. The data concerning immigrants admitted to the United States summarized by decade as raw frequency distributions in Table 2.1 were taken from the 2002 Yearbook of Immigration Statistics, USCIS, (www.uscis.gov). Immigrants for whom the country of last residence was unknown are omitted. For this example a unit is an individual immigrant and these data correspond to a census of the entire population of immigrants, for whom the country of last residence was known, for these decades. Because the region of last residence of an immigrant is a nominal variable and its values do not have an inherent ordering, the values in the bar graphs (and relative frequency distributions) in Figure 2.1 have been arranged so that the percentages for the 1931–1940 decade are in decreasing order.

Table 2.1 Region of last residence for immigrants to USA.				
	period			
region	1931 - 1940	1961 - 1970	1991 - 2000	
Europe	$347,\!566$	$1,\!123,\!492$	$1,\!359,\!737$	
Asia	$16,\!595$	$427,\!692$	2,795,672	
North America	130,871	886, 891	2,441,448	
Caribbean	15,502	470,213	978,787	
Central America	$5,\!861$	$101,\!330$	$526,\!915$	
South America	$7,\!803$	$257,\!940$	$539,\!656$	
Africa	1,750	28,954	354,939	
Oceania	$2,\!483$	$25,\!122$	55,845	
total	$528,\!431$	$3,\!321,\!634$	9,052,999	

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Two aspects of the distributions of region of origin of immigrants which are apparent in these bar graphs are: The decrease in the proportion of immigrants from Europe; and, the increase in the proportion of immigrants from Asia. In 1931–1940 a large majority (65.77%) of the immigrants were from Europe but for the later decades this proportion steadily decreases. On the other hand, the proportion of Asians (only 3.14% in 1931-1940) steadily increases to 30.88% in 1991–2000. Also note that the proportion of immigrants from North America is reasonably constant for these three decades. The patterns we observe in these distributions may be attributable to several causes. Political, social, and economic pressures in the region of origin of these people will clearly have an impact on their desire to immigrate to the US. Furthermore, political pressures within the US have effects on immigration quotas and the availability of visas.

2.3 Qualitative data 19

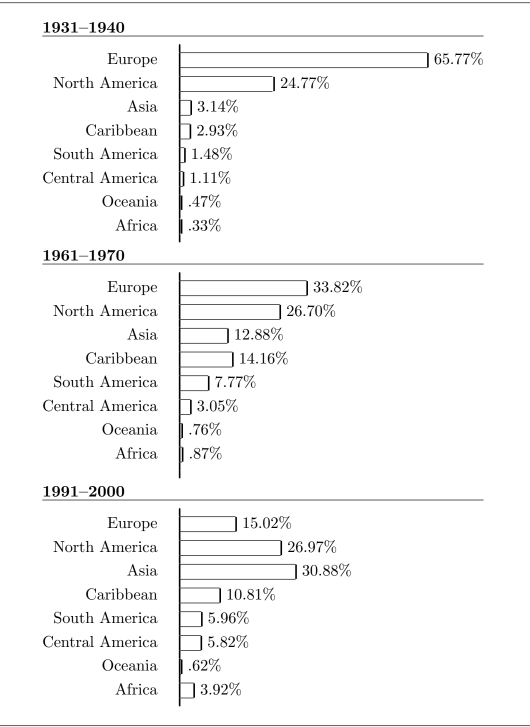


Figure 2.1 Region of last residence for immigrants to USA, by decade.

2.4 Quantitative data

The tabular representations used to summarize the distribution of a discrete quantitative variable, *i.e.*, the frequency and relative frequency distributions, are defined the same as they were for qualitative data. Since the values of a quantitative variable can be viewed as points on the number line, we need to indicate this structure in a tabular representation. In the frequency or relative frequency distribution the values of the variable are listed in order and all possible values within the range of the data are listed even if they do not appear in the data.

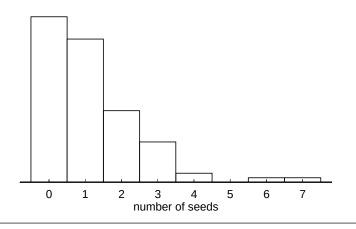
We will use a graphical representation called a histogram to summarize the distribution of a discrete quantitative variable. Like the bar graph we used to represent the distribution of a qualitative variable, the histogram provides a representation of the distribution of a quantitative variable using area to represent relative frequency. A **histogram** is basically a bar graph modified to indicate the location of the observed values of the variable on the number line. For ease of discussion we will describe histograms for situations where the possible values of the discrete quantitative variable are equally spaced (the distance between any two adjacent possible values is always the same). We will use the following weed seed example to illustrate the methodology.

Example 2.2 Weed seeds. C. W. Leggatt counted the number of seeds of the weed potentilla found in 98 quarter-ounce batches of the grass Phleum praetense. This example is taken from Snedecor and Cochran, Statistical Methods, Iowa State, (1980), 198; the original source is C. W. Leggatt, Comptes rendus de l'association international d'essais de semences, 5 (1935), 27. The 98 observed numbers of weed seeds, which varied from 0 to 7, are summarized in the relative frequency distribution of Table 2.2 and the histogram of Figure 2.2. In this example a unit is a batch of grass and the number of seeds in a batch is a discrete quantitative variable with possible values of 0, 1, 2, ...

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Table 2.2 Weed seeds relative frequency distribution.			
number	frequency	relative	
of seeds		frequency	
0	37	.3776	
1	32	.3265	
2	16	.1633	
3	9	.0918	
4	2	.0204	
5	0	.0000	
6	1	.0102	
7	1	.0102	
total	98	1.0000	

Figure 2.2 Histogram for number of weed seeds.



Consider the histogram for the number of weed seeds in a batch of grass of Figure 2.2. This histogram is made up of rectangles of equal width, centered at the observed values of the variable. The heights of these rectangles are chosen so that the area of a rectangle is the relative frequency of the corresponding value of the variable. There is not a gap between two adjacent rectangles in the histogram unless there is an unobserved possible value of the variable between the corresponding adjacent observed values. For this example there is a gap at 5 since none of the batches had exactly 5 weed seeds.

In this histogram we are using an interval of values on the number line to indicate a single value of the variable. For example, the rectangle centered over 1 in the histogram of Figure 2.2 represents the relative frequency that a batch of grass contains exactly 1 weed seed;

22 2.4 Quantitative data

but, its base extends from .5 to 1.5 on the number line. Because it is impossible for the number of weed seeds to be strictly between 0 and 1 or strictly between 1 and 2, we are identifying the entire interval from .5 to 1.5 on the number line with the actual value of 1. This identification of an interval of values with the possible value at the center of the interval eliminates gaps in the histogram that would incorrectly suggest the presence of unobserved, possible values.

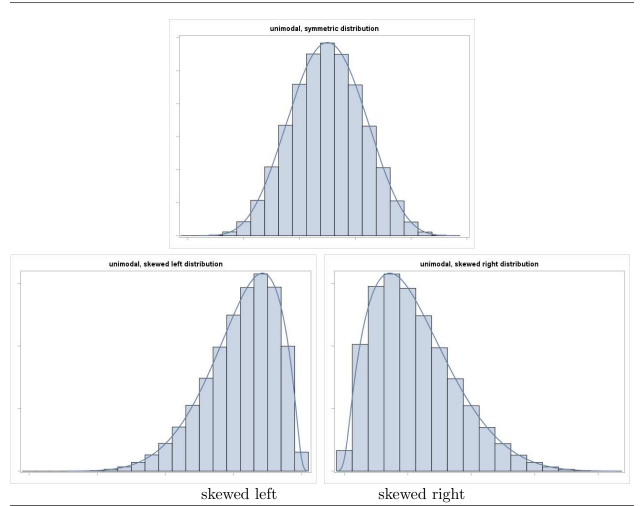
The histogram for the distribution of the number of weed seeds in Figure 2.2 has a mound shaped appearance with a single peak at zero, indicating that the most common number of weed seeds is zero. In fact, 37.76% of the batches of grass contain no weed seeds. Among the batches that do contain weed seeds we see that 32.65% contain one weed seed and 16.33% contain two. Thus, 86.74% of the 98 batches of grass contain two or fewer weed seeds and 95.92% contain three or fewer weed seeds. In summary, the majority of these batches of grass have a small number of weed seeds; but, there are a few batches with relatively high numbers of weed seeds.

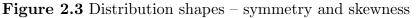
The histogram of Figure 2.2, or the associated distribution, is not symmetric. That is, the histogram (distribution) is not the same on the left side (smaller values) as it is on the right side (larger values). This histogram or distribution is said to be skewed to the right. The concept of a distribution being skewed to the right is often explained by saying that the right "tail" of the distribution is "longer" than the left "tail". That is, the area in the histogram is more spread out along the number line on the right than it is on the left. For this example, the smallest 25% of the observed values are zeros and ones while the largest 25% of the observed values ranging from two to seven. In the present example we might say that there is essentially no left tail in the distribution.

The number of weed seeds histogram provides an example of a very common type of histogram (distribution) which is mound shaped and has a single peak. (A distribution with a single peak is said to be unimodal.) This type of distribution arises when there is a single value (or a few adjacent values) which occurs with highest relative frequency, causing the histogram to have a single peak at this location, and when the relative frequencies of the other values taper off (decrease) as we move away from the location of the peak.

Three examples of mound shaped distributions with a single peak are provided in Figure 2.3. For these illustrations a smooth curve is used to indicate the shape of the histogram. The **symmetric** distribution is such that the histogram has two mirror image halves. The

skewed distributions are more spread out along the number line on one side (the direction of the skewness) than they are on the other side.





There is a fundamental difference between summarizing and describing the distribution of a discrete quantitative variable and summarizing and describing the distribution of a continuous quantitative variable. Since a continuous quantitative variable has an infinite number of possible values, it is not possible to list all of these values. Therefore, some changes to the tabular and graphical summaries used for discrete variables are required.

In practice, the observed values of a continuous quantitative variable are discretized, *i.e.*, the values are rounded so that they can be written down. Therefore, there is really no difference between summarizing the distribution of a continuous variable and summarizing

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the distribution of a discrete variable with a large number of possible values. In either case, it may be impossible or undesirable to actually list all of the possible values of the variable within the range of the observed data. Thus, when summarizing the distribution of a continuous variable, we will group the possible values into intervals.

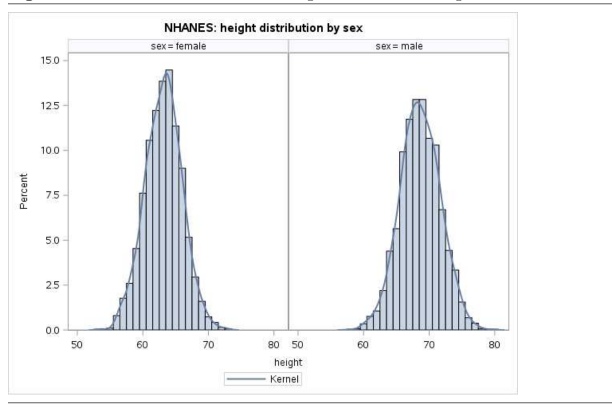


Figure 2.4 NHANES 2013–2014 adult height distribution histograms.

Example 1.1 NHANES (revisited). We will use some data from the 2013–2014 NHANES to illustrate the basic ideas we are discussing. For this application we will use all of the 5588 adults (age 20 and over) in the 2013–2014 NHANES for whom a height or weight measurement is available. This group forms a random sample from the population of all adults (age 20 and over) in the US at the time of the survey. Histograms for the heights and weights of the adult participants in the 2013–2014 NHANES, grouped by sex, are given in Figures 2.4 and 2.5. A smooth version of each histogram (smooth curve) is also provided. The height distributions are both unimodal and reasonably symmetric. The weight distributions are both unimodal and skewed to the right. We will discuss these distributions in more depth shortly.

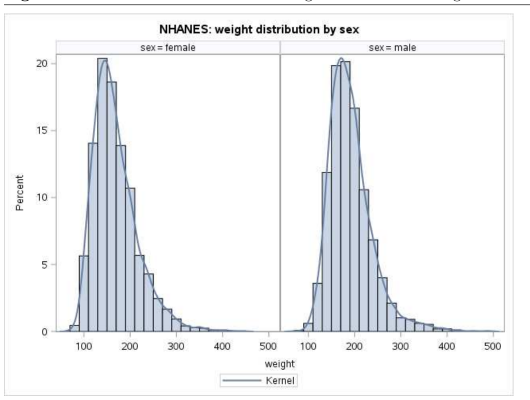


Figure 2.5 NHANES 2013–2014 adult weight distribution histograms.

Notice that the data have been grouped into intervals in order to construct these histograms. For the height distributions the intervals are of length one inch. For the weight distributions the intervals are of length 20 pounds. (In the context of histograms these intervals are also know as bins.) For example, in the weight histograms the area of the rectangle centered over 100 is the proportion of the individuals in the group who had a weight between 90 and 110 pounds.

2.5 Numerical summary

For many purposes a few well-chosen numerical summary values (statistics) will suffice as a description of the distribution of a quantitative variable. A **statistic** is a numerical characteristic of a sample. More formally, a statistic is a numerical quantity computed from the values of a variable, or variables, corresponding to the units in a sample. Thus a statistic serves to quantify some interesting aspect of the distribution of a variable in a sample. Summary statistics are particularly useful for comparing and contrasting the distribution of a variable for two different samples.

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If we plan to use a small number of summary statistics to characterize a distribution or to compare two distributions, then we first need to decide which aspects of the distribution are of primary interest. If the distributions of interest are essentially mound shaped with a single peak (unimodal), then there are three aspects of the distribution which are often of primary interest. The first aspect of the distribution is its location on the number line. Generally, when speaking of the location of a distribution we are referring to the location of the "center" of the distribution. The location of the center of a symmetric, mound shaped distribution is clearly the point of symmetry. There is some ambiguity in specifying the location of the center of an asymmetric, mound shaped distribution and we shall see that there are at least two standard ways to quantify location in this context. The second aspect of the distribution is the amount of variability or dispersion in the distribution. Roughly speaking, we would say that a distribution exhibits low variability if the observed values tend to be close together on the number line and exhibits high variability if the observed values tend to be more spread out in some sense. For example, the female height distribution histogram of Figure 2.4 is more peaked than the male height distribution histogram, which is somewhat flatter or more spread out. This indicates that, for this NHANES data, there is less variability among the heights of the females than there is among the heights of the males. The weight distribution histograms of Figure 2.5 suggest that the variability among the weights of the females is similar to the variability among the weights of the males. The third aspect is the shape of the distribution and in particular the degree of skewness in the distribution.

As a starting point consider the **minimum** (smallest observed value) and **maximum** (largest observed value) as statistics. We know that all of the data values lie between the minimum and the maximum, therefore, the minimum and the maximum provide a crude quantification of location and variability. In particular, we know that all of the values of the variable are restricted to the interval from the minimum to the maximum; however, the minimum and the maximum alone tell us nothing about how the data values are distributed within this interval. If the distribution is reasonably symmetric and mound shaped, then the **midrange**, defined as the average of the minimum and the maximum, may provide a suitable quantification of the location of the center of the distribution. The median and mean, which are defined below, are generally better measures of the center of a distribution.

The **range**, defined as the distance from the minimum to the maximum can be used to quantify the amount of variability in the distribution. Note that the range is the positive number obtained by subtracting the minimum from the maximum. When comparing two distributions the distribution with the larger range will generally have more variability than the distribution with the smaller range; however, the range is very sensitive to extreme observations so that one or a few unusually large or small values can lead to a very large range.

We will now consider an approach to the quantification of the shape, location, and variability of a distribution based on the division of the histogram of the distribution into sections of equal area. This is equivalent to dividing the data into groups, each containing the same number of values. We will first use a division of the histogram into halves. We will then use a division of the histogram into fourths.

The median is used to quantify the location of the center of the distribution. In terms of area, the **median** is the number (point on the number line) with the property that the area in the histogram to the left of the median is equal to the area to the right of the median. Here and in the sequel we will use a lower case n to denote the sample size, *i.e.*, n will denote the number of units in the sample. In terms of the n observations, the **median** is the number with the property that at least n/2 of the observed values are less than or equal to the median and at least n/2 of the observed values are greater than or equal to the median.

A simple procedure for finding the median, which is easily generalized to fractions other that 1/2, is outlined below.

Median computation procedure.

step 1. Arrange the data (observations) in increasing order from the smallest (obs. no. 1) to the largest (obs. no. n). Be sure to include all n values in this list, including repeats if there are any.

step 2. Compute the quantity n/2.

step 3a. If n/2 is not a whole number, round it up to the next largest integer. The observation at the location indicated by the rounded-up value in the ordered listing of the data is the median.

step 3b. If n/2 is a whole number, then we need to average two values to get the median. The two observations to be averaged are obs. no. n/2 and the next observation (obs. no.

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n/2 + 1) in the ordered listing of the data. Find these two observations and average them to get the median.

We can use the distance between the minimum and the median and the distance between the median and the maximum to quantify the amount of skewness in the distribution. The distance between the minimum and the median is the range of the lower (left) half of the distribution, and the distance between the median and the maximum is the range of the upper (right) half of the distribution. If the distribution is symmetric, then these two distances (median – minimum) and (maximum – median) will be equal. If the distribution is skewed, then we would expect to observe a larger range (indicating more variability) for the half of the distribution in the direction of the skewness. Thus if the distribution is skewed to the left, then we would expect (median – minimum) to be greater than (maximum – median). On the other hand, if the distribution is skewed to the right, then we would expect (maximum – median) to be greater than (median – minimum).

Example 2.2 Weed seeds (revisited). Recall that this example is concerned with the number of weed seeds found in n = 98 quarter-ounce batches of grass. Since 98/2 = 49, the median for this example is the average of observations 49 and 50. Referring to Table 2.2 we find that the minimum number of weed seeds is 0, the maximum is 7, and the median is 1, since observations 49 and 50 are each 1. The range for this distribution is 7 - 0 = 7. Notice that the range of the right half of this distribution (maximum - median) = 7 - 1 = 6 is much larger than the range of the left half (median - minimum) = 1 - 0 = 1 confirming our observation that this distribution is strongly skewed to the right.

A more detailed quantification of the shape and variability of a distribution can be obtained from a division of the distribution into fourths. In order to divide a distribution into fourths, we need to specify three numbers or points on the number line. These statistics are called **quartiles**, since they divide the distribution into quarters. In terms of area, the **first quartile**, denoted by Q_1 (read this as Q sub one), is the number (point on the number line) with the property that the area in the histogram to the left of Q_1 is equal to one fourth and the area to the right of Q_1 is equal to three fourths. The **second quartile**, denoted by Q_2 , is the median. The **third quartile**, denoted by Q_3 , is the number (point on the number line) with the property that the area in the histogram to the left of Q_3 is equal to three fourths and the area to the right of Q_3 is equal to one fourth. In terms of the *n* observations, Q_1 is the number with the property that at least n/4 of the observed values are less than or equal to Q_1 and at least 3n/4 of the observed values are greater than or equal to Q_1 . Similarly, Q_3 is the number with the property that at least 3n/4 of the observed values are less than or equal to Q_3 and at least n/4 of the observed values are greater than or equal to Q_3 .

The method for finding the median given above is readily modified for finding the first and third quartiles. For Q_1 , we simply replace n/2 by n/4 and replace the words 'the median' by Q_1 . To find Q_3 , use exactly the same method but count down from the largest value instead of counting up from the smallest value. Some calculators and computer programs use variations of the methods given above for finding Q_1 and Q_3 . These variations may give slightly different values for Q_1 and Q_3 .

Example 2.1 Weed seeds (revisited). Since 98/4 = 24.5, the quartiles Q_1 and Q_3 for this example are the observations located at position 25 counting up for Q_1 and counting down for Q_3 . Referring to Table 2.2 we find that $Q_1 = 0$ and $Q_3 = 2$. Notice that the range of the lower three-fourths of this distribution, Q_3 – minimum, is 2 while the range of the upper fourth, maximum – Q_3 is 5. This indicates that 75% (a large proportion) of the batches of grass have relatively few weed seeds, and the skewness in this distribution is due to the high amount of variability among the numbers of weed seeds in the 25% of the batches with between 2 and 7 weed seeds.

Previously we introduced the range as a measure of variability. An alternative measure of variability is provided by the interquartile range. The **interquartile range** (IQR) is the distance between the first quartile Q_1 and the third quartile Q_3 , *i.e.*, the interquartile range is the positive number obtained by subtracting Q_1 from Q_3 . Notice that the **interquartile range** is the range of the middle half of the distribution. The interquartile range is less sensitive to the presence of a few extreme observations in the data than is the range. For example, if there are one or two unusually large or unusually small values, then these values may have the effect of making the range much larger than it would be if these unusual values were not present. In such a situation, we might argue that the range is too large to be deemed an appropriate overall measure of the variability of the distribution. The interquartile range is not affected by a few unusual values, since it only depends on the middle half of the data. We could use the range of a larger part of the middle of the distribution, say the middle 75% or 90%, as a compromise between the range and the interquartile range.

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The five summary statistics: the minimum (min), the first quartile (Q_1) , the median (med), the third quartile (Q_3) , and the maximum (max), constitute the **five number summary** of the distribution. Each of these five statistics provides a quantification of a particular aspect of the distribution. They quantify where the distribution begins, where the first quarter of the distribution ends, and so on. Furthermore, the distances between these five statistics can be used to quantify the shape (skewness) of the distribution.

The four distances: $(Q_1 - \min)$, $(\text{med} - Q_1)$, $(Q_3 - \text{med})$, and $(\max - Q_3)$, are the ranges of the first, second, third, and fourth quarters of the distribution, respectively. These distances can be used to quantify the amount of variability in the corresponding parts of the distribution. Comparisons of appropriate pairs of these distances provide indications of certain aspects of the shape of the distribution. The relationship between $(\text{med} - Q_1)$ and $(Q_3 - \text{med})$ can be used to quantify the shape (skewness) of the middle half of the distribution. Since $(Q_1 - \min)$ and $(\max - Q_3)$ are the lengths of the tails (lower and upper fourths) of the distribution, the relationship between these numbers can be used to quantify skewness in the tails of the distribution.

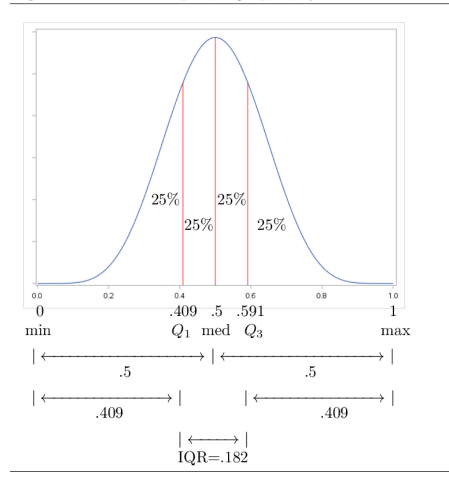
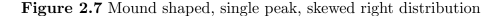
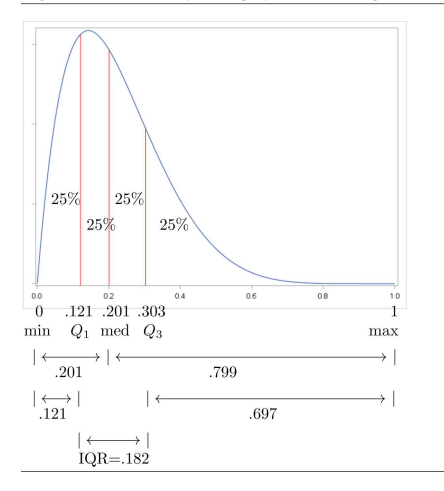


Figure 2.6. Mound shaped, single peak, symmetric distribution

The distribution of Figure 2.6 is mound shaped with a single peak (mode) at .5. This distribution is symmetric. Since this distribution is symmetric we see that the range of the left half of this distribution .5 is equal to the range of the right half; the range of the left tail .409 is equal to the range of the right tail; and, the median .5 is exactly half way between $Q_1 = .409$ and $Q_3 = .591$.

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The distribution of Figure 2.7 is mound shaped with a single peak (mode) around .15. This distribution is clearly skewed right. The fact that the range of the right half of this distribution .799 is about 4 times .201 the range of the left half shows extreme skewness to the right. This skewness is mostly due to the fact that the range of the right tail .697 is almost 6 times .121 the range of the left tail. Notice that the middle half of the distribution is reasonably symmetric since the median .201 is about half way between $Q_1 = .121$ and $Q_3 = .303$.

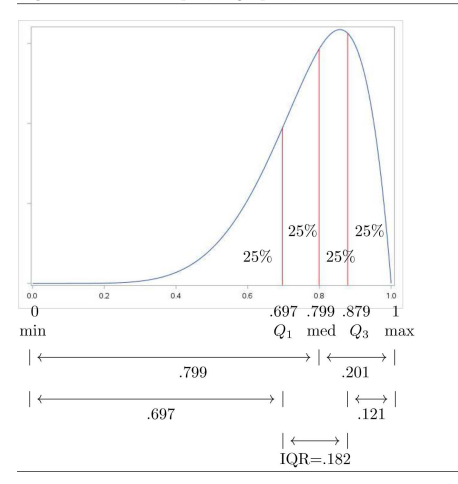


Figure 2.8 Mound shaped, single peak, skewed left distribution

The distribution of Figure 2.8 is mound shaped with a single peak (mode) around .85. This distribution is clearly skewed left. The fact that the range of the left half of this distribution .799 is about 4 times .201 the range of the right half shows extreme skewness to the left. This skewness is mostly due to the fact that the range of the left tail .697 is almost 6 times .121 the range of the right tail. Notice that the middle half of the distribution is reasonably symmetric since the median .799 is about half way between $Q_1 = .697$ and $Q_3 = .879$.

We can use the five number summary values to form a simple graphical representation of a distribution known as a **boxplot** or a box and whiskers plot. A boxplot provides a useful graphical impression of the shape of the distribution as well as its location and variability. Simple boxplots for unimodal mound shaped distributions similar to the distributions of Figures 2.6, 2.7, and 2.8 are provided in Figure 2.9.

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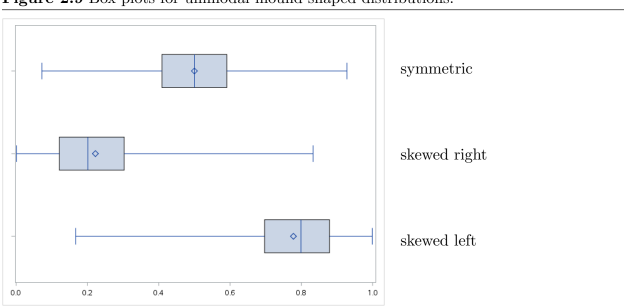


Figure 2.9 Box plots for unimodal mound shaped distributions.

Each boxplot has five vertical marks indicating the locations of the five number summary values. The box which extends from the first quartile to the third quartile and is divided into two parts by the median gives an impression of the distribution of the values in the middle half of the distribution. In particular, a glance at this box indicates whether the middle half of the distribution is skewed or symmetric and indicates the magnitude of the interquartile range (the length of the box). The line segments (whiskers) which extend from the ends of the box to the extreme values (the minimum and the maximum) give an impression of the distribution of the values in the tails of the distribution. The relative lengths of the whiskers indicate the contribution of the tails of the distribution to the symmetry or skewness of the distribution.

Example 1.1 NHANES (revisited). Boxplots for the NHANES height and weight distributions are given in Figures 2.10 and 2.11. In these boxplots the whiskers are modified to show extreme observations (indicated by o's in these plots) in the tails of the distribution. Summary information for the NHANES height and weight distributions is given in Tables 2.3 and 2.4.

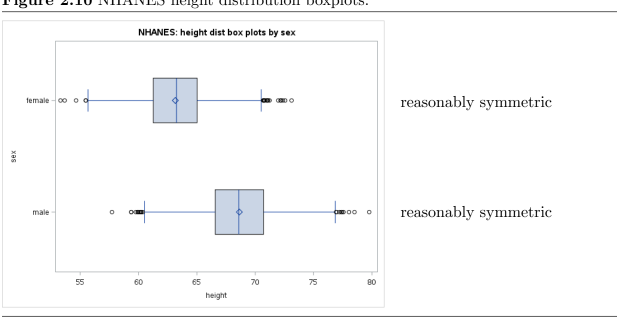


Figure 2.10 NHANES height distribution boxplots.

Table 2.3 NHANES height distribution summary information. (height in inches)

emales 2888 63.15 63.25 2.82 7.97 19.80 3.76	males 2642 68.63 68.58 3.06 9.39 22.05 4.13
63.1563.252.827.9719.80	68.63 68.58 3.06 9.39 22.05
63.25 2.82 7.97 19.80	$\begin{array}{c} 68.58 \\ 3.06 \\ 9.39 \\ 22.05 \end{array}$
63.25 2.82 7.97 19.80	$\begin{array}{c} 68.58 \\ 3.06 \\ 9.39 \\ 22.05 \end{array}$
$2.82 \\ 7.97 \\ 19.80$	$3.06 \\ 9.39 \\ 22.05$
$7.97 \\ 19.80$	$9.39 \\ 22.05$
$7.97 \\ 19.80$	$9.39 \\ 22.05$
19.80	22.05
3.76	/ 13
	4.10
53.31	57.72
61.26	66.57
63.25	68.58
65.02	70.71
73.11	79.76
7.95	8.85
1 00	2.01
1.99	2.13
$1.99 \\ 1.77$	
	1.99

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As noted earlier, the height distribution histograms for the adult males and the adult females of Figure 2.4 are both unimodal and reasonably symmetric. The boxplots of Figure 2.10 also indicate reasonable symmetry. We will now use the distances from Table 2.3 to quantify this claim of reasonable symmetry. For the adult males we see that the range of the left tail 7.95 is almost the same as 8.09 the range of the right tail. Looking at the middle half of the male height distribution we find that the range of the left side $med - Q_1 = 1.99$ is only slightly larger than the range of the right side $Q_3 - med = 1.77$. Similarly, for the adult females the range of the left tail 8.85 is almost the same as 9.05 the range of the right tail; and, $med - Q_1 = 2.01$ is only slightly smaller than $Q_3 - med = 2.13$.

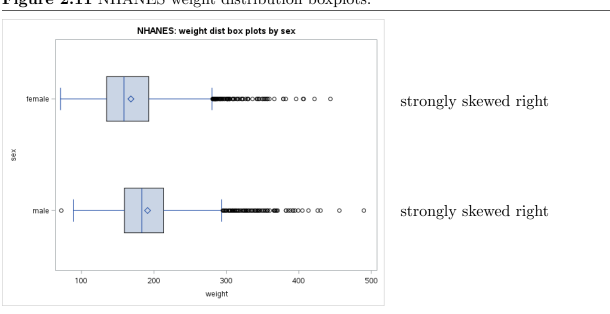


Figure 2.11 NHANES weight distribution boxplots.

Turning to the weight distributions, recall that the weight distribution histograms for the adult males and the adult females of Figure 2.5 are both unimodal and strongly skewed to the right. The boxplots of Figure 2.11 also indicate strong skewness to the right. From each of these boxplots it is clear that in both cases the skewness is due to the extreme variability in the right tail of the distribution, that is, due to high variability among the weights of the heaviest 25% of the group. Note also that in each case the middle half of the distribution is reasonably symmetric. The distances in Table 2.4 readily support these observations. For the male weights $\max -Q_3 = 250.58$ is much larger than $Q_1 - \min = 63.58$, while, relatively speaking, med $-Q_1 = 23.98$ is only slightly smaller than $Q_3 - \text{med} = 34.32$. Similarly, for the female weights max $-Q_3 = 276.32$ is much larger than $Q_1 - \min = 87.12$, while, relatively speaking, med $-Q_1 = 23.98$ is only slightly smaller than $Q_3 - \text{med} = 30.14$.

	gro	oup
statistic	females	males
n	2888	2645
location		
mean	168.23	191.21
median	158.62	183.26
variability		
std deviation	47.70	46.80
variance	2275	2190
range	372.46	417.56
IQR	58.30	54.12
5 number summary		
min	71.06	72.16
Q_1	134.64	159.28
median	158.62	183.26
Q_3	192.94	213.40
max	443.52	489.72
distances		
Q_1 -min	63.58	87.12
med - Q_1	23.98	23.98
Q_3 -med	34.32	30.14
$\max -Q_3$	250.58	276.32

Table 2.4 NHANES	weight	distribution sum	mary inform	nation.	(weight in	pounds)
------------------	--------	------------------	-------------	---------	------------	---------

2.6 Quantiles and percentiles in general

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We will now provide an extension of the method we used to compute the median and quartiles to allow an arbitrary fraction. Given a proportion p (a fraction between zero and one), the *pth* **quantile** ($p \times 100th$ **percentile**) of the distribution of X is the value Q_p with the property that if we choose a value of X at random, then X will be less than Q_p with probability p and X will be greater than Q_p with probability 1 - p, *i.e.*, $p \times 100\%$ of the time X will be less than Q_p and $(1 - p) \times 100\%$ of the time X will be greater than Q_p . In terms of the histogram of the distribution of X this means that the area in the histogram to the left of Q_p is p and the area to the right of Q_p is 1 - p. Note that the first quartile is the 25th percentile, the median is the 50th percentile, and the third quartile is the 75th percentile.

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In terms of a sample of n values, the pth quantile is the number Q_p with the property that at least pn of the sample values are less than or equal to Q_p and at least (1 - p)nof the sample values are greater than or equal to Q_p . In order to compute a quantile we first need to sort (order) the data. Let x_1, x_2, \ldots, x_n denote the (unordered) data. Let $x_{(1)}, x_{(2)}, \ldots, x_{(n)}$ denote the ordered values with $x_{(1)} \leq x_{(2)} \leq \cdots \leq x_{(n)}$. These ordered values are know as **order statistics**.

Quantile computation procedure. For 0 , the*pth* $quantile <math>Q_p$ of the *n* sample values x_1, \ldots, x_n of X is computed as follows. Let $x_{(1)} \leq x_{(2)} \leq \cdots \leq x_{(n)}$ denote the sample values ordered from smallest to largest. There are two cases case 1: If there is an integer k such that pn = k, then $Q_p = \frac{x_{(k)} + x_{(k+1)}}{2}$ case 2: If there is an integer k such that k < pn < k + 1, then $Q_p = x_{(k+1)}$.

2.7 The mean and standard deviation

The approach that we have been using to form summary statistics is to select a single representative value from the observed values of the variable (or the average of two adjacent observed values) to quantify a particular aspect of the distribution. We have also considered statistics that are distances between two such representative values.

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An alternative approach to forming a summary statistic is to combine all of the observed values to get a suitable statistic. The first statistic of this type that we consider is the mean. The **mean**, which is the simple arithmetic average of the n data values, is used to quantify the location of the center of the distribution. You could compute the mean by adding all n data values together and dividing this sum by n; however, it is better to use a calculator or a computer. The sample mean is often denoted by the symbol \overline{X} (read this as X bar).

Recall that the median is the number (point on the number line) with the property that the area in the histogram to the left of the median is equal to the area to the right of the median. The mean is the number (point on the number line) where the histogram would balance. To understand what we mean by the balance point, imagine the histogram as being cut out of a piece of cardboard. The mean is located at the point along the number line side of this cutout where the histogram cutout would balance. These geometric characterizations of the mean and the median imply that when the distribution is symmetric the mean will be equal to the median. Furthermore, if the distribution is skewed to the right, then the mean (the balance point) will be larger than the median (to the right of the median).

Similarly, if the distribution is skewed to the left, then the mean (the balance point) will be smaller than the median (to the left of the median).

The primary use of the mean, like the median, is to quantify the location of the center of a distribution and to compare the locations (centers) of two distributions. Since both the mean and the median can be used to quantify the location of the center of a distribution, it seems reasonable to ask which is more appropriate. If the distribution is approximately symmetric, then the mean and the median will be approximately equal. On the other hand, if the distribution is not symmetric, then the median is likely to provide a better indication of the center of the distribution. For example, if the distribution is strongly skewed to the right, then the mean may be much larger than the median and the mean may not be a good indication of the center of the distribution. For a specific application it is a good idea to mark the locations of the mean and the mean and the median on a histogram of the distribution and consider which seems more reasonable as an indicator of the center of the distribution.

The two measures of variability we discussed earlier, the range and the interquartile range, are distances between two representative values, the minimum and maximum for the range and the first and third quartiles for the interquartile range. We will now discuss a more complex measure of variability which is based on the distances between each of the observations and a single representative value. If the mean \overline{X} is deemed suitable as a measure of the center of the distribution of X, then the deviations $(X - \overline{X})$ of the observed values of X from their mean \overline{X} contain information about the amount of variability in the distribution. If there is little variability (the observed values of X are close together and they are close to the mean \overline{X}), then the deviations $(X - \overline{X})$ will tend to be small in magnitude (absolute value). On the other hand, if there is a lot of variability (at least some of the observed values of X are far apart and they are not all close to the mean \overline{X}), then the deviations $(X - \overline{X})$ will tend to be large in magnitude. It is this observation which suggests that a summary statistic based on the distances between each of the observed values of the variable and their mean can be used to measure the variability in the distribution. These deviations from the mean are illustrated, for a small sample of values, in Figure 2.12.

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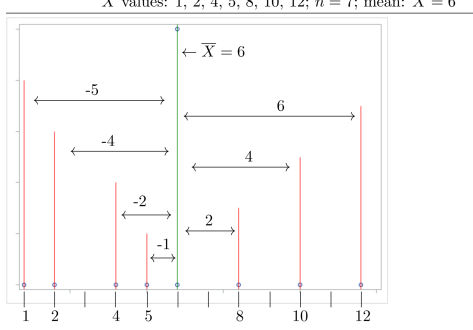


Figure 2.12 An example showing deviations from the mean. X values: 1, 2, 4, 5, 8, 10, 12; n = 7; mean: $\overline{X} = 6$

The most commonly used measure of variability based on these deviation from the mean is the **standard deviation**. The **standard deviation** is the square root of the "average" of the squared deviations of the observed values of the variable from their mean. The formula for the standard deviation given below is not intended for computation purposes; you should use a calculator or a computer to compute the standard deviation. The standard notation for the **sample standard deviation** of the distribution of the variable X is S_X (read this as S sub X). The defining formula is

$$S_X = \sqrt{\frac{\Sigma(X - \overline{X})^2}{n - 1}}$$

In this formula the capital Greek letter sigma, Σ , represents the statement "the sum of", and $(X-\overline{X})^2$ denotes the square of the distance from the observed value X to the mean \overline{X} . Therefore, as mentioned above, the expression under the square root sign in the formula is the "average" of the squared deviations of the observed values of the variable from their mean. The reason for the square root is so that the standard deviation of X and the variable X are in the same units of measurement. The quantity

$$S_X^2 = \frac{\Sigma (X - X)^2}{n - 1}$$

is the **sample variance** of the distribution of the variable X. The calculations corresponding to these formulae are demonstrated below for the sample used in Figure 2.12.

Example 2.3 Computation of the standard deviation. Consider the sample of seven (n = 7) X values: 1, 2, 4, 5, 8, 10, 12. The sample mean is

$$\overline{X} = \frac{1+2+4+5+8+10+12}{7} = \frac{42}{7} = 6.$$

The sample variance is

$$\begin{split} S_X^2 &= \frac{(1-6)^2 + (2-6)^2 + (4-6)^2 + (5-6)^2 + (8-6)^2 + (10-6)^2 + (12-6)^2}{7-1} \\ &= \frac{(-5)^2 + (-4)^2 + (-2)^2 + (-1)^2 + (2)^2 + (4)^2 + (6)^2}{6} \\ &= \frac{25+16+4+1+4+16+36}{6} = \frac{102}{6} = 17. \end{split}$$

The sample standard deviation is

$$S_X = \sqrt{17}.$$

The standard deviation is positive, unless there is no variability at all in the data. That is, unless all of the observations are exactly the same, the standard deviation is a positive number. The standard deviation is a very widely used measure of variability. Unfortunately, the standard deviation does not have a simple, direct interpretation. The important thing to remember is that larger values of the standard deviation indicate that there is more variability in the data.

There are quotation marks around the word average in the definition of the sample standard deviation because we divided by n - 1 even though there are n squared deviations in the average. When the standard deviation (variance) is computed for the population this divisor is changed to n and a lower case Greek sigma is used instead of an S. That is, the **population standard deviation** is defined as

$$\sigma_X = \sqrt{\frac{\Sigma(X - \overline{X})^2}{n}}$$

and the **population variance** is σ_X^2 .

Example 1.1 NHANES (revisited). We noted earlier that the shapes of the female and male height distributions are very similar and that the shapes of the female and male weight

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distributions are also very similar. We will now use summary statistics from Tables 2.3 and 2.4 to compare and contrast the locations and the variability in these distributions.

We will first look at the height distributions. As you would expect the males tend to be taller than the females. On average the males are about 5.5 inches taller than the females (male mean height of 68.63 inches versus female mean height of 63.15 inches). Since the height distributions are reasonably symmetric the median heights are very similar to the mean heights. In terms of variability, there is slightly more variability among the height standard deviation of 3.06 inches verses female height standard deviation of 2.82 inches; male height range of 22.05 inches verses female height range of 19.80 inches; and, male height interquartile range of 4.13 inches verses female height interquartile range of 3.76 inches.

Now we will look at the weight distributions. Again, as you would expect, the males tend to be heavier than the females. On average the males are about 23 pounds heavier than the females (male mean weight of 191.21 pounds versus female mean weight of 168.23 pounds). Since the both weight distributions are strongly skewed right each median weight is smaller than the corresponding mean weight; but, as it turns out, the difference between the median weights is similar to the difference between the mean weights (the median weight difference is about 25 pounds). The weight distributions are quite similar in terms of variability.

2.8 A measure of relative position

Percentiles can be used to quantify the location of a particular value of X relative to a group. Another widely used measure of the relative position of a value within a group is its Z-score. The Z-score of X quantifies the location of X relative to the mean \overline{X} of the sample in terms of the standard deviation S_X of the sample. Since the Z-score is based on \overline{X} and S_X , the Z-score is only appropriate when \overline{X} and S_X are appropriate measures of the center and variability in the sample, respectively. We will develop the Z-score in two stages.

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First, we need a measure of the location of X relative to the center of the distribution as determined by the mean \overline{X} . The deviation, $X - \overline{X}$, of X from the mean \overline{X} is such a measure. The deviation $X - \overline{X}$ is the signed distance from the particular value X to the mean \overline{X} . If $X - \overline{X}$ is negative, then X is below (smaller than) the mean. If $X - \overline{X}$ is positive, then X is above (larger than) the mean. In summary, the sign of the deviation $X - \overline{X}$ indicates the location of X relative to the mean \overline{X} ; and the magnitude of the deviation $|X - \overline{X}|$ is the distance from X to the mean \overline{X} , measured in the units of measurement used for the observation X.

Second, we want a measure of the location of X relative to the mean \overline{X} which takes the amount of variability in the data into account. We will obtain such a measure by using the standard deviation S_X of the sample to standardize the deviation $X - \overline{X}$. Given a particular value X, the sample mean \overline{X} , and the sample standard deviation S_X , the **Z**-score corresponding to X is

$$Z = \frac{X - \overline{X}}{S_X}.$$

The sign of the Z-score indicates the location of X relative to the mean \overline{X} and the magnitude of the Z-score is the distance from X to the mean \overline{X} in terms of standard deviation units. For example, if Z = 2, then X is two standard deviation units above the mean $(X = \overline{X} + 2S_X)$, and, if Z = -2, then X is two standard deviation units below the mean $(X = \overline{X} - 2S_X)$.

Interpretation of a Z-score requires some knowledge of the connection between Z-scores and percentiles. The 68%-95%-99.7% rule given below allows us to associate a percentage with a Z-score. This rule works best for distributions that are unimodal (single peaked), mound shaped, and symmetric. A formal statement of the rule follows.

The 68%-95%-99.7% rule. For a distribution that is unimodal (has a single peak), mound shaped, and reasonably symmetric:

i) Approximately 68% of the observed values will be within one standard deviation unit of the mean. That is, approximately 68% of the observed values will have a Z-score that is between -1 and 1.

ii) Approximately 95% of the observed values will be within two standard deviation units of the mean. That is, approximately 95% of the observed values will have a Z-score that is between -2 and 2.

iii) Approximately 99.7% of the observed values will be within three standard deviation units of the mean. That is, approximately 99.7% of the observed values will have a Z-score that is between -3 and 3. Notice that this indicates that almost all of the observed values will be within three standard deviations of the mean.

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When it is applicable, the 68% - 95% - 99.7% rule, can be used to determine the relative position of a particular value of a variable based on the corresponding Z-score. Notice that this rule indicates that a fairly large proportion (68%) of the sample will lie within one standard deviation of the mean; a very large proportion (95%) of the sample will lie within two standard deviations of the mean; and, almost all (99.7%) of the sample will lie within three standard deviations of the mean.

An aside – Chebyshev's rule

Another connection between Z-scores and percentages is provided by Chebyshev's rule. Chebyshev's rule is a mathematical fact that is true for any distribution. Unfortunately, the universal applicability of Chebyshev's rule forces its conclusions to be of more theoretical than practical interest. That is, the conclusions of Chebyshev's rule are valid for any distribution; but, they are often so imprecise that they are of limited practical use.

Chebyshev's rule. For any distribution:

i) At least 75% of the observed values will be within two standard deviation units of the mean. That is, at least 75% of the observed values will have a Z-score that is between -2 and 2.

ii) At least 89% of the observed values will be within three standard deviation units of the mean. That is, at least 89% of the observed values will have a Z-score that is between -3 and 3.

iii) In general, given a number k > 1, at least $[1 - (1/k^2)]100\%$ of the observed values will be within k standard deviation units of the mean, i.e., at least this percentage of the observed values will have a Z-score that is between -k and k.

2.9 Problems

The problem sets and solutions can be found on the book web page at https://www.ucs.louisiana.edu/~jcb0773/.

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3 Probability

3.1 The setting

Probability theory is used to model the behavior of random experiments. In this context a **random experiment** is any process of observation or experimentation for which the particular outcome is not known with certainty in advance of performance of the experiment. For example we might consider the experiment of tossing a coin or a die, tossing a pair of dice, dealing a hand of cards from a deck of cards, drawing balls from a box of balls, selecting a person from a population and measuring the person's height, age, or weight, or testing a machine to determine if it works properly.

Our goal is to formulate a theory which can be used to specify a formal model for the randomness in the outcomes of the experiment by assigning probabilities to events associated with the experiment. An **event** is a description of the outcome of the experiment. It is useful to distinguish between simple events and compound events. A **simple event** is an event which cannot be decomposed into simpler events. We will refer to simple events as **elementary outcomes**. A **compound event** is an event which can be decomposed into two or more events. For example, if we toss a die once, then the elementary outcomes, "observe a 1", "observe a 2", *etc.*, can be represented by the integers 1, 2, 3, 4, 5, and 6. The compound event "observe an even number" is the collection (set) $\{2, 4, 6\}$ of the three elementary outcomes corresponding to even numbers.

The first step in forming a probability model for a particular experiment is the specification of a **sample space** containing all the possible elementary outcomes of the experiment. Relevant events can then be viewed as subsets of the sample space. The elementary outcomes (simple events) are the singleton sets (sets containing a single elementary outcome) and compound events are sets containing two or more elementary outcomes.

3.2 Some illustrative examples

Example 3.1 Tossing a die. As noted above, we can represent the 6 elementary outcomes of a single toss of a die by the integers 1, 2, 3, 4, 5, and 6. Now suppose that a die is tossed twice. The 36 elementary outcomes for this experiment can be represented by ordered pairs of the integers $1, \ldots, 6$. For example the ordered pair (1,3) indicates that the first toss yielded a 1 and the second yielded a 3. The sample space is the set containing the 36 elementary outcomes (ordered pairs) shown in Table 3.1.

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	second toss												
irst toss	1	2	3	4	5	6							
1	(1,1)	(1,2)	(1,3)	(1,4)	(1,5)	(1,6)							
2	(2,1)	(2,2)	(2,3)	(2,4)	(2,5)	(2,6)							
3	(3,1)	(3,2)	(3,3)	(3,4)	(3,5)	(3,6)							
4	(4,1)	(4,2)	(4,3)	(4,4)	(4,5)	(4,6)							
5	(5,1)	(5,2)	(5,3)	(5,4)	(5,5)	(5,6)							
6	(6,1)	(6,2)	(6,3)	(6,4)	(6,5)	(6,6)							

Table 3.1 Elementary outcomes (ordered pairs) when a die is tossed twice.

Two events of potential interest are:

1. The event that the sum of the two numbers observed is 7. The ordered pairs on the upper right to lower left diagonal of the table are favorable for this event. Thus the event that the sum is 7 is represented by the set

$$\{(1,6), (2,5), (3,4), (4,3), (5,2), (6,1)\}.$$

Read this as: the set containing the ordered pairs (1, 6), (2, 5), (3, 4), (4, 3), (5, 2), and (6, 1).

2. The event that the number on the first toss is even. The ordered pairs in the second, fourth, and sixth rows of the table are favorable to this event. Thus the event that the number on the first toss is even is represented by the set

$$\left\{ \begin{array}{l} (2,1), (2,2), (2,3), (2,4), (2,5), (2,6), \\ (4,1), (4,2), (4,3), (4,4), (4,5), (4,6), \\ (6,1), (6,2), (6,3), (6,4), (6,5), (6,6) \end{array} \right\}.$$

Example 3.2 Tossing a coin. If we toss a coin once, then, letting H denote "heads" and T denote "tails", the sample space is the set containing the 2 elementary outcomes $\{H, T\}$. Letting H and T denote the values of a dichotomy, such as the sex of a person (male or female), the quality of an item (acceptable or unacceptable), or the outcome of a medical procedure (successful or not), we can use the H and T notation of this and the three toss example below to represent a wide variety of experiments.

If we toss a coin three times, then, letting H denote "heads" and T denote "tails", we can use an ordered triple (with the order indicating the outcomes of the first, second, and

third tosses) to represent an elementary outcome. The sample space is the set containing the 8 elementary outcomes (ordered triples):

$\{HHH, HHT, HTH, HTT, THH, THT, TTH, TTT\}.$

Two events of potential interest are:

1. The event that there are exactly two heads. The ordered triples with two H's and one T are favorable to this event. Thus the event is represented by the set

$\{HHT, HTH, THH\}.$

2. The event that there are at least two heads. The ordered triples with two H's and one T or three H's are favorable to this event. Thus the event is represented by the set

```
\{HHH, HHT, HTH, THH\}.
```

An aside – playing cards, bridge, and poker

A standard deck of playing cards, as shown in Figure 3.1, contains 52 cards. The cards appear in four suits, hearts \heartsuit , diamonds \diamondsuit , clubs \clubsuit , and spades \bigstar , of these the heart and diamond cards are colored red and the club and spade cards are colored black. The 13 cards of each suit have face values (ranks) of 2, 3, ..., 10, jack, queen, king, and ace. An ace is often treated as having a face value (rank) of one. Cards with the same face value are said to be of the same kind. The jacks, queens, and kings, which typically (as with the deck in Figure 3.1) have "faces" on them are called face cards.

Bridge is a four-player partnership game. The four players in bridge are known as North, East, South, and West. North and South form one team and East and West form the second team. All 52 cards are dealt with each player receiving 13 cards. Thus, for our purposes, bridge means dividing the 52 cards into four hands (sets) of 13 cards and a bridge hand is a collection of 13 cards. There are many variations of poker. For our purposes, a poker hand is a collection of five cards.

A ‡	*	*	2	* *	÷2	3 *	* * *	A.‡	4 * ◆	* **	5 *	*	+ + * *	6 * *	*	74	*	* * *	8 •	****	9	* * *.* ***	* 6		*	, t
A •	٨	*	2	♠ ♥	Ż	3 ♠	♠ ♠ ♥	Ψ	4 ♠♠ ♥	♠ ♥₩	5 ♠	♠ ₩	ب ۹ 9¢	6	♠ ♠ ♥		* *	♠ ♠ ♥Ľ	8 ♠ ♠ ♥		9	* * *** * *				
€	٠	\$	2	•	2	3	*	¢ 2	4 ♥	♥ ♠♣	5 ♥	*	• • •	€ ♥ ♥	•			¥ ◆ į́	8 • •	8	9		6			
A ◆	٠	•	2	* •	ŧ	3 ◆	* * *	¢ 2	4 ★ ◆	◆ ◆;	5 ◆	•	• • • •	6 ↓ ◆ ↓	• • •			• • <u>;</u>	8 • •	8	9 •		•			, k

Figure 3.1 A standard deck of 52 playing cards

Example 3.3 Bridge or poker. If we assign the numbers from 1 to 52 as labels for the 52 cards in the deck (for example we might label the cards according to their positions in Figure 3.1), then we can use an ordered arrangement of the numbers from 1 to 52 to represent a particular deal of cards for a bridge game with the first 13 numbers corresponding to North, the next 13 to East, and so on. If we are only concerned with characteristics of one hand we could use a collection of 13 different numbers selected from the labels 1 to 52. Similarly, we could use a collection (set) of 5 different numbers to represent a poker hand.

Consider the event that a poker hand contains exactly two aces. The sets of five cards containing two aces and three non-aces are favorable for this event. There are too many such hands to list here. In case you wondered, using notation defined in Section 5, there are $\binom{4}{2}\binom{48}{3} = 103,776$ hands containing exactly two aces.

Example 3.4 Tossing a coin or die repeatedly. Suppose that a coin is tossed repeatedly until a head occurs. In this application we can represent an elementary outcome by a sequence terminating with an H and with enough T's to indicate the outcome. For example; Hindicates that we got heads on the first toss, TH indicates that we first got heads on the second toss, TTH indicates that we first got heads on the third toss, and so on. Thus the sample space is the countably infinite set

$$\{H, TH, TTH, TTTH, \ldots\}$$

As noted earlier, we can use the H and T notation of this example to represent a wide variety of experiments. Three events of potential interest are: 1. The event that exactly three tosses are required to get a head. Since TTH is the only outcome favorable for this event, it is represented by the set

$\{TTH\}.$

2. The event that at most three tosses are required to get a head. The three outcomes favorable for this event, heads first, second, or third, are the elements of the set

$$\{H, TH, TTH\}.$$

3. The event that more than three tosses are required to get a head. The representation of this event is provided by the infinite set

$\{TTTH, TTTTH, TTTTTH, \ldots\}$

containing all sequences which consist of a sequence of three or more T's followed by a single H.

If a die is tossed repeatedly until a 1 (an ace) occurs, then this same representation can be used with H indicating that a 1 occurs and T indicating that a 2, 3, 4, 5, or 6 occurs.

Similarly, if parts selected from a production line are tested sequentially until a defective part is found, then this same representation can be used with H indicating that the part is defective and T indicating that the part is acceptable.

Example 3.5 Sampling for a numerical value. Suppose that we are interested in the distribution of weights among individuals in a particular population. Further suppose that we choose one individual from this population and determine the weight of this individual. We can use a positive real number x to represent an elementary outcome (the weight of the individual). If we know the weights of all of the individuals in the population we can use these weights as the sample space. Since it is unlikely that we would know the weights of all the individuals, it is more convenient to use an interval of values on the number line as our sample space. If we know that no one in the population weights more than 300 pounds, then we can use the interval from zero to 300 ((0, 300] in interval notation) as our sample space. If we do not want to specify a maximal weight, we can use the positive part of the number line ((0, ∞) in interval notation) to represent the sample space. Events can be represented by appropriate intervals. For example, the event "the individual weights

between 100 and 150 pounds" corresponds to the interval [100, 150] and the event "the individual weighs at least 100 pounds" corresponds to the interval $[100, \infty)$ (or [100, 300) with the sample space (0, 300)).

3.3 Sample spaces and events

The examples given above illustrate the process of defining an elementary outcome, a sample space, and events for several specific experiments. Recall that for our purposes an experiment is a process of observation or experimentation which results in one of several possible elementary outcomes. We will now begin a more formal treatment of these aspects of an experiment. We will assume that all of the possible elementary outcomes of the experiment are known in advance but the actual outcome of the experiment will not be known with certainty until after the experiment is performed.

The sample space Ω of a particular experiment is the collection of all possible elementary outcomes for the experiment. Recall that an elementary outcome or simple event is an event which cannot be decomposed into simpler events. We will assume that these elementary outcomes are mutually exclusive so that two distinct elementary outcomes cannot occur at the same time. A generic elementary outcome will be denoted ω . In set terminology the objects which form a set are called elements, thus, we may refer to an elementary outcome as an element of the sample space or of an event. (Ω and ω are the upper and lower case versions of the Greek letter omega.)

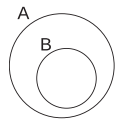
An event A is a collection of elementary outcomes, *i.e.*, a subset of Ω . If the experiment is conducted and the elementary outcome ω occurs, then if ω is an element of A (in symbols, $\omega \in A$) we say that event A has occurred. On the other hand, if ω is not an element of A (in symbols, $\omega \notin A$) we say that A has not occurred. For example, in the die tossing example, if we observe $\omega = 2$, then $2 \in \{2, 4, 6\}$ and the event observe an even number occurred. But, $2 \notin \{1, 3, 5\}$ so the event observe an odd number did not occur.

For convenience we define the **null event** (empty set), \emptyset , as the event with no elements. Viewed as an event, the sample space Ω is the **sure event**, since it contains all possible outcomes and therefore must occur.

Given two events A and B we write $B \subset A$ (B is contained in A or B is a subset of A) when every element of B is also an element of A. More formally, $B \subset A$ means that if $\omega \in B$, then $\omega \in A$. This is illustrated in Figure 3.2. Notice that if $B \subset A$, then the occurrence of B implies the occurrence of A. In particular, if B is a proper subset of A, that is, if A contains some elements that are not in B, then the event B is a "special case" of the event A. Also note that for every event $A, A \subset \Omega$ (A is a subset of the sample space Ω), $A \subset A$ (A is a subset of itself), and $\emptyset \subset A$ (the empty set is a subset of A).

Figure 3.2 An event and a subevent.

In this Venn diagram event B is a subevent (subset) of event A.



Example 3.6a Tossing a die once. For one toss of a die we have $\Omega = \{1, 2, 3, 4, 5, 6\}$. Consider the following events:

- A: "the number observed is 4 or less" $A = \{1, 2, 3, 4\}$
- *B*: "the number observed is 1, 3, or 4" $B = \{1, 3, 4\}$
- C: "the number observed is 5 or 6" $C = \{5, 6\}$

D: "the number observed is 3 or more" $D = \{3, 4, 5, 6\}$.

For these events, $B \subset A$, since observing a 1, 3, or 4 implies that we observed a number which is 4 or less, and $C \subset D$ since observing 5 or 6 implies that we we observed a number which is 3 or larger.

Example 3.6b Tossing a coin three times. For three tosses of a coin we have

 $\Omega = \{HHH, HHT, HTH, HTT, THH, THT, TTH, TTT\}$. Consider the following events:

- A: "the first toss yields a head" $A = \{HHH, HHT, HTH, HTT\}$
- B: "the first two tosses yield heads" $B = \{HHH, HHT\}$
- C: "there are exactly two heads" $C = \{HHT, HTH, THH\}$

D: "there are at least two heads" $D = \{HHH, HHT, HTH, THH\}$.

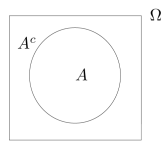
For these events, $B \subset A$, since observing heads on the first two tosses is a special case of observing heads on the first toss, $B \subset D$, since observing heads on the first two tosses is a special case observing at least two heads, and $C \subset D$ since observing exactly two heads is a special case of observing at least two heads.

52 3.3 Sample spaces and events

Given an event A the complementary event A^c (the **complement** of A) is the event containing all elements of Ω that do not belong to A. This is illustrated in Figure 3.3. In symbols, $A^c = \{\omega \in \Omega : \omega \notin A\}$ Read this as the set of ω in Ω such that ω is not in A. In words, A^c (A complement) is the event that A does not occur.

Figure 3.3 An event A and its complement A^c .

In this Venn diagram, the rectangle represents the sample space Ω , the interior of the circle the event A, and the region outside the circle A^c , the complement of A.



Example 3.6a Tossing a die once (continued). The complements of the events $A = \{1, 2, 3, 4\}, B = \{1, 3, 4\}, C = \{5, 6\}, \text{ and } D = \{3, 4, 5, 6\}$ are: $A^c = C = \{5, 6\}, B^c = \{2, 5, 6\}, C^c = A$, and $D^c = \{1, 2\}$

Example 3.6b Tossing a coin three times (continued). The complements of the events $A = \{HHH, HHT, HTH, HTT\}, B = \{HHH, HHT\}, C = \{HHT, HTH, THH\}, and D = \{HHH, HHT, HTH, THH\}$ are:

 $A^{c} = \{THH, THT, TTH, TTT\}$ "the first toss yields tails"

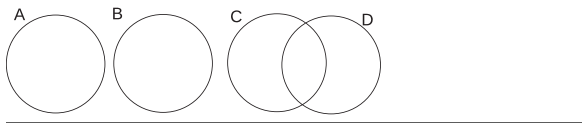
 $B^c = \{HTH, HTT, THH, THT, TTH, TTT\}$ "at least one of the first two tosses yields tails"

 $C^{c} = \{HHH, HTT, THT, TTH, TTT\}$ "there are zero, one, or three heads" $D^{c} = \{HTT, THT, TTH, TTT\}$ "there is at most one head".

Note that, for any event A, events A and A^c are mutually exclusive in the sense that they cannot both occur at the same time. Furthermore, since they are complementary, exactly one of A and A^c must occur. Note also that $\Omega^c = \emptyset$ (the complement of everything is nothing) and $\emptyset^c = \Omega$ (the complement of nothing is everything).

In general, two events are said to be **mutually exclusive** or **disjoint** if they do not have any elements in common. In other words, two events are mutually exclusive (disjoint) if they cannot occur at the same time. This is illustrated in Figure 3.4.

Figure 3.4 Mutually exclusive (disjoint) events. In this Venn diagram events A and B are mutually exclusive or disjoint while events C and D are not mutually exclusive.

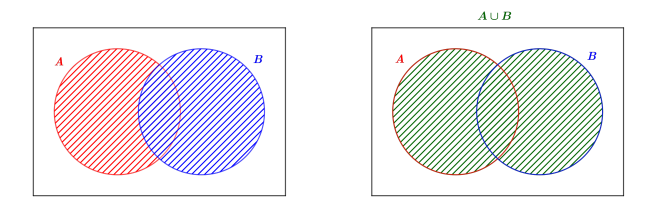


Example 3.6a Tossing a die once (continued). There are two pairing of the events $A = \{1, 2, 3, 4\}, B = \{1, 3, 4\}, C = \{5, 6\}, \text{ and } D = \{3, 4, 5, 6\}$ which give mutually exclusive events. A and C are mutually exclusive and B and C are mutually exclusive.

Example 3.6b Tossing a coin three times (continued). Each of the events

 $A = \{HHH, HHT, HTH, HTT\}, B = \{HHH, HHT\}, C = \{HHT, HTH, THH\}, and D = \{HHH, HHT, HTH, THH\}$ contains HHT; thus, no pairing of these events gives mutually exclusive events. On the other hand, the event $E = \{TTT, TTH, THT, HTT\}$ (observe at least two tails) shares no elements with B, C, or D; thus, E and B are mutually exclusive, E and C are mutually exclusive, and E and D are mutually exclusive.

Figure 3.5 The union of A and B.



The **union** of events A and B, denoted $A \cup B$, is the collection of elementary outcomes which belong to A or B. The *or* in this statement is the *logical or* meaning one or the other or both. That is, when we say A or B we mean A alone or B alone or both A and B. This is illustrated in Figure 3.5.

In symbols, $A \cup B = \{\omega \in \Omega : \omega \in A \text{ or } \omega \in B\}$

In words, $A \cup B$ is the event "A or B" in the sense that A or B or both A and B occur.

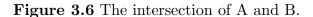
Example 3.6a Tossing a die once (continued). Recall that $A = \{1, 2, 3, 4\}$, $B = \{1, 3, 4\}, C = \{5, 6\}, \text{ and } D = \{3, 4, 5, 6\}$ The unions of these events are: $A \cup B = \{1, 2, 3, 4\} \cup \{1, 3, 4\} = \{1, 2, 3, 4\} = A$ (B is a subset of A) $A \cup C = \{1, 2, 3, 4\} \cup \{5, 6\} = \{1, 2, 3, 4, 5, 6\}$ $A \cup D = \{1, 2, 3, 4\} \cup \{3, 4, 5, 6\} = \{1, 2, 3, 4, 5, 6\}$ $B \cup C = \{1, 3, 4\} \cup \{5, 6\} = \{1, 3, 4, 5, 6\}$ $B \cup D = \{1, 3, 4\} \cup \{3, 4, 5, 6\} = \{1, 3, 4, 5, 6\}$ $C \cup D = \{5, 6\} \cup \{3, 4, 5, 6\} = \{3, 4, 5, 6\} = D$ (C is a subset of D)

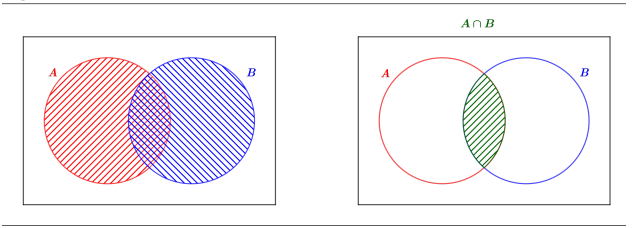
Example 3.6b Tossing a coin three times (continued). Recall that $A = \{HHH, HHT, HTH, HTT\}, B = \{HHH, HHT\}, C = \{HHT, HTH, THH\},$ $D = \{HHH, HHT, HTH, HTH\}$ The unions of these events are: $A \cup B = \{HHH, HHT, HTH, HTT\} \cup \{HHH, HHT\} = A (B \text{ is a subset of } A)$ $A \cup C = \{HHH, HHT, HTH, HTT\} \cup \{HHT, HTH, THH\} = A (C \text{ is a subset of } A)$ $A \cup D = \{HHH, HHT, HTH, HTT\} \cup \{HHH, HHT, HTH, THH\}$ $= \{HHH, HHT, HTH, HTT, THH\}$ $B \cup C = \{HHH, HHT\} \cup \{HHT, HTH, THH\} = \{HHH, HHT, HTH, THH\}$ $B \cup D = \{HHH, HHT\} \cup \{HHH, HHT, HTH, THH\} = D (B \text{ is a subset of } D)$

 $C \cup D = \{HHT, HTH, THH\} \cup \{HHH, HHT, HTH, THH\} = D \ (C \text{ is a subset of } D)$

The **intersection** of events A and B, denoted $A \cap B$, is the collection of elementary outcomes which belong to both A and B. This is illustrated in Figure 3.6. In symbols, $A \cap B = \{\omega \in \Omega : \omega \in A \text{ and } \omega \in B\}$

In words, $A \cap B$ is the event "A and B" in the sense that both A and B occur.





Example 3.6a Tossing a die once (continued). Recall that $A = \{1, 2, 3, 4\}$, $B = \{1, 3, 4\}, C = \{5, 6\}, \text{ and } D = \{3, 4, 5, 6\}$. The intersections of these events are: $A \cap B = \{1, 2, 3, 4\} \cap \{1, 3, 4\} = \{1, 3, 4\} = B$ (B is a subset of A) $A \cap C = \{1, 2, 3, 4\} \cap \{1, 3, 4\} = \{1, 3, 4\} = B$ (B is a subset of A) $A \cap D = \{1, 2, 3, 4\} \cap \{5, 6\} = \emptyset$ (A and C are mutually exclusive (disjoint)). $A \cap D = \{1, 2, 3, 4\} \cap \{3, 4, 5, 6\} = \{3, 4\}$ $B \cap C = \{1, 3, 4\} \cap \{3, 4, 5, 6\} = \{3, 4\}$ $C \cap D = \{5, 6\} \cap \{3, 4, 5, 6\} = \{5, 6\} = C$ (C is a subset of D) Example 3.6b Tossing a coin three times (continued). Recall that $A = \{HHH, HHT, HTH, HTT\}, B = \{HHH, HHT\}, C = \{HHT, HTH, THH\},$ $D = \{HHH, HHT, HTH, HTT\} \cap \{HHH, HHT\} = B$ (B is a subset of A)

 $A \cap C = \{HHH, HHT, HTH, HTT\} \cap \{HHT, HTH, THH\} = C \ (C \text{ is a subset of } A)$ $A \cap D = \{HHH, HHT, HTH, HTT\} \cap \{HHH, HHT, HTH, THH\} = \{HHH, HHT, HTH\}$

 $B \cap C = \{HHH, HHT\} \cap \{HHT, HTH, THH\} = \{HHT\}$

 $B \cap D = \{HHH, HHT\} \cap \{HHH, HHT, HTH, THH\} = B$ (B is a subset of D)

 $C \cap D = \{HHT, HTH, THH\} \cap \{HHH, HHT, HTH, THH\} = C \ (C \text{ is a subset of } D)$

Substantial simplifications of probability calculations are often possible using the representations of the complement of a union and the complement of an intersection contained in DeMorgan's laws.

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DeMorgan's laws. For any pair of events A and B:

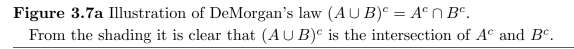
(1) The complement of the union of the events is the intersection of their complements. This is illustrated in Figure 3.7. In symbols, $(A \cup B)^c = A^c \cap B^c$

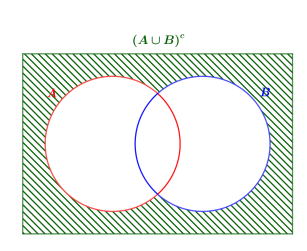
(2) The complement of the intersection of the events is the union of their complements. In symbols, $(A \cap B)^c = A^c \cup B^c$.

Example 3.6 Tossing a die once (continued). Recall that $\Omega = \{1, 2, 3, 4, 5, 6\}, A = \{1, 2, 3, 4\}, B = \{1, 3, 4\}, C = \{5, 6\}, \text{ and } D = \{3, 4, 5, 6\}.$ We will use A and B to demonstrate DeMorgan's laws. The complements are $A^c = \{5, 6\}, B^c = \{2, 5, 6\}.$

First consider $A \cup B = \{1, 2, 3, 4\}$. For these events the complement of the union is $(A \cup B)^c = \{1, 2, 3, 4\}^c = \{5, 6\}$ and the intersection of the complements is $A^c \cap B^c = \{5, 6\} \cap \{2, 5, 6\} = \{5, 6\}$.

Next consider $A \cap B = \{1, 3, 4\}$. For these events the complement of the intersection is $(A \cap B)^c = \{1, 3, 4\}^c = \{2, 5, 6\}$ and the union of the complements is $A^c \cup B^c = \{5, 6\} \cup \{2, 5, 6\} = \{2, 5, 6\}$.





 A^c is red lower left to upper right hatched B^c is blue upper left to lower right hatched $A^c \cap B^c$ is cross-hatched

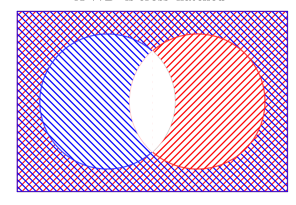
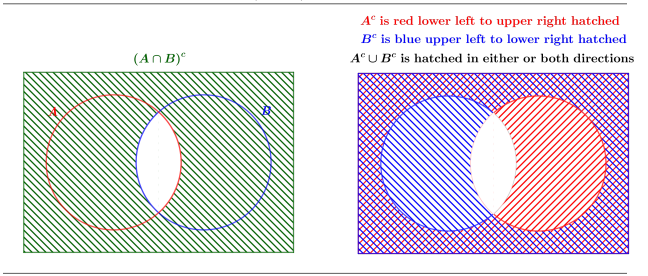


Figure 3.7b Illustration of DeMorgan's law $(A \cap B)^c = A^c \cup B^c$. From the shading it is clear that $(A \cap B)^c$ is the union of A^c and B^c .



Example 3.6b Tossing a coin three times (continued). We will use A and D to demonstrate DeMorgan's laws. Recall that $A = \{HHH, HHT, HTH, HTT\}$

and $D = \{HHH, HHT, HTH, THH\}.$

First consider $A \cup D = \{HHH, HHT, HTH, HTT, THH\}$ "the first toss yields a head OR there are at least two heads". The complement of this union is $(A \cup D)^c = \{THT, TTH, TTT\}$, that is, "the first toss does not yield a head AND there are less than two heads". Notice that this is the intersection of the complements $A^c = \{THH, THT, TTH, TTT\}$ "the first toss yields tails (not heads)" and $D^c = \{HTT, THT, TTH, TTT\}$: "there are less than two heads" Thus, $(A \cup D)^c = A^c \cap D^c$.

Next consider $A \cap D = \{HHH, HHT, HTH\}$ "the first toss yields a head AND there are at least two heads". The complement of this intersection is

 $(A \cap D)^c = \{HTT, THH, THT, TTH, TTT\}$ "the first toss does not yield a head OR there are less than two heads". Notice that this is the union of the complements $A^c = \{THH, THT, TTH, TTT\}$ "the first toss yields tails (not heads)"

and $D^c = \{HTT, THT, TTH, TTT\}$: "there are less than two heads" Thus, $(A \cap D)^c = A^c \cup D^c$.

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Some properties of set operations

Some basic properties of set operations are summarized here for completeness and ease of reference.

Commutative properties. For any events A and B

(a) $A \cup B = B \cup A$ (b) $A \cap B = B \cap A$

Associative properties. For any events A, B, and C

(a) $(A \cup B) \cup C = A \cup (B \cup C)$ (thus the notation $A \cup B \cup C$ is unambiguous) (b) $(A \cap B) \cap C = A \cap (B \cap C)$ (thus the notation $A \cap B \cap C$ is unambiguous)

Distributive properties. For any events A, B, and C (a) $A \cap (B \cup C) = (A \cap B) \cup (A \cap C)$

 $(b) \ A \cup (B \cap C) = (A \cup B) \cap (A \cup C)$

Additional properties.

For any event A: $A \cap A = A$, $A \cup A = A$, $A \cap \Omega = A$, $A \cup \Omega = \Omega$, $A \cap \emptyset = \emptyset$, $A \cup \emptyset = A$, $(A^c)^c = A$, $A \cap A^c = \emptyset$, and $A \cup A^c = \Omega$. If $A \subset B$, then $A \cap B = A$ and $A \cup B = B$.

3.4 Partitioning an event

Given two events, we will often find it useful to use one of the events to decompose the other event into disjoint subevents. A decomposition of an event into disjoint subevents yields a **partition** of the event. Given events A and B, we can partition (decompose) A into the subevents $A \cap B$ and $A \cap B^c$. These two subevents form a partition of A, since they are disjoint (they cannot share any outcomes since $A \cap B$ is a subset of B while $A \cap B^c$ is a subset of B^c) and their union is A (each outcome in A is either an element of B or an element of its complement B^c).

In symbols, $A = (A \cap B) \cup (A \cap B^c)$ and $(A \cap B) \cap (A \cap B^c) = \emptyset$.

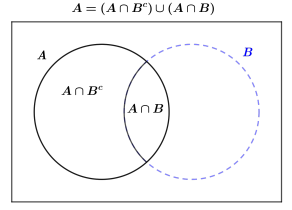
In words, there are two mutually exclusive ways in which the event A can occur:

(1) Event A and event B both occur $(A \cap B \text{ occurs})$; or,

(2) Event A occurs but event B does not occur $(A \cap B^c \text{ occurs})$.

This basic decomposition of A is illustrated in Figure 3.8. Figures 3.9 and 3.10 illustrate how this relationship can be used to partition a union of two or three events.

Figure 3.8 Use of event B to decompose event A into two disjoint parts.



Example 3.7 Characteristics of a person. Consider the experiment of selecting a person from a population of adults. Let A denote the event that the person selected is a college student and let B denote the event that the person is 20 years old. The decomposition $A = (A \cap B) \cup (A \cap B^c)$ indicates that a college student is either 20 years old, event $A \cap B$, or some other age, event $A \cap B^c$. The fact that this is a partition of A simply indicates that there is no other option, *i.e.* the college student is either 20 or not.

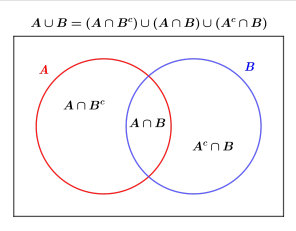
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The Venn diagram of Figure 3.9 shows a partition (decomposition) of $A \cup B$ into three disjoint parts. For this partition we have $A \cup B = (A \cap B^c) \cup (A \cap B) \cup (A^c \cap B)$. This shows the three mutually exclusive ways in which the event $A \cup B$ can occur:

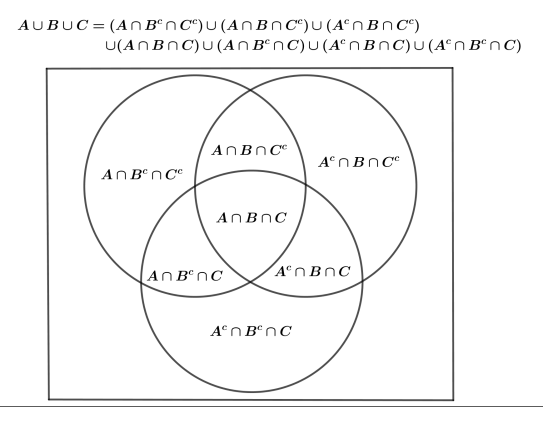
- (1) (event $A \cap B^c$) A occurs and B does not occur;
- (2) (event $A \cap B$) both A and B occur; or,
- (3) (event $A^c \cap B$) B occurs and A does not occur.

Figure 3.9 Decomposition of $A \cup B$ into three disjoint parts.



Example 3.7 Characteristics of a person, revisited. As before, consider the experiment of selecting a person from a population of adults. Let A denote the event that the person selected is a college student and let B denote the event that the person is 20 years old. The union $A \cup B$ is the event that the person selected is a college student or age 20 or both. This union is partitioned into the three disjoint events: event $A \cap B^c$ the person is a college student but is not 20 years old, event $A \cap B$ the person is a college student and is also 20 years old, and event $A^c \cap B$ the person is 20 years old but is not a college student.

Figure 3.10 Decomposition of $A \cup B \cup C$ into seven disjoint parts.



The Venn diagram of Figure 3.10 shows a partition (decomposition) of $A \cup B \cup C$ into seven disjoint parts. The seven mutually exclusive ways in which the event $A \cup B \cup C$ can occur are:

- (1) (event $A \cap B^c \cap C^c$) A occurs but B and C do not occur;
- (2) (event $A \cap B \cap C^c$) A and B occur but C does not occur;
- (3) (event $A^c \cap B \cap C^c$) B occurs but A and C do not occur;
- (4) (event $A \cap B \cap C$) A and B and C all occur;
- (5) (event $A \cap B^c \cap C$) A and C occur but B does not occur;
- (6) (event $A^c \cap B \cap C$) B and C occur but A does not occur; or,
- (7) (event $A^c \cap B^c \cap C$) C occurs but A and B do not occur.

Example 3.7 Characteristics of a person, revisited. As before, consider the experiment of selecting a person from a population of adults. Let A denote the event that the person selected is a college student, let B denote the event that the person is 20 years old, and let C denote the event that the person owns a car. The union $A \cup B \cup C$ is the event that the person selected is a college student or age 20 or owns a car, *i.e.*, the person possesses

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at least one of these three characteristics. This union is partitioned into the seven disjoint events:

- (1) (event $A \cap B^c \cap C^c$) is a college student, is not age 20, and does not own a car;
- (2) (event $A \cap B \cap C^c$) is a college student, is age 20, and does not own a car;
- (3) (event $A^c \cap B \cap C^c$) is not a college student, is age 20, and does not own a car;
- (4) (event $A \cap B \cap C$) is a college student, is age 20, and owns a car;
- (5) (event $A \cap B^c \cap C$) is a college student, is not age 20, and owns a car;
- (6) (event $A^c \cap B \cap C$) is not a college student, is age 20, and owns a car;
- (7) (event $A^c \cap B^c \cap C$) is not a college student, is not age 20, and owns a car;

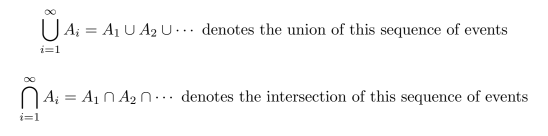
An aside – notation for more complicated unions and intersections

On occasion it is useful to have a compact notation for the union or intersection of a collection of events.

Given a set of n events $\{A_1, \ldots, A_n\}$:

$$\bigcup_{i=1}^{n} A_{i} = A_{1} \cup A_{2} \cup \dots \cup A_{n} \text{ denotes the union of these events}$$
$$\bigcap_{i=1}^{n} A_{i} = A_{1} \cap A_{2} \cap \dots \cap A_{n} \text{ denotes the intersection of these events}$$

Given an infinite sequence of events $\{A_1, A_2, \ldots\}$:



3.5 Problems

The problem sets and solutions can be found on the book web page at https://www.ucs.louisiana.edu/~jcb0773/.

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4 Probability measure

4.1 Definition of a probability measure

Given an experiment and an event A we need to associate a probability P(A) with the event. The formal (axiomatic) definition of a **probability measure** below indicates the restrictions we will impose on any such assignment of probabilities to events.

Definition. A probability measure P is a function which assigns probabilities to events (subsets of Ω) and satisfies the following axioms.

Axiom 1: For every event $A, 0 \leq P(A) \leq 1$. Axiom 2: $P(\Omega) = 1$. Axiom 3: For every finite collection of mutually exclusive events A_1, A_2, \ldots, A_n , $P(A_1 \cup A_2 \cup \cdots \cup A_n) = P(A_1) + P(A_2) + \cdots + P(A_n)$. Furthermore, for every infinite sequence of mutually exclusive events $\{A_1, A_2, \ldots\}$, $P(A_1 \cup A_2 \cup \cdots) = P(A_1) + P(A_2) + \cdots$, more formally $P(\bigcup_{i=1}^{\infty} A_i) = \sum_{i=1}^{\infty} P(A_i)$.

The meaning and desirability of these axioms is straightforward. The first two axioms place some basic restrictions on the possible values of probabilities. Axiom 1 simply requires that the probability of an event must be a number between zero and one. (If we think of probabilities as percentages, this says that a probability must be between zero and 100 percent.) By assigning the value 1 (100%) to the sample space, Axiom 2 simply requires that something must happen when the experiment is conducted. The third axiom requires a certain type of consistency in the assignment of probabilities. Recall that when events are mutually exclusive (disjoint), they cannot occur at the same time. Axiom 3 states that if an event can be partitioned (decomposed) into a collection of mutually exclusive subevents (the A_i), then the probability of the event must be equal to the sum of the probabilities of the mutually exclusive subevents of the partition.

4.2 Properties of probability measures

In this section we will provide some basic properties of probability measures.

The null event has probability zero

As noted above, Axiom 2, $P(\Omega) = 1$, indicates that something must occur when the experiment is conducted. In other words, the sure event Ω must occur. As you would

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expect, this implies that the null event \emptyset , will not occur. Formally this is indicated by saying that the null event has probability zero. That is,

$$P(\emptyset) = 0.$$

(Technical remark: This result follows from the infinite sequence part of axiom 3.)

The probability of the complement of an event

For any event A, the sample space can be partitioned as $\Omega = A \cup A^c$. Thus, by axioms 2 and 3 we see that for any event A, $P(\Omega) = P(A) + P(A^c) = 1$. It follows that the probability of the complement of an event is one minus the probability of the event. In symbols,

$$P(A^c) = 1 - P(A).$$

An important decomposition of the probability of an event

In Figure 3.8 we illustrated the use of one event (event B) to partition another event (event A), viz, $A = (A \cap B) \cup (A \cap B^c)$. This partition and axiom 3 imply that for any events A and B,

$$P(A) = P(A \cap B) + P(A \cap B^c).$$

Probabilities of nested events

Recall that if $A \subset B$, then the occurrence of A implies the occurrence of B. Therefore, if A is a subset of B, then the probability of B cannot be less than the probability of A. More formally, if $A \subset B$, then $A \cap B = A$ and the decomposition we just discussed becomes $P(B) = P(A \cap B) + P(A^c \cap B) = P(A) + P(A^c \cap B)$. Since $P(A^c \cap B) \ge 0$, it follows that

if
$$A \subset B$$
, then $P(A) \leq P(B)$.

We will now provide some important expressions for **the probability of a union** in terms of the probabilities of certain subevents.

The probability of the union of 2 events

In Figure 3.9 we illustrated the partition of $A \cup B$ into three disjoint parts, *viz*,

 $A \cup B = (A \cap B^c) \cup (A \cap B) \cup (A^c \cap B)$. This partition implies that

 $P(A\cup B)=P(A\cap B^c)+P(A\cap B)+P(A^c\cap B). \text{ The partitions } A=(A\cap B)\cup (A\cap B^c)$

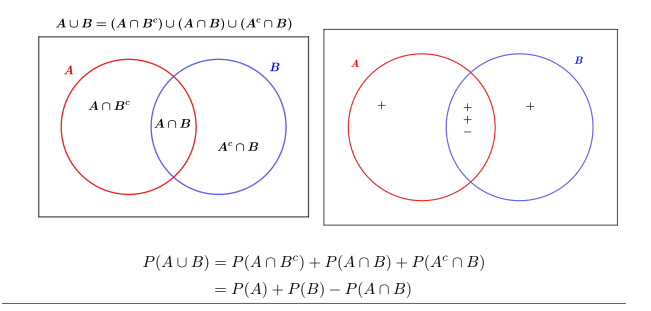
and $B = (A \cap B) \cup (A^c \cap B)$ imply that $P(A) = P(A \cap B^c) + P(A \cap B)$ and

 $P(B) = P(A \cap B) + P(A^c \cap B)$. These observations lead to the expression

$$P(A \cup B) = P(A) + P(B) - P(A \cap B)$$

for the union of any two events A and B. This decomposition is illustrated in Figure 4.1.

Figure 4.1 Illustration – **the union of 2 events** The pluses and minuses indicate how the probabilities for each section enter into and are removed from the sum.



Example 4.1 Dogs and cats. Among households in the United States (in 2006), 44% of the households have a dog, 29% have a cat, 17% have both, and 44% do not have a dog or a cat. Let D denote the event that a household has a dog and let C denote the event that a household has a dog and let C denote the event that a household at random, so that each household has the same chance of being selected, then P(D) = .44, P(C) = .29, and $P(D \cap C) = .17$. We will now find $P(D \cup C)$, the probability that a household has a dog or a cat. Application of the formula for the probability of a union yields $P(D \cup C) = P(D) + P(C) - P(D \cap C) = .44 + .29 - .17 = .56$, that is, 56% have a dog or a cat or both.

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Some readers may find the following detailed derivation of this probability instructive. Since $P(D) = P(D \cap C) + P(D \cap C^c)$, we find that $P(D \cap C^c) = P(D) - P(D \cap C) = .44 - .17 = .27$, that is, 27% have a dog but not a cat. Similarly $P(C) = P(D \cap C) + P(D^c \cap C)$, so that $P(D^c \cap C) = P(C) - P(D \cap C) = .29 - .17 = .12$, that is, 12% have a cat but not a dog. Combining these results we have

 $P(D \cup C) = P(D \cap C^{c}) + P(D \cap C) + P(D^{c} \cap C) = .27 + .17 + .12 = .56.$

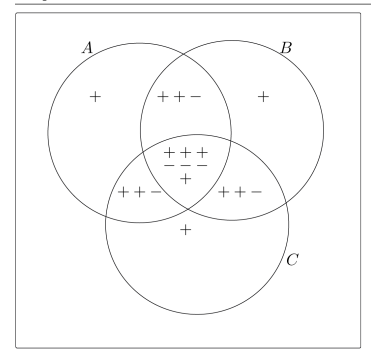
The probability of the union of 3 events

In Figure 3.10 we illustrated the partition of $A \cup B \cup C$ into seven disjoint parts, viz, $A \cup B \cup C = (A \cap B^c \cap C^c) \cup (A^c \cap B \cap C^c) \cup (A \cap B \cap C^c) \cup (A \cap B \cap C) \cup (A \cap B^c \cap C) \cup (A^c \cap B^c \cap C)) \cup (A^c \cap B^c \cap C)$. This partition implies that $P(A \cup B \cup C) = P(A \cap B^c \cap C^c) + P(A^c \cap B \cap C^c) + P(A \cap B \cap C^c) + P(A \cap B \cap C^c) + P(A \cap B \cap C)) + P(A^c \cap B \cap C) + P(A^c \cap B \cap C)) + P(A^c \cap B \cap C) + P(A^c \cap B \cap C))$. Similar decompositions for the probabilities of the events $A, B, C, A \cap B, A \cap C, B \cap C$, and $A \cap B \cap C$, and a bit of algebra yields the decomposition

$$P(A \cup B \cup C) = P(A) + P(B) + P(C) - P(A \cap B) - P(A \cap C) - P(B \cap C) + P(A \cap B \cap C).$$

This decomposition is illustrated in Figure 4.2.

Figure 4.2 Illustration – **the union of 3 events** The pluses and minuses indicate how the probabilities for each section enter into and are removed from the sum.



 $P(A \cup B \cup C) = P(A) + P(B) + P(C) - P(A \cap B) - P(A \cap C) - P(B \cap C) + P(A \cap B \cap C)$

An aside – The probability of the union of many events

The decomposition of the probability of 3 events above can be extended to 4 or more events. This extension is straightforward (but tedious) with alternating inclusions (pluses) and exclusions (minuses). Here is the expression for n events. For any events A_1, \ldots, A_n ,

$$P(A_1 \cup A_2 \cup \dots \cup A_n) = \sum_i P(A_i) - \sum_{i < j} P(A_i \cap A_j) + \sum_{i < j < k} P(A_i \cap A_j \cap A_k)$$
$$+ \dots + (-1)^{n+1} P(A_1 \cap \dots \cap A_n).$$

This expression indicates that the probability of the union is obtained by first adding the probabilities of each event, then subtracting the probabilities of each intersection of 2 events, then adding the probabilities of each intersection of 3 events, with this process continuing (alternating adding and subtracting) until the probability of the intersection of all n events is entered.

4.3 Probabilities on discrete sample spaces

A sample space Ω is said to be **discrete** if it contains a finite or countably infinite number of elementary outcomes. That is, either $\Omega = \{\omega_1, \ldots, \omega_N\}$ for some positive integer N or the elements of Ω can be arranged in a sequence $\Omega = \{\omega_1, \omega_2, \ldots\}$.

A probability distribution on a finite sample space $\Omega = \{\omega_1, \ldots, \omega_N\}$ is an assignment of probabilities to the elementary outcomes (elements) of Ω . More formally, given a finite sample space $\Omega = \{\omega_1, \ldots, \omega_N\}$, a collection of probabilities p_1, \ldots, p_N , with $0 \le p_i \le 1$ and $p_1 + \cdots + p_N = 1$, determines a probability distribution on Ω with $P(\omega_i) = p_i$ for $i = 1, \ldots, N$. Note that in most situations we can remove any elements with zero probability and there are at least two elements with positive probability, thus, we can assume that $0 < p_i < 1$.

Similarly, if $\Omega = \{\omega_1, \omega_2, \ldots\}$ is countably infinite, then a **probability distribution** on Ω is a sequence p_1, p_2, \ldots of probabilities $(P(\omega_i) = p_i)$ with $0 \le p_i \le 1$ for all i and $\sum_{i=1}^{\infty} p_i = p_1 + p_2 + \cdots = 1$.

Given an event A, *i.e.*, given $A \subset \Omega$, the probability of the event A is the sum of the probabilities of the elementary outcomes which belong to A, *i.e.*, if the elements of Ω are labeled so that $A = \{\omega_1, \omega_2, \ldots, \omega_m\}$, then

$$P(A) = P(\omega_1) + P(\omega_2) + \dots + P(\omega_m) = p_1 + p_2 + \dots + p_m.$$

The simplest way to assign probabilities to the elements of a finite sample space $\Omega = \{\omega_1, \ldots, \omega_N\}$ is to assume that the N elementary outcomes are **equally probable (equally likely**) so that $P(\omega_i) = 1/N$ for $i = 1, \ldots, N$. When the N elementary outcomes are assumed equally probable, the probability of an event A is P(A) = N(A)/N, where N(A) is the number of elementary outcomes which belong to A. In other words, with equally probable outcomes, the probability of event A is the ratio of the number of outcomes "favorable" for A to the number of "possible" outcomes. This simple situation is convenient for demonstrating concepts; but, the usefulness of the assumption of a finite sample space with equally probable outcomes as a model for an idealized version of reality is limited.

Example 4.2 Fiber breaks (from Schervish and DeGroot). Consider an experiment in which five fibers having different lengths are subjected to a testing procedure to see which will

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break first. Suppose that the lengths of these five fibers are 1, 2, 3, 4, and 5 inches, respectively. Suppose also that the probability that any given fiber will be the first to break is proportional to its length. We will find the probability that the length of the first fiber to break is no more than 3 inches.

For i = 1, ..., 5, let ω_i be the outcome that the fiber of length *i* inches breaks first. Then $\Omega = \{\omega_1, ..., \omega_5\}$ and $p_i = ki$ for i = 1, ..., 5, where *k* is a proportionality factor. Since we need $p_1 + ... + p_5 = 1$ and $k_1 + k_2 + k_3 + k_4 + k_5 = 15k$, we know that k = 1/15. Let *A* denote the event that the length of the first fiber to break is no more than 3 inches. Then $A = \{\omega_1, \omega_2, \omega_3\}$, and

$$P(A) = p_1 + p_2 + p_3 = \frac{1}{15} + \frac{2}{15} + \frac{3}{15} = \frac{2}{5}.$$

4.4 Problems

The problem sets and solutions can be found on the book web page at https://www.ucs.louisiana.edu/~jcb0773/.

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5 Combinatorics – counting

5.1 Counting basics

We will start with a basic (illustrative) counting problem. Consider a box containing 5 balls labeled with the integers 1, 2, 3, 4, 5. If we select one ball from the box, then, obviously, there are 5 possible outcomes. Let's consider a slightly more interesting setup. Suppose we select 2 balls from the box in the following way, first we select a ball and note its number, then we return the ball to the box and make a second selection. How many possible outcomes are there now? We can think of this selection process as an experiment with two stages and we can represent the possible outcomes as ordered pairs of the form (a, b)where a denotes the number on the first ball selected and b the number on the second. The answer is that there are 5 times 5 equals 25 possible outcomes. These 25 possible outcomes are listed in Table 5.1a. In this table there are 5 rows, one for each of the 5 possibilities at the first stage, and 5 columns, one for each of the 5 possibilities at the second stage.

Table 5.1a Ordered pairs (a)	b) with a	$b, b \in \{1,$	2, 3, 4, 5	5} (repeats allowed).	
(2,1) (3,1) (4,1)	$ \begin{array}{c} (1,2) \\ (2,2) \\ (3,2) \\ (4,2) \\ (5,2) \end{array} $	(2,3) (3,3) (4,3)	(2,4) (3,4) (4,4)	(2,5) (3,5) (4,5)	

How does the answer change if we do not return the first ball to the box before selecting the second? In this case, regardless of which ball was selected first, there are only 4 possibilities at the second stage. As shown in Table 5.1b, in terms of the listing in Table 5.1a, the ordered pairs on the diagonal, (1,1), (2,2), *etc.*, are not possible. Thus, in this case, there are 5 times 4 equals 20 possible outcomes. We will now formalize this fundamental rule of counting.

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Table 5.1bOrdered pairs (a, b)) with a	$a, b \in \{1,$	2, 3, 4, 5	5} and $a \neq b$ (repeats not allowed).
$(9 \ 1)$	(1,2)	(1,3)	· · ·	(1,5) (2,5)
(2,1) (3,1)		(2,3)		(2,5) (3,5)
(4,1)	(,)	(4,3) (5,3)	(5.4)	(4,5)
(0,1)	(0,2)	(0, 3)	(0,4)	

Fundamental rule of counting – multiplication rule for counting. If an experiment consists of two stages (parts) a first stage which can be performed in m ways, and, regardless of the particular outcome of the first stage, a second stage which can be performed in n ways, then the experiment itself can be performed in mn ways.

If we use ordered pairs to represent the possible outcomes of the experiment, as we did in Tables 5.1a and 5.1b, we have the following alternate statement.

Alternate statement of the fundamental rule of counting. Given *m* labels (outcomes) a_1, \ldots, a_m and *n* labels (outcomes) b_1, \ldots, b_n , there are *mn* ordered pairs of the form (a_i, b_j) with $i = 1, 2, \ldots, m$ and $j = 1, 2, \ldots, n$.

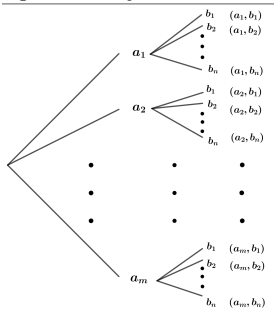


Figure 5.1 Multiplication rule: a tree diagram with mn paths.

We can also envision the choices in the two stages of the experiment as branches in the tree diagram of Figure 5.1. Here a path (choice of an *a* branch, say a_i , and choice of a *b* branch, say b_j) corresponds to the outcome associated with the label (a_i, b_j) . The fundamental rule of counting says that there are mn paths through this tree diagram. That is, there are mn paths through a tree diagram with m branches at the first level (stage) and n branches at the second level (stage).

Example 5.1 Some basic counting problems.

1. Each year starts on one of the seven days (Sunday through Saturday). Each year is either a leap year (*i.e.*, it includes February 29) or not. How many different calendars are possible for a year?

2. John has 4 clean shirts and 2 clean pairs of jeans. How many clean shirt / clean jeans combinations does John have to choose from?

solutions

1. There are two steps in forming a calendar. First we need to select a day of the week for January 1 (7 choices) and then we need to decide whether to include February 29 (2 choices). Thus there are $7 \times 2 = 14$ possible calendars.

2. John has 4 choices for his shirt and 2 choices for his jeans. Therefore, John has $4 \times 2 = 8$ clean shirt / clean jeans combinations to choose from.

Extension of the fundamental rule of counting. If an experiment consists of k stages, so that, for i = 1, ..., k, regardless of the exact outcomes of the other stages, the i^{th} stage can be performed in n_i ways, then the experiment itself can be performed in $n_1n_2 \cdots n_k$ ways. In other words, thinking of the possible outcomes as ordered k-tuples, there are $n_1n_2 \cdots n_k$ ways in which an ordered k-tuple can be formed when there are n_1 choices for the first element, n_2 choices for the second element, and so on ending with n_k choices for the k^{th} element.

Example 5.2 Some more basic counting problems. Suppose that a license plate "number" consists of a combination of three letters followed by four digits, such as ABC-1234.

1. How many such license plate "numbers" are possible if there is no restriction on the choices for the letters or digits?

2. How many such license plate "numbers" are possible if the three letters must be distinct

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(three different letters) but there is no restriction on the choices the digits?

3. How many such license plate "numbers" are possible if the three letters must be distinct and the four digits must be distinct?

solutions

There are 26 letters (A, B, \ldots, Z) and ten digits $(0, 1, \ldots, 9)$.

1. With no restrictions, there are 26 choices for each letter and 10 for each digit resulting in $26 \cdot 26 \cdot 26 \cdot 10 \cdot 10 \cdot 10 = 175,760,000$ possible license plate "numbers".

2. If the letters must be distinct, then at each letter selection stage there is one fewer choices. Hence, there are $26 \cdot 25 \cdot 24 \cdot 10 \cdot 10 \cdot 10 = 156,000,000$ possible license plate "numbers".

3. Similarly, if the letters must be distinct and the digits must be distinct, then there are $26 \cdot 25 \cdot 24 \cdot 10 \cdot 9 \cdot 8 \cdot 7 = 78,624,000$ possible license plate "numbers".

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5.2 Ordered samples

Consider a population (collection) consisting of N distinct objects.

An ordered sample of size n is an ordered collection of n objects selected from the N objects in the population. An ordered sample of this type can be represented by an ordered n-tuple. The multiplication rule can be used to determine the number of possible ordered samples. We can think of the selection of an ordered sample of size n as a sequence of n steps, where the first element of the sample is selected at step one, the second element is selected at step two, and so on until all n elements have been selected. Since the selection of the sample will entail n such steps the multiplication rule will yield the number of possible samples as a product of n values (one for each of the n steps).

Before we start counting, notice that there are two ways in which a sample can be selected. The elements of the sample can be **selected with replacement**, so that at each step the element is selected from the entire population, or the elements of the sample can be **selected without replacement**, so that at each step an object is removed from the population and there is one less object in the population for the next selection. When sampling with replacement a particular object can appear more than once in the sample and there is no restriction on the sample size n. When sampling without replacement no particular object can appear more than once in the sample size n clearly cannot exceed the population size N.

When sampling with replacement there are N choices at each of the n steps. Thus there are N^n possible ordered samples of size n when the sample is selected with replacement from a population of size N. This result is illustrated for a population of size N = 4 and a sample of size n = 3 in Table 5.2.

Table 5.2	The $4^3 = 6$	64 ordered	samples o	f size $n = 3$, sel	ected with	ı replacem	ent
	from $\{a, b\}$	$, c, d\}$					
(a, a, a)	(a, a, b)	(a, a, c)	(a, a, d)	(a,b,a)	(a, b, b)	(a,b,c)	(a, b, d)
(a,c,a)	(a,c,b)	(a,c,c)	(a, c, d)	(a,d,a)	(a,d,b)	(a, d, c)	(a, d, d)
(b, a, a)	(b, a, b)	(b, a, c)	(b, a, d)	(b,b,a)	(b,b,b)	(b,b,c)	(b, b, d)
(b,c,a)	(b,c,b)	(b,c,c)	(b,c,d)	(b,d,a)	(b,d,b)	(b, d, c)	(b, d, d)
(c, a, a)	(c, a, b)	(c, a, c)	(c, a, d)	(c, b, a)	(c,b,b)	(c, b, c)	(c, b, d)
(c,c,a)	(c,c,b)	(c,c,c)	(c,c,d)	(c,d,a)	(c,d,b)	(c,d,c)	(c, d, d)
(d, a, a)	(d, a, b)	(d, a, c)	(d, a, d)	(d, b, a)	(d, b, b)	(d, b, c)	(d, b, d)
(d, c, a)	(d, c, b)	(d, c, c)	(d, c, d)	(d, d, a)	(d, d, b)	(d, d, c)	(d, d, d)

When sampling without replacement there are N choices at the first step, N-1 choices at the second step, and so on, with N - n + 1 choices at the n^{th} step (assuming that $n \leq N$). Thus, assuming that $n \leq N$, there are $N(N-1)\cdots(N-n+1)$ possible ordered samples of size n when the sample is selected without replacement from a population of size N. Notice that, as before, there are n terms in this product. This result is illustrated for a population of size N = 4 and a sample of size n = 3 in Table 5.3.

	fr	om $\{a, b, c$	$,d\}$	I I		,
_	(a, b, c)	(a, b, d)	(a, c, b)	(a, c, d)	(a, d, b)	(a,d,c)
	(b, a, c)	(b, a, d)	(b,c,a)	(b,c,d)	(b,d,a)	(b,d,c)
	(c, a, b)	(c, a, d)	(c, b, a)	(c,b,d)	(c,d,a)	(c,d,b)
	(d, a, b)	(d, a, c)	(d, b, a)	(d, b, c)	(d, c, a)	(d,c,b)

Table 5.3 The $4 \cdot 3 \cdot 2 = 24$ ordered samples of size n = 3, selected without replacement

For ease of reference these counting results are summarized below.

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The number of ordered samples. Refer to the ordered samples of Tables 5.2 and 5.3 for an illustration of this result. For a population of size N:

(1) There are N^n ordered samples of size n when the sample is selected with replacement. (2) For $n \leq N$, there are $N(N-1)\cdots(N-n+1)$ ordered samples of size n when the sample is selected without replacement.

Example 5.3 Some basic ordered sample counting problems. Consider a club consisting of 35 members.

1. In how many ways can this club select three officers (a president, a secretary, and a treasurer), if the three officers are required to be three different members?

2. How does the answer to part 1 change if we allow one person to hold two or even three offices?

solutions

We can use an ordered triple (a, b, c) to represent the choice of the officers (a is president, b is secretary, c is treasurer). Since there are 35 members and we need 3 officers, N = 35and n = 3.

1. If a member can hold at most one office, then there are $N(N-1)(N-2) = 35 \cdot 34 \cdot 33 = 39,270$ ways to select the three officers.

2. If a member is allowed to hold two or more offices, then there are $N^n = 35^3 = 42,875$ ways to select the three officers.

An ordered sample selected without replacement can also be viewed as an ordered arrangement of a set of objects. A **permutation of a set** is an ordered arrangement of the elements of the set. Note that if an ordered sample of size n = N is selected without replacement from a population of size N, then the ordered sample is a permutation (ordered arrangement) of the N objects which comprise the population. There are $N! = N(N-1)\cdots 1$ (read this as N factorial) permutations of N objects. By convention 0! = 1.

The $24 = 4 \cdot 3 \cdot 2$ ordered samples of size n = 3 selected without replacement from $\{a, b, c, d\}$ listed in Table 5.3 are the permutations of the four letters $\{a, b, c, d\}$ taken three at a time. In general, the **permutations of** N **objects taken** n **at a time** are the ordered samples of size n selected without replacement from a population of N objects. Thus there are $N(N-1)\cdots(N-n+1)$ permutations of N objects taken n at a time. This result is restated below for ease of reference.

The number of permutations – ordered samples. The number of permutations of N objects taken n at a time, which is also the number of ordered samples of size n selected without replacement from a population of size N, is equal to

$$_{N}P_{n} = \frac{N!}{(N-n)!} = N(N-1)\cdots(N-n+1).$$

Notice that there are n terms in the product determining the value of $_NP_n$.

Example 5.4 Arranging people.

1. In how many ways can 6 people be seated in a row of 6 theater seats?

We can represent a seating arrangement by a 6-tuple $(x_1, x_2, x_3, x_4, x_5, x_6)$, where x_i is the label of the seat assigned to person *i*. There are 6 choices for the first person's seat, 5 choices for the second person's seat, and so on, down to 1 choice for the sixth person's seat. Thus, the number of ways to seat 6 people in a row of 6 theater seats is $6! = 6 \cdot 5 \cdot 4 \cdot 3 \cdot 2 \cdot 1 = 720$. Notice that this is the number of permutations of 6 objects and can also be written as $_6P_6$.

2. Now suppose that there are 12 seats in the row. In how many ways can 6 people be seated in a row of 12 theater seats?

Arguing as before, in this situation there are 12 choices for the first person's seat, 11 choices for the second person's seat, and so on, down to 7 choices for the sixth person's seat. Thus, the number of ways to seat 6 people in a row of 12 theater seats is ${}_{12}P_6 = 12 \cdot 11 \cdot 10 \cdot 9 \cdot 8 \cdot 7 = 665, 280$.

Calculator aside: Many calculators use the notation nPr (the number of permutations of n objects taken r at a time). For these calculators 6! = (6 nPr 6) and ${}_{12}P_6 = 12 \cdot 11 \cdot 10 \cdot 9 \cdot 8 \cdot 7 = (12 nPr 6)$.

Example 5.5 Choosing different numbers.

Consider a box containing N balls, numbered 1, 2, ..., N. Suppose a sample of n balls is selected at random with replacement from this box. Assuming that the N^n possible ordered samples of size n are equally likely, we will find the probability P that the numbers on the n balls are all different? If n > N, then P = 0, since the sample size is greater than the

78 5.2 Ordered samples

number of balls in the box and at least one number must occur more than once. If $n \leq N$, then the $_N P_n$ permutations of N objects taken n at a time are the samples of size n for which the the numbers on the n balls are all different. Hence, the probability of interest is

$$P = \frac{{}_{N}P_{n}}{N^{n}} = \frac{N!}{(N-n)! N^{n}} = \frac{N(N-1)\cdots(N-n+1)}{N^{n}}.$$

We will consider an interesting application of this result in the next example.

Example 5.6 The birthday problem. In this example we will find the probability P_n that at least two people in a group of n people have the same birthday (month and day). We need to make a few simplifying assumptions. First assume that there are no twins in the group and assume that no one in the group was born on February 29. Next assume that the 365 birthdays are equally likely. We will use the argument from the preceding example to solve the birthday problem. The complement of the event that at least two people in the group of n have the same birthday is the event that no two people have the same birthday. Since there are N = 365 possible birthdays, we have

$$P_n = 1 - \frac{365P_n}{365^n} = 1 - \frac{365!}{(365-n)! \ 365^n}$$

Numerical values of this probability for several choices of n are given in Table 5.4. If you have never encountered this problem before, you may find these numbers surprisingly large. If fact, you might reasonably guess that we would need a large number of people, say n > 100, in order for P_n to be greater than 1/2. However, you can see that we only need n = 23 people in the group to get $P_n > 1/2$. Notice that for n = 50 people there is a 97% chance that at least two people have the same birthday!

Table 5.4 The probability P_n that at least two people in a group of n people					
will have the same birth	day.				
n	P_n	n	P_n		
5	0.027	23	0.507		
10	0.117	25	0.569		
15	0.253	30	0.706		
20	0.411	40	0.891		
22	0.476	50	0.970		

5.3 Unordered samples

An unordered sample of size n is a collection (set) of n objects selected without replacement from the N objects in the population. In other words, an unordered sample is a subset or subpopulation of the original population. We can use set notation to represent an unordered sample. We will now determine, for fixed n < N, the number of unordered samples of size n that can be formed from the objects in a population of size N. Note that the selection of a unordered sample of size n can also be viewed as a partitioning (division) of the population into two complementary subpopulations — one of size n (the objects in the sample) and the other of size N - n (the objects not in the sample).

A combination of N objects taken n at a time is an unordered sample of size n selected without replacement from a population of N objects, *i.e.*, a subpopulation (subset) of size n. We will now find the number of such combinations.

We know that there are ${}_{N}P_{n} = N(N-1)\cdots(N-n+1)$ ordered samples of size n selected without replacement from a population of size N, *i.e.*, there are ${}_{N}P_{n}$ permutations of N objects taken n at a time. Each of these ordered samples of size n can be ordered (permuted) in n! ways, *i.e.*, there are n! permutations of a particular set of n distinct objects. This connection is illustrated in the table of Example 5.7 where the ${}_{4}P_{3} = 24$ permutations of $\{a, b, c, d\}$ taken 3 at a time are arranged in rows of 3! = 6 permutations, one row for each of the 24/6 = 4 combinations.

Example 5.7 Permutations and combinations. The connection between the permutations of the four letters $\{a, b, c, d\}$ taken three at a time and the combinations of the four letters $\{a, b, c, d\}$ taken three at a time is indicated in this table. Within each row, the first 6 columns contain the 6 permutations of the combination in column 7.

The permutations and combinations of the four letters $\{a, b, c, d\}$ taken three at a time.						
permutations						combinations
	(•	ordered s	amples)			(unordered samples)
(a,b,c)	(a,c,b)	(b, a, c)	(b, c, a)	(c, a, b)	(c,b,a)	$\{a,b,c\}$
(a, b, d)	(a,d,b)	(b, a, d)	(b, d, a)	(d, a, b)	(d,b,a)	$\{a,b,d\}$
(a, c, d)	(a, d, c)	(c, a, d)	(c, d, a)	(d, a, c)	(d,c,a)	$\{a,c,d\}$
(b,c,d)	(b, d, c)	(c, b, d)	(c,d,b)	(d,b,c)	(d,c,b)	$\{b,c,d\}$

toc

If we divide the number of ordered samples of size n by the number of ways that each set of n objects can be ordered, then we obtain the number of unordered samples. For example, there are four combinations in the table of Example 5.7 since ${}_{4}P_{3}/3! = 24/6 = 4$. A general version of this expression is given below.

The number of combinations – unordered samples. The number of combinations of N objects taken n at a time, which is also the number of unordered samples of size n selected without replacement (number of subpopulations of size n) from a population of size N, is given by the binomial coefficient (read $\binom{N}{n}$ as N choose n)

$$\binom{N}{n} = \frac{N(N-1)\cdots(N-n+1)}{n(n-1)\cdots1} = \frac{N!}{n!(N-n)!}$$

The terminology N choose n indicates that $\binom{N}{n}$ is the number of ways to choose n objects without replacement from N objects. The binomial coefficient $\binom{N}{n}$ is sometimes denoted by ${}_{N}C_{n}$ with C indicating combinations. Note that in the ratio of products expression for $\binom{N}{n}$ the numerator product $N(N-1)\cdots(N-n+1)$ and the denominator product n! both contain n terms.

Calculator aside: Many calculators use the notation nCr (the number of combinations of n objects taken r at a time). For these calculators $\binom{N}{n} = {}_{N}C_{n} = (N \ nCr \ n), \ e.g.,$ $\binom{7}{3} = \frac{7 \cdot 6 \cdot 5}{3 \cdot 2 \cdot 1} = 35 = (7 \ nCr \ 3).$

As noted above, a combination of N objects taken n at a time can also be viewed as a subpopulation of size n selected from a population of size N. Thus the **binomial coefficient** $\binom{N}{n}$ is the number of subpopulations of size n from a population of size N. Note also that selecting a subpopulation of size n from a population of size N is equivalent to selecting a collection of N - n objects which are excluded from the subpopulation. Thus, recalling the convention 0! = 1, which leads to the convention $\binom{N}{0} = \binom{N}{N} = 1$, for $0 \le n \le N$, we have the identity

$$\binom{N}{n} = \binom{N}{N-n}.$$

Continuing along this line of thought we see that there are $\binom{N}{n}$ ways to partition a population of size N into a primary subpopulation of size n and a complementary subpopulation

of size N - n. Note that this method of counting partitions takes the order of the subpopulations into account. For example, with N = 4 and n = 2 the six ordered partitions, expressed as ordered pairs, are:

 $(\{a,b\},\{c,d\}),(\{a,c\},\{b,d\}),(\{a,d\},\{b,c\}),(\{b,c\},\{a,d\}),(\{b,d\},\{a,c\}),(\{c,d\},\{a,b\}).$

We will now extend the binomial coefficient to provide a general expression for the number of partitions of a population. For $m \ge 2$ and n_1, \ldots, n_m such that $n_1 + \cdots + n_m = N$, we might ask: In how many ways can a population of size N be partitioned into msubpopulations of respective sizes n_1, \ldots, n_m ? (Note that, as with the case when m = 2and $n_1 = n_2$, there is an implied ordering of the subpopulations in this statement.) To answer this question consider the formation of such a partition via a sequence of m steps. At the first step there are N objects to choose from and $\binom{N}{n_1}$ ways to select the first subpopulation. At the second step there are $N - n_1$ objects to choose from and $\binom{N-n_1}{n_2}$ ways to select the second subpopulation. Continuing with this reasoning when we reach the last (m^{th}) step we find that there are $N - n_1 - \cdots - n_{m-1} = n_m$ objects to choose from and $\binom{n_m}{n_m} = 1$ ways to select the final subpopulation. The answer to our question is the **multinomial coefficient** given in the following result.

The number of partitions. Consider a population of size N. For n_1, \ldots, n_m such that each $n_i \ge 1$ and $n_1 + \cdots + n_m = N$, the number of partitions of the population into m subpopulations of respective sizes n_1, \ldots, n_m is given by the multinomial coefficient

$$\binom{N}{n_1, n_2, \dots, n_m} = \binom{N}{n_1} \binom{N-n_1}{n_2} \binom{N-n_1-n_2}{n_3} \cdots \binom{n_m}{n_m} = \frac{N!}{n_1!n_2!\cdots n_m!}$$

Note that if we allow $n_i = 0$ in this theorem, then the expression is still valid but the number of nontrivial subpopulations in the partition is reduced by the number of *i* for which $n_i = 0$. Note also that a binomial coefficient is a special case of a multinomial coefficient. In particular,

$$\binom{N}{n,N-n} = \binom{N}{n}.$$

Example 5.7a An application. A child has 20 colored beads, of which 9 are red, 5 are green, 4 are blue, and 2 are black. If the child puts the beads on a string to form a necklace, how many arrangements are possible?

We will assume that the beads of the same color are indistinguishable (all look the same). The elementary outcomes can be represented by ordered sequences of the form

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(*RRRRRRRGGGGGBBBBKK*), where *R*, *G*, *B*, and *K* denote the respective colors red, green, blue, and black. As noted above, switching the positions of two beads of the same color does not result in a new outcome since such a change would be undetectable once the beads were on the string. We will count by selecting positions in the sequence (on the string) sequentially for the four colors. Initially there are 20 positions to choose from. Thus, there are $\binom{20}{9}$ choices for the 9 red beads. Once this is done, there are 11 positions to choose from, so we have $\binom{11}{5}$ choices for the green beads. Continuing in this way, there are $\binom{6}{4}$ position choices for the blue beads and there is $\binom{2}{2} = 1$ choice for the black beads. Hence, there are

$$\binom{20}{9,5,4,2} = \binom{20}{9} \binom{11}{5} \binom{6}{4} \binom{2}{2}$$

= 167,960 \cdot 462 \cdot 15 \cdot 1 = 1,163,962,800

possible arrangements of the 20 beads.

The next few sections are devoted to some probabilistic applications of the combinatorial techniques we have been discussing.

5.4 Sampling with replacement – The binomial distribution toc

We will first consider probabilities associated with the number of objects of a specified type in a random sample selected with replacement.

We will use the selection of balls from a box of suitably labeled balls to represent a random experiment. Consider a box (the population) containing $N = N_1 + N_2$ balls (individual objects) of which N_1 are red (of one type) and $N_2 = N - N_1$ are blue (of a second type). Suppose that a simple random sample of n balls is selected at random with replacement from this population. For an integer x, between 0 and n, we will find the probability P(x) that the sample contains exactly x red balls (and consequently exactly n - x blue balls).

At first glance the use of a box of balls as a model for a population and the drawing of balls from the box as sampling from the population may not appear to lend itself to great generality in application. As we shall see, this is not the case. The simple device of using the selection of balls from a box — a so-called box model — can be adapted to a very wide spectrum of applications.

Figure 5.2 A box containing 5 red and 15 blue balls.

	$N_1 = 5$ balls of the first type (red) $N_2 = 15$ balls of the second type (blue)
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Let's start with a simple example. Consider the box (population), illustrated in figure 5.2, containing N = 20 balls (objects) of which $N_1 = 5$ are red (of one type) and $N_2 = 15$ are blue (of a second type). Suppose that a simple random sample of n = 3 balls is selected at random with replacement from this population. The characteristic of interest here is the number of red balls among the n = 3 balls in the sample. The possible numbers of red balls in a sample of n = 3 are 0, 1, 2, 3.

Before we start counting, consider a representation of the possible outcomes of the three draws by ordered triples of the letters R for red and B for blue. Note well that these 8 elementary outcomes are not equally likely!

 $\Omega = \{BBB, RBB, BRB, BBR, RRB, RBR, BRR, RRR\}$

Since we are drawing with replacement, there are always 5 choices for each R and 15 choices for each B in these outcomes. Furthermore, there are always a total of 20 choices for each draw. Thus, the probabilities for these outcomes are as shown below.

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$$P(BBB) = \frac{15 \cdot 15 \cdot 15}{20 \cdot 20 \cdot 20} = \left(\frac{15}{20}\right)^{3}$$

$$P(RBB) = \frac{5 \cdot 15 \cdot 15}{20 \cdot 20 \cdot 20} = \left(\frac{5}{20}\right)^{1} \left(\frac{15}{20}\right)^{2}$$

$$P(BRB) = \frac{15 \cdot 5 \cdot 15}{20 \cdot 20 \cdot 20} = \left(\frac{5}{20}\right)^{1} \left(\frac{15}{20}\right)^{2}$$

$$P(BBR) = \frac{15 \cdot 15 \cdot 5}{20 \cdot 20 \cdot 20} = \left(\frac{5}{20}\right)^{1} \left(\frac{15}{20}\right)^{2}$$

$$P(RRB) = \frac{5 \cdot 5 \cdot 15}{20 \cdot 20 \cdot 20} = \left(\frac{5}{20}\right)^{2} \left(\frac{15}{20}\right)^{1}$$

$$P(RBR) = \frac{5 \cdot 15 \cdot 5}{20 \cdot 20 \cdot 20} = \left(\frac{5}{20}\right)^{2} \left(\frac{15}{20}\right)^{1}$$

$$P(BRR) = \frac{15 \cdot 5 \cdot 5}{20 \cdot 20 \cdot 20} = \left(\frac{5}{20}\right)^{2} \left(\frac{15}{20}\right)^{1}$$

$$P(RRR) = \frac{15 \cdot 5 \cdot 5}{20 \cdot 20 \cdot 20} = \left(\frac{5}{20}\right)^{2} \left(\frac{15}{20}\right)^{1}$$

$$P(RRR) = \frac{5 \cdot 5 \cdot 5}{20 \cdot 20 \cdot 20} = \left(\frac{5}{20}\right)^{3}$$

Notice that there is $\binom{3}{0} = 1$ way to get x = 0 red (BBB); there are $\binom{3}{1} = 3$ ways to get x = 1 red (RBB, BRB, BBR); there are $\binom{3}{2} = 3$ ways to get x = 2 red (RRB, RBR, BRR); and, there is $\binom{3}{3} = 1$ way to get x = 3 red (RRR). Also notice that for x = 1 red and x = 2 red, the three probabilities corresponding to the three ways to order the R's and B's are the same. These observations lead to the probabilities P(x) as summarized below.

These probabilities depend on $N_1 = 5$ and $N_2 = 15$ only through the proportions $p_1 = \frac{5}{20} = \frac{1}{4}$ (proportion red) and $p_2 = 1 - p_1 = \frac{15}{20} = \frac{3}{4}$ (proportion blue). This implies that if we had used a box with any values of N_1 and N_2 that yielded these proportions, then the probabilities would be the same!

$$P(0) = P(0 \text{ red}) = 1 \cdot \frac{15 \cdot 15 \cdot 15}{20 \cdot 20 \cdot 20} = {\binom{3}{0}} \left(\frac{5}{20}\right)^0 \left(\frac{15}{20}\right)^3 = {\binom{3}{0}} \left(\frac{1}{4}\right)^0 \left(\frac{3}{4}\right)^3 = \frac{27}{64}$$

$$P(1) = P(1 \text{ red}) = 3 \cdot \frac{5 \cdot 15 \cdot 15}{20 \cdot 20 \cdot 20} = {\binom{3}{1}} \left(\frac{5}{20}\right)^1 \left(\frac{15}{20}\right)^2 = {\binom{3}{1}} \left(\frac{1}{4}\right)^1 \left(\frac{3}{4}\right)^2 = \frac{27}{64}$$

$$P(2) = P(2 \text{ red}) = 3 \cdot \frac{5 \cdot 5 \cdot 15}{20 \cdot 20 \cdot 20} = {\binom{3}{2}} \left(\frac{5}{20}\right)^2 \left(\frac{15}{20}\right)^1 = {\binom{3}{2}} \left(\frac{1}{4}\right)^2 \left(\frac{3}{4}\right)^1 = \frac{9}{64}$$

$$P(3) = P(3 \text{ red}) = 1 \cdot \frac{5 \cdot 5 \cdot 5}{20 \cdot 20 \cdot 20} = {\binom{3}{3}} \left(\frac{5}{20}\right)^3 \left(\frac{15}{20}\right)^0 = {\binom{3}{3}} \left(\frac{1}{4}\right)^3 \left(\frac{3}{4}\right)^0 = \frac{1}{64}$$

We are now ready to generalize the expression for P(x) from our specific example with $N_1 = 5$ red, $N_2 = 15$ blue, and a sample of size n = 3 to the general situation.

In general, we have a population containing $N = N_1 + N_2$ individual objects (the balls in the box) of which N_1 are of one type (red) and $N_2 = N - N_1$ are of a second type (blue). The proportion of individual objects of the first type (red) is $p_1 = N_1/N$ and the proportion of individual objects of the second type (blue) is $p_2 = N_2/N = 1 - p_1$. We are assuming that a simple random sample of n individual objects is selected at random with replacement from this population.

For x = 0, 1, ..., n, the probability that the sample of n objects contains exactly x objects of the first type (red) is

$$P(x) = \binom{n}{x} \left(\frac{N_1}{N}\right)^x \left(\frac{N_2}{N}\right)^{n-x}$$
$$= \binom{n}{x} p_1^x p_2^{n-x} = \binom{n}{x} p_1^x (1-p_1)^{n-x}.$$

Notice that this expression depends on N, N_1 , and N_2 only through $p_1 = \frac{N_1}{N}$, the probability that a single object (ball) selected at random is of the first type (red), and $1 - p_1 = \frac{N_2}{N} = \frac{N-N_1}{N}$, the probability that a single object (ball) selected at random is of the second type (blue). Hence, for fixed values of n and x, the probability P(x) does not depend on the size of the population; it only depends on the proportion of red balls in the population.

Let's review the terms in this expression for the binomial probability P(x). It is helpful to split this expression into two terms.

$$P(x) = \binom{n}{x} \left(p_1^x p_2^{n-x} \right) = \binom{n}{x} \left(\frac{N_1^x \cdot N_2^{n-x}}{N^x} \right)$$

(1) $\binom{n}{x}$ is counting the number of ways that x R's and n - x B's can be arranged in a sequence of length n. This is analogous to counting the number of ways to string x red beads and n - x blue beads like we did in Example 5.7a.

(2) $p_1^x p_2^{n-x} = \frac{N_1^x \cdot N_2^{n-x}}{N^x}$ is the probability that we observed x R's followed by n-x B's, expressed as the ratio N(A)/N of the number of favorable outcomes to the number of possible outcomes. As noted above, this probability is the same for each of the $\binom{n}{x}$ ways to order x R's and n-x B's.

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Expressing the probability P(x) as the product of terms (1) and (2) is equivalent to summing the probabilities (as expressed in term (2)) for each of the $\binom{n}{x}$ ways that x R's and n - x B's can be arranged in a sequence of length n.

We will reconsider this binomial distribution and discuss it in more detail in a later section.

Example 5.8 Tossing a die Tossing a fair (balanced) die and noting the number on the upturned face is abstractly the same as selecting a number (ball) at random from the set $\{1, 2, 3, 4, 5, 6\}$. Tossing the die *n* times is equivalent to selecting a random sample of size *n* with replacement from the set $\{1, 2, 3, 4, 5, 6\}$.

If a fair die is tossed 10 times, then, for x = 0, 1, ..., 10, the probability of observing exactly x aces (ones) is

$$P(x) = {\binom{10}{x}} \left(\frac{1}{6}\right)^x \left(\frac{5}{6}\right)^{10-x}$$

In particular:

The probability of observing exactly 0 aces is $P(0) = {\binom{10}{0}} (\frac{1}{6})^0 (\frac{5}{6})^{10} = \frac{1 \cdot 5^{10}}{6^{10}} \approx .1615$. The probability of observing exactly 1 ace is $P(1) = {\binom{10}{1}} (\frac{1}{6})^1 (\frac{5}{6})^9 = \frac{10 \cdot 5^9}{6^{10}} \approx .3230$. The probability of observing exactly 2 aces is $P(2) = {\binom{10}{2}} (\frac{1}{6})^2 (\frac{5}{6})^8 = \frac{45 \cdot 5^8}{6^{10}} \approx .2907$. The probability of observing exactly 3 aces is $P(3) = {\binom{10}{3}} (\frac{1}{6})^3 (\frac{5}{6})^7 = \frac{120 \cdot 5^7}{6^{10}} \approx .1550$.

Example 5.9 Sampling from a box of balls Consider a box containing 600 balls of which 100 are red and 500 are blue. Since p = 1/6 of the balls in the box are red and the probability of getting an ace when a fair die is tossed once is p = 1/6, selecting 10 balls at random with replacement from these 600 balls and counting the number of red balls is equivalent to tossing a fair die 10 times and counting the number of aces tossed. Thus the probabilities of Example 5.8 apply here as well. That is, if 10 balls are selected at random with replacement from these 600 balls, then, for x = 0, 1, ..., 10, the probability of observing exactly x red balls is

$$P(x) = {\binom{10}{x}} \left(\frac{1}{6}\right)^x \left(\frac{5}{6}\right)^{10-x}.$$

5.5 Sampling without replacement – The hypergeometric distribution toc We will now consider probabilities associated with the number of objects of a specified type in a random sample selected without replacement.

As before, we will use the selection of balls from a box of suitably labeled balls to represent a random experiment and we will consider a box (the population) containing $N = N_1 + N_2$ balls (individual objects) of which N_1 are red (of one type) and $N_2 = N - N_1$ are blue (of a second type).

This time, however, suppose that a simple random sample of n balls is selected at random without replacement from this population. For an integer x, between 0 and n, we will find the probability P(x) that the sample contains exactly x red balls (and consequently exactly n - x blue balls). Note that, since the population contains N_1 red balls and $N_2 = N - N_1$ blue balls and we are sampling without replacement certain values of xmay not be attainable. In particular, we must have P(x) = 0 when $x > N_1$ and when $n - x > N - N_1$, since these values of x are not possible.

Let's start with the same simple example as we used when introducing the binomial distribution. Consider the box (population), illustrated in figure 5.2, containing N = 20balls (objects) of which $N_1 = 5$ are red (of one type) and $N_2 = 15$ are blue (of a second type). Suppose that a simple random sample of n = 3 balls is selected at random without replacement from this population. The characteristic of interest here is the number of red balls among the n = 3 balls in the sample. The possible numbers of red balls in a sample of n = 3 are 0, 1, 2, 3. In this application, since the box contains more than 3 balls of both colors, all values of x are attainable.

For the probabilities we want to compute, when sampling without replacement we can think of selecting all n balls at once and we can use unordered outcomes in our representation of the sample space. There are $\binom{20}{3} = 1,140$ ways to choose 3 balls (without keeping track of order) from a box containing 20 balls. In order to formally list these elementary outcomes we would need to add labels to the 20 balls so that they would be distinguishable. This is tedious and not very helpful so let's just consider the counting argument. Before we start counting, note that there are three basic types of elementary outcomes here, using R for red and B for blue, the possibilities are: a set of the form $\{B, B, B\}$ (three distinct blue balls); a set of the form $\{R, B, B\}$ (one red ball and two distinct blue balls); a set of the form $\{R, R, B\}$ (two distinct red balls and one blue ball); and, a set of the form $\{R, R, R\}$

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(three distinct red balls). Note well that, as when sampling with replacement, these 4 types of elementary outcomes are not equally likely!

It is helpful to think of the process of choosing the balls as series of three stages (select the desired number, x, of red balls, select the desired number, n - x, of blue balls, and select the desired number, n, of balls). These three selections are illustrated in Figure 5.3.

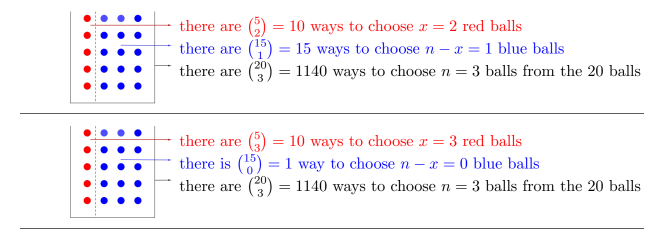
Figure 5.3 Choosing balls from a box containing 5 red and 15 blue balls.
choose x balls from the $N_1 = 5$ balls of the first type (red) choose $n - x$ balls from the $N_2 = 15$ balls of the second type (blue) choose n balls from the $N = N_1 + N_2 = 20$ balls in the box

There is $\binom{5}{0} = 1$ way to choose x = 0 red from 5 red balls; there are $\binom{5}{1} = 5$ ways to choose x = 1 red from 5 red balls; there are $\binom{5}{2} = 10$ ways to choose x = 2 red from 5 red balls; and, there are $\binom{5}{3} = 10$ ways to choose x = 3 red from 5 red balls. Similarly, there are $\binom{15}{3} = 455$ ways to choose n - x = 3 - 0 = 3 blue from 15 blue balls; there are $\binom{15}{2} = 105$ ways to choose n - x = 3 - 1 = 2 blue from 15 blue balls; there are $\binom{15}{1} = 15$ ways to choose n - x = 3 - 1 = 2 blue from 15 blue balls; there are $\binom{15}{1} = 15$ ways to choose n - x = 3 - 2 = 1 blue from 15 blue balls; and, there is $\binom{15}{0} = 1$ way to choose n - x = 3 - 3 = 0 blue from 15 blue balls. Finally, as noted earlier, there are $\binom{20}{3} = 1,140$ ways to choose n = 3 balls from the N = 20 balls in the box. These observations are represented graphically in Figure 5.4.

Figure 5.4 Choosing balls from a box containing 5 red and 15 blue balls.

there is $\binom{5}{0} = 1$ way to choose x = 0 red balls there are $\binom{15}{3} = 455$ ways to choose n - x = 3 blue balls there are $\binom{20}{3} = 1140$ ways to choose n = 3 balls from the 20 balls

there are $\binom{5}{1} = 5$ ways to choose x = 1 red balls there are $\binom{15}{2} = 105$ ways to choose n - x = 2 blue balls there are $\binom{20}{3} = 1140$ ways to choose n = 3 balls from the 20 balls



Combining these observations and computations yields the probabilities P(x) as summarized below.

$$P(0) = P(0 \text{ red}) = \frac{\binom{5}{0}\binom{15}{3}}{\binom{20}{5}} = \frac{1 \cdot 455}{1140} = \frac{91}{228}$$
$$P(1) = P(1 \text{ red}) = \frac{\binom{5}{1}\binom{15}{2}}{\binom{20}{5}} = \frac{5 \cdot 105}{1140} = \frac{105}{228}$$
$$P(2) = P(2 \text{ red}) = \frac{\binom{5}{2}\binom{15}{1}}{\binom{20}{5}} = \frac{10 \cdot 15}{1140} = \frac{30}{228}$$
$$P(3) = P(3 \text{ red}) = \frac{\binom{5}{3}\binom{15}{0}}{\binom{20}{5}} = \frac{10 \cdot 1}{1140} = \frac{2}{228}$$

We are now ready to generalize the expression for P(x) from our specific example with $N_1 = 5$ red, $N_2 = 15$ blue, and a sample of size n = 3 to the general situation.

In general, we have a population containing $N = N_1 + N_2$ individual objects (the balls in the box) of which N_1 are of one type (red) and $N_2 = N - N_1$ are of a second type (blue). We are assuming that a simple random sample of n individual objects is selected at random without replacement from this population.

For x = 0, 1, ..., n (subject to the restrictions described below) the probability that the sample of n objects contains exactly x objects of the first type (red) is

$$P(x) = \frac{\binom{N_1}{x}\binom{N-N_1}{n-x}}{\binom{N}{n}} = \frac{\binom{N_1}{x}\binom{N_2}{n-x}}{\binom{N_1+N_2}{n}}.$$

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The restriction mentioned above says that this hypergeometric probability formula only works when $x \leq N_1$ and $n-x \leq N_2$. This restriction reflects the fact that we are sampling without replacement and that it may be possible to run out of objects (balls) of one type (color) before we obtain a sample of the desired size n.

Notice the pattern in the binomial coefficients in the expression for the hypergeometric probability P(x). The "top" numbers, N_1 and N_2 , in the numerator binomial coefficients sum to give the "top" number, N, in the denominator binomial coefficient. Similarly, the "bottom" numbers, x and n - x, in the numerator binomial coefficients sum to give the "bottom" number, n, in the denominator binomial coefficient. This pattern reflects the facts that the N objects (balls) are comprised of N_1 objects of the first type (red balls) and N_2 objects of the second type (blue balls). And, the n objects (balls) in the sample are comprised of x objects of the first type (red balls) and n-x objects of the second type (blue balls).

Also notice that, unlike the binomial probability formula, this hypergeometric probability formula does depend on the values of N_1 and N_2 . Therefore, these hypergeometric probabilities cannot be computed unless we know the sizes of the two subpopulations.

We will reconsider this hypergeometric distribution and discuss it in more detail in a later section.

Example 5.10 Poker Consider the number of aces in a 5-card poker hand. Dealing a 5-card hand is abstractly the same as selecting five cards at random without replacement from a collection of 52 balls (cards). For this problem, the relevant partition of the 52 cards in the deck is into the 4 aces and the 48 non-aces. Since there are only 4 aces the x values of interest are x = 0, 1, 2, 3, 4. For x = 0, 1, 2, 3, 4, the probability that the 5-card hand contains exactly x aces is

$$P(x) = \frac{\binom{4}{x}\binom{48}{5-x}}{\binom{52}{5}}.$$

In particular:

The probability that the hand contains exactly 0 aces is $P(0) = [\binom{4}{0}\binom{48}{5}]/\binom{52}{5} \approx .6588$ The probability that the hand contains exactly 1 ace is $P(1) = [\binom{4}{1}\binom{48}{4}]/\binom{52}{5} \approx .2995$ The probability that the hand contains exactly 2 aces is $P(2) = [\binom{4}{2}\binom{48}{3}]/\binom{52}{5} \approx .0399$ The probability that the hand contains exactly 3 aces is $P(3) = [\binom{4}{3}\binom{48}{2}]/\binom{52}{5} \approx .0017$ The probability that the hand contains exactly 4 aces is $P(4) = [\binom{4}{4}\binom{48}{1}]/\binom{52}{5} \approx .00002$

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We will now extend the binomial distribution by allowing three types of objects. As before, suppose that a simple random sample of n balls (objects) is selected at random with replacement from a population of N balls (objects). However, now suppose that N_1 balls are red (of one type), N_2 are blue (of a second type), and N_3 are magenta (of a third type), with $N = N_1 + N_2 + N_3$.

For $n = x_1 + x_2 + x_3$, we will find the probability $P(x_1, x_2, x_3)$ that the sample contains x_1 red balls, x_2 blue balls, and x_3 magenta balls. As in the binomial case, we will think of obtaining the balls in the sample one at a time and use ordered outcomes in our representation of the sample space.

As we did with the binomial distribution, let's start with a simple example. Consider the box (population), illustrated in Figure 5.5, containing N = 30 balls (objects) of which $N_1 = 5$ are red (of one type), $N_2 = 15$ are blue (of a second type), and $N_3 = 10$ are magenta (of a third type). Suppose that a simple random sample of n = 6 balls is selected at random with replacement from this population. The characteristics of interest here are the number of balls of each of the three colors (objects of each of the three types) among the n = 6 balls in the sample. There are 28 ways in which a sample of n = 6 balls can be distributed among the three colors. For example, we might observe ($x_1 = 0$ red, $x_2 = 0$ blue, and $x_3 = 6$ magenta) or ($x_1 = 1$ red, $x_2 = 2$ blue, and $x_3 = 3$ magenta) or ($x_1 = 2$ red, $x_2 = 2$ blue, and $x_3 = 2$ magenta).

I Igure 0.0 II box com	taining o rea, 10 blue, and 10 magenta bans.
• • • • •	
•••••	$N_1 = 5$ balls of the first type (red)
• • • • •	$N_2 = 15$ balls of the second type (blue)
• • • • • •	$N_3 = 10$ balls of the third type (magenta)

Figure 5.5 A	box	containing 5	5 red.	, 15 blue	e, and 10	magenta balls.
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Before we start counting, consider a representation of the possible outcomes of the six draws by ordered 6-tuples of the letters R for red, B for blue, and M for magenta. Note

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well that these 28 elementary outcomes are not equally likely!

$\label{eq:sphere:sphe$

Since we are drawing with replacement, there are always 5 choices for each R, 15 choices for each B, and 10 choices for each M in these outcomes. Furthermore, there are always a total of 30 choices for each draw. The probabilities for a few of these outcomes are

$P(RRBBMM) = \frac{5 \cdot 5 \cdot 15 \cdot 15 \cdot 10 \cdot 10}{30 \cdot 30 \cdot 30 \cdot 30 \cdot 30 \cdot 30} = \left(\frac{5}{30}\right)^2 \left(\frac{15}{30}\right)^2 \left(\frac{10}{30}\right)^2$
$P(RBBMMM) = \frac{5 \cdot 15 \cdot 15 \cdot 10 \cdot 10 \cdot 10}{30 \cdot 30 \cdot 30 \cdot 30 \cdot 30 \cdot 30} = \left(\frac{5}{30}\right)^1 \left(\frac{15}{30}\right)^2 \left(\frac{10}{30}\right)^3$
$P(RRRBBB) = \frac{5 \cdot 5 \cdot 5 \cdot 15 \cdot 15 \cdot 15}{30 \cdot 30 \cdot 30 \cdot 30 \cdot 30 \cdot 30} = \left(\frac{5}{30}\right)^3 \left(\frac{15}{30}\right)^3 \left(\frac{10}{30}\right)^0$
$P(RRRRR) = \frac{5 \cdot 5 \cdot 5 \cdot 5 \cdot 5 \cdot 5}{30 \cdot 30 \cdot 30 \cdot 30 \cdot 30 \cdot 30} = \left(\frac{5}{30}\right)^6 \left(\frac{15}{30}\right)^0 \left(\frac{10}{30}\right)^0$

For each collection of values x_1, x_2, x_3 with $x_1 + x_2 + x_3 = 6$, we now need to determine how many ways we can order a collection of $x_1 R's$, $x_2 B's$, and $x_3 M's$. As we noted in the binomial section, this is analogous to asking in how many ways can we string a collection of x_1 red beads, x_2 blue beads, and x_3 magenta beads (when $x_1 + x_2 + x_3 = 6$). In other words, we need count the number of ways to partition the n = 6 draws (elements of the 6-tuple) into a group of x_1 when a red ball is drawn, a group of x_2 when a blue ball is drawn, and a group of x_3 when a magenta ball is drawn. The solution is provided by the multinomial coefficient

$$\binom{6}{x_1, x_2, x_3} = \binom{6}{x_1} \binom{6 - x_1}{x_2} \binom{6 - x_1 - x_2}{x_3}.$$

As in the binomial case, we can now combine these observations and provide the following expression for these multinomial probabilities in the application when n = 6, $p_1 = N_1/N = 5/30$, $p_2 = N_2/N = 15/30$, and $p_3 = N_3/N = 10/30$. For values of x_1, x_2, x_3 between 0

and 6 with $x_1 + x_2 + x_3 = 6$, the probability of observing x_1 red balls, x_2 blue balls, and x_3 magenta balls in a sample of size n = 6 is

$$P(x_1, x_2, x_3) = \begin{pmatrix} 6\\x_1, x_2, x_3 \end{pmatrix} \left(\frac{5}{30}\right)^{x_1} \left(\frac{15}{30}\right)^{x_2} \left(\frac{10}{30}\right)^{x_3} \\ = \begin{pmatrix} 6\\x_1 \end{pmatrix} \begin{pmatrix} 6-x_1\\x_2 \end{pmatrix} \begin{pmatrix} 6-x_1-x_2\\x_3 \end{pmatrix} \left(\frac{5}{30}\right)^{x_1} \left(\frac{15}{30}\right)^{x_2} \left(\frac{10}{30}\right)^{x_3}$$

Generalizing this expression for $P(x_1, x_2, x_3)$ from our specific example with $N_1 = 5$ red, $N_2 = 15$ blue, $N_3 = 10$ magenta, and a sample of size n = 6 to the general situation is straightforward.

In general, we have a population containing $N = N_1 + N_2 + N_3$ individual objects (the balls in the box) of which N_1 are of one type (red) and N_2 are of a second type (blue), and N_3 are of the third type (magenta). The proportion of individual objects of the first type (red) is $p_1 = N_1/N$, the proportion of individual objects of the second type (blue) is $p_2 = N_2/N$, and the proportion of individual objects of the third type (magenta) is $p_3 = N_3/N$. Note well that $p_1 + p_2 + p_3 = 1$, since $N_1 + N_2 + N_3 = N$. We are assuming that a simple random sample of n individual objects is selected at random with replacement from this population.

For values of x_1, x_2, x_3 between 0 and n with $x_1 + x_2 + x_3 = n$, the probability of observing x_1 objects of the first type (red balls), x_2 objects of the second type (blue balls), and x_3 objects of the third type (magenta balls) in a sample of size n is

$$P(x_1, x_2, x_3) = \binom{n}{x_1, x_2, x_3} \left(\frac{N_1}{N}\right)^{x_1} \left(\frac{N_2}{N}\right)^{x_2} \left(\frac{N_3}{N}\right)^{x_3} = \binom{n}{x_1} \binom{n - x_1}{x_2} \binom{n - x_1 - x_2}{x_3} \left(\frac{N_1}{N}\right)^{x_1} \left(\frac{N_2}{N}\right)^{x_2} \left(\frac{N_3}{N}\right)^{x_3} = \binom{n}{x_1} \binom{n - x_1}{x_2} \binom{n - x_1 - x_2}{x_3} p_1^{x_1} p_2^{x_2} p_3^{x_3}$$

Notice that this expression depends on N, N_1, N_2 and N_3 only through $p_1 = \frac{N_1}{N}$, the probability that a single object (ball) selected at random is of the first type (red), $p_2 = \frac{N_2}{N}$, the probability that a single object (ball) selected at random is of the second type (blue), and $p_3 = \frac{N_3}{N}$, the probability that a single object (ball) selected at random is of the third type (magenta). Hence, for fixed values of n, x_1, x_2 , and x_3 , the probability $P(x_1, x_2, x_3)$ does not depend on the size of the population; it only depends on the proportions of

objects (balls) in the three subpopulations. Computational aside: $n - x_1 - x_2 = x_3$ and thus $\binom{n-x_1-x_2}{x_3} = \binom{x_3}{x_3} = 1$.

As you might expect, this is a trinomial distribution, since we have three types of objects (colors). The extension of this expression to give multinomial probabilities when there are four or more types of objects should be reasonably clear. We will reconsider this multinomial distribution and discuss it in more detail in a later section.

Example 5.11 Tossing a die Suppose a fair die is tossed 10 times. Since the population (possible outcomes for one toss) $\{1, 2, 3, 4, 5, 6\}$ contains six values, in order to apply the multinomial probability formula, we need to partition these six values into a fixed number of values ("colors").

First consider the probability of observing exactly 3 ones and exactly 4 twos. Since 3+4=7 we also need 3 numbers other than one or two. Thus we need the partition $\{1, 2, 3, 4, 5, 6\} = \{1\} \cup \{2\} \cup \{3, 4, 5, 6\}$. Here we have $N_1 = 1$, $N_2 = 1$ and $N_3 = 4$ and $x_1 = 3$, $x_2 = 4$, and $x_3 = 3$. The probability of observing exactly 3 ones and exactly 4 twos is

$$P(3 \text{ ones and } 4 \text{ twos}) = P(3 \text{ ones, } 4 \text{ twos, and } 3 \text{ others})$$
$$= {\binom{10}{3}} {\binom{7}{4}} {\binom{3}{3}} {\binom{1}{6}}^3 {\binom{1}{6}}^4 {\binom{4}{6}}^3 = \frac{120 \cdot 35 \cdot 1 \cdot 4^3}{6^{10}} \approx .0044.$$

Now let's consider an event which involves four values. In particular, let's find the probability of observing exactly 2 ones, exactly 3 twos, and exactly 4 threes. Since 2+3+4=9we also need 1 number other than one, two, or three. Here we have $N_1 = 1$, $N_2 = 1$, $N_3 = 1$, and $N_4 = 3$ and $x_1 = 2$, $x_2 = 3$, $x_3 = 4$, and $x_4 = 1$. The probability of observing exactly 2 ones, exactly 3 twos, and exactly 4 threes is

$$P(2 \text{ ones, } 3 \text{ twos, and } 4 \text{ threes}) = P(2 \text{ ones, } 3 \text{ twos, } 4 \text{ threes, and } 1 \text{ other})$$
$$= {\binom{10}{2}} {\binom{8}{3}} {\binom{5}{4}} {\binom{1}{1}} {\binom{1}{6}}^2 {\binom{1}{6}}^3 {\binom{1}{6}}^4 {\binom{3}{6}}^1$$
$$= \frac{45 \cdot 56 \cdot 5 \cdot 1 \cdot 3^1}{6^{10}} \approx .00062.$$

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The hypergeometric distribution is also readily generalized to allow for balls of three or more colors. As before, suppose that a simple random sample of n balls (objects) is selected at random without replacement from a box (population) of N balls (objects). However, now suppose that N_1 balls are red (of one type), N_2 are blue (of a second type), and N_3 are magenta (of a third type), with $N = N_1 + N_2 + N_3$.

As we did with the hypergeometric distribution, let's start with a simple example. Consider the box (population), illustrated in Figure 5.6, containing N = 30 balls (objects) of which $N_1 = 5$ are red (of one type), $N_2 = 15$ are blue (of a second type), and $N_3 = 10$ are magenta (of a third type). Suppose that a simple random sample of n = 6 balls is selected at random without replacement from this population.

Figure 5.6 Choosing balls from a box containing 5 red and 15 blue balls.

••••	choose x_1 balls from the $N_1 = 5$ balls of the first type (red)
	\rightarrow choose x_2 balls from the $N_2 = 15$ balls of the second type (blue)
••••	- choose x_3 balls from the $N_3 = 10$ balls of the third type (magenta)
••••	choose $n = 6$ balls from the $N = N_1 + N_2 + N_3 = 30$
	balls in the box

In this application there are $\binom{30}{6} = 593,775$ equally likely possible outcomes (subsets of size n = 6). Our task is to determine how many of these possible outcomes are favorable for each event of the form "the sample contains x_1 red balls, x_2 blue balls, and x_3 magenta balls" so that we can compute the corresponding probability $P(x_1, x_2, x_3)$.

The ingredients we need are:

1. There are $\binom{5}{x_1}$ ways to choose x_1 red balls from the $N_1 = 5$ red balls in the box.

2. There are $\binom{15}{x_2}$ ways to choose x_2 blue balls from the $N_2 = 15$ blue balls in the box.

3. There are $\binom{10}{x_3}$ ways to choose x_3 magenta balls from the $N_3 = 10$ magenta balls in the box.

The obvious extension of the hypergeometric probability expression yields

$$P(x_1, x_2, x_3) = \frac{\binom{5}{x_1}\binom{15}{x_2}\binom{10}{x_3}}{\binom{30}{6}}$$

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Note well that the sizes of the subpopulations (numbers of balls of each color in the box) place some restrictions on which values of x_1 , x_2 , and x_3 , will work in this expression. Since there are only 5 red balls we must have $x_1 \in \{0, 1, 2, 3, 4, 5\}$. Since there are more than six blue balls and more than six magenta balls, there is no restriction on the values of x_2 and x_3 other than the requirement that $x_1 + x_2 + x_3 = 6$.

Generalizing this expression for $P(x_1, x_2, x_3)$ from our specific example with $N_1 = 5$ red, $N_2 = 15$ blue, $N_3 = 10$ magenta, and a sample of size n = 6 to the general situation is straightforward.

In general, we have a population containing $N = N_1 + N_2 + N_3$ individual objects (the balls in the box) of which N_1 are of one type (red) and N_2 are of a second type (blue), and N_3 are of the third type (magenta). We are assuming that a simple random sample of *n* individual objects is selected at random without replacement from this population.

For values of x_1, x_2, x_3 between 0 and n with $x_1 + x_2 + x_3 = n$ (subject to the restrictions that $x_1 \leq N_1, x_2 \leq N_2$, and $x_3 \leq N_3$), the probability of observing x_1 objects of the first type (red balls), x_2 objects of the second type (blue balls), and x_3 objects of the third type (magenta balls) in a sample of size n is

$$P(x_1, x_2, x_3) = \frac{\binom{N_1}{x_1}\binom{N_2}{x_2}\binom{N_3}{x_3}}{\binom{N_1 + N_2 + N_3}{x_1 + x_2 + x_3}} = \frac{\binom{N_1}{x_1}\binom{N_2}{x_2}\binom{N_3}{x_3}}{\binom{N}{n}}$$

As with the hypergeometric distribution, notice the pattern in the binomial coefficients in the expression for the multiple hypergeometric probability $P(x_1, x_2, x_3)$. The "top" numbers, N_1, N_2 , and N_3 , in the numerator binomial coefficients sum to give the "top" number, N, in the denominator binomial coefficient. Similarly, the "bottom" numbers, x_1, x_2 , and x_3 , in the numerator binomial coefficients sum to give the "bottom" number, n, in the denominator binomial coefficient. This pattern reflects the facts that the N objects (balls) are comprised of N_1 objects of the first type (red balls), N_2 objects of the second type (blue balls), and N_3 objects or the third type (magenta balls). And, the n objects (balls) in the sample are comprised of x_1 objects of the first type (red balls), x_2 objects of the second type (blue balls) and x_3 objects of the third type (magenta balls). Also notice that, unlike the multinomial probability formula, this multiple hypergeometric probability formula does depend on the values of N_1, N_2 , and N_3 . Therefore, as with hypergeometric 5.7 Sampling without replacement – The multiple hypergeometric distribution 97

probabilities, these multiple hypergeometric probabilities cannot be computed unless we know the sizes of the three subpopulations.

The extension to a population with four or more types of objects (balls) should be clear.

Example 5.12 Poker Now we can find the probabilities of some more interesting poker hands. The 52 card deck contains 4 aces, 4 kings, and 44 other cards. Using this partition yields the following probabilities.

The probability that a poker hand contains exactly 2 aces and exactly 1 king is

$$\frac{\binom{4}{2}\binom{4}{1}\binom{44}{2}}{\binom{52}{5}} \approx .0087.$$

The probability that a poker hand contains exactly 2 aces and exactly 2 kings is

$$\frac{\binom{4}{2}\binom{4}{2}\binom{44}{1}}{\binom{52}{5}} \approx .0006.$$

The probability that a poker hand contains exactly 2 aces and exactly 3 kings is

$$\frac{\binom{4}{2}\binom{4}{3}\binom{44}{0}}{\binom{52}{5}} \approx .000009.$$

5.8 Problems

The problem sets and solutions can be found on the book web page at https://www.ucs.louisiana.edu/~jcb0773/.

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6 Conditional probability and independence

6.1 Conditional probability

In many situations we have partial information about the outcome of an experiment and we need to update the probability measure to reflect this additional information. More formally, consider two events A and B which can occur at the same time, if we know that event A has occurred, then we need to determine how to update the probability of event B to take this information about A into account. That is, we need to determine the conditional probability of event B conditioning on the fact that event A has occurred. A couple of simple examples will help motivate the formal definition of conditional probability given below.

Example 6.1 Selecting one card Suppose we select 1 card at random from a 52 card deck. The deck of 52 cards contains 4 kings. Thus, the probability that we will select a king is P(king) = 4/52 = 1/13.

Now suppose we know that the card selected is a face card. How does this partial information about the card affect the probability that it is a king? That is, what is the conditional probability that the card is a king given the information that it is a face card? Since there are 12 face cards (the 4 kings, 4 queens, and 4 jacks) and we know that one of these 12 cards has been selected at random, there is a 4 out of 12 probability that the card is a king. Thus, when we know that the card is a face card the probability that it is a king (conditioning on the fact that it is a face card) is P(king given the card is a face card) = 4/12 = 1/3.

Next suppose that we know that the card selected is a spade. How does this partial information about the card affect the probability that it is a king? That is, what is the conditional probability that the card is a king given the information that it is a spade? Since there are 13 spades and we know that one of these 13 spades has been selected at random, and there is only one king of spades, there is a 1 out of 13 probability that the card is a king (conditioning on the fact that it is a spade) is P(king given the card is a spade) = 1/13.

It is helpful to consider the three probabilities we just computed in terms of the representation of the 52 cards in the image of Figure 3.1. The first probability we computed, the

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unconditional probability that the card is a king, is 1/13 because we are selecting at random from all 52 cards and 4 of these are kings. For the second probability, we conditioned on the event that the card was a face card. That is, we restricted our selection to the 12 cards in the last three columns of Figure 3.1. Since 1/3 of the cards in these columns are kings, we argued that the conditional probability of selecting a king given that the card is a face card is 1/3. For the third probability, we conditioned on the event that the card was a spade. That is, we restricted our selection to the 13 spades in the third row of Figure 3.1. Since one of the cards in the spade row is a king, we argued that the conditional probability of selecting a king given that the card is a spade is 1/13.

Notice that the unconditional probability of selecting a king changes when we condition on the event that the card is a face card; but, it does not change when we condition on the event that the card is a spade. When two events, say A and B, have the property that the unconditional probability of A is the same as the conditional probability of A given that B has occurred, the events are said to be independent. We will discuss independence in Section 6.2

Example 6.2 Bachelor's degree majors In 2004–2005, degree–granting institutions in the US conferred N = 1,439,264 bachelor's degrees (see Table 6.1). Of these degrees, N(S) = 174,980 had a Science or Math major. Thus, if we select a bachelor's degree recipient at random, then the probability that that person is a Science or Math major is P(Science or Math major) = N(S)/N = .1216. As you might expect this probability will change if we know that the person selected is a woman. For these N degrees, N(F) = 826,264 were awarded to women and out of these N(F) degrees $N(S \cap F) = 77,380$ had a Science or Math major. Thus, if we know that a woman was selected, then the conditional probability that she has a Science or Math major is

 $P(\text{Science or Math major given the person is a woman}) = N(S \cap F)/N(F) = .0936$. We will return to this example after we define some relevant notation. (source: National Center for Education Statistics (NCES))

Consider a population of N objects and two subpopulations (events) A and B. Let N(A) denote the number of objects in subpopulation A and let N(B) denote the number of objects in subpopulation B. If an object is selected at random from this population, then P(A) = N(A)/N and P(B) = N(B)/N. Now suppose that it is known that event A has

occurred. That is suppose that we know that the object belongs to subpopulation (event) A. Using subpopulation A as our reference space (new sample space) we note that the event B occurs if and only if the object is the intersection $A \cap B$. Thus letting $N(A \cap B)$ denote the number of objects in the population which belong to subpopulation A and subpopulation B, we see that the conditional probability of the selected object belonging to subpopulation A is

$$P(B|A) = \frac{N(A \cap B)}{N(A)} = \frac{N(A \cap B)/N}{N(A)/N} = \frac{P(A \cap B)}{P(A)}$$

Definition – Conditional probability. Given events A and B with P(A) > 0, the conditional probability of B given A is

$$P(B|A) = \frac{P(A \cap B)}{P(A)}$$

Example 6.2 Bachelor's degree majors revisited The N = 1,439,264 bachelor's degrees conferred by degree–granting institutions in the US in 2004–2005 are classified by field of major and sex in Table 6.1.

	se		
Major	male	female	total
Business	162,669	176,358	339,027
Social Sciences	91,533	220,410	311,943
Humanities	80,163	126,431	206,594
Science/Math	97,600	77,380	174,980
Art	61,901	101,006	162,907
Education	34,215	94,124	128,339
Engineering	84,919	30,555	115,474
total	613,000	826,264	1,439,264

Table 6.1. Bachelor's degrees conferred in 2004-05, by sex and major. (cou	Table 6.1.	Bachelor's degrees	s conferred in	2004-05, b	ov sex and ma	jor. (counts)
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Suppose that a person is selected at random from this group of N = 1,439,264 bachelor's degree recipients. The (unconditional) probabilities for each major-sex combination,

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obtained by dividing the entries in Table 6.1 by N, are given in Table 6.2. The margins of this table contain the (unconditional) probabilities for each major (the column of row probabilities) and for each sex (the row of column probabilities).

	S		
Major	male	female	total
Business	0.1130	0.1225	0.2356
Social Sciences	0.0636	0.1531	0.2167
Humanities	0.0557	0.0878	0.1435
Science/Math	0.0678	0.0538	0.1216
Art	0.0430	0.0702	0.1132
Education	0.0238	0.0654	0.0892
Engineering	0.0590	0.0212	0.0802
total	0.4259	0.5741	1

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Table 6.2.	Bacholor's c	logroog (contorrod i	n 2007 05	by gov	ond.	maior	proportiong	1
	Datheror S t	ICEICCS (Johnerieu i	II <u>2004</u> -00.	UV 952	. anu	major, i	DIODOI 010119	,

Of these N bachelor's degrees, N(M) = 613,000 were awarded to men and N(F) = 826,264 were awarded to women. We will use conditional probabilities to explore some differences and similarities in the distributions of the tabulated majors among these groups of men and women. Note that we can use the counts in Table 6.1 or the probabilities (proportions) in Table 6.2 to compute conditional probabilities.

First consider education majors. Let E denote the event that the person selected had an education major. There are N(E) = 128,339 education majors and $P(E) = \frac{128,339}{1,439,264} = 0.0892$. Thus, there is a 8.92% chance of selecting an education major. Let F denote the event that the person selected is a woman, and let M denote the event that the person selected is a man. From Table 6.2 we see that P(F) = 0.5741 and P(M) = 0.4259. Thus, there is a 57.41% chance of selecting a woman and a 42.59% chance of selecting a man. Now consider how the probability of selecting an education major changes when we know the sex of the selected person. In order to compute the desired conditional probabilities we need the counts or probabilities of the intersections $E \cap F$ and $E \cap M$.

First consider education majors among women. In terms of the counts in the "Female" column of Table 6.1 we have $N(E \cap F) = 94,124$ and N(F) = 826,264. Thus

$$P(E|F) = \frac{N(E \cap F)}{N(F)} = \frac{94,124}{826,264} = 0.1139.$$

In terms of the probabilities in the "Female" column of Table 6.2 we have $P(E \cap F) = 0.0654$ and P(F) = 0.5741. Thus

$$P(E|F) = \frac{0.0654}{0.5741} = 0.1139.$$

Now consider education majors among men. In terms of the counts in the "Male" column of Table 6.1 we have $N(E \cap M) = 34,215$ and N(M) = 613,000. Thus

$$P(E|M) = \frac{N(E \cap M)}{N(M)} = \frac{34,215}{613,000} = 0.0558.$$

In terms of the probabilities in the "Male" column of Table 6.2 we have $P(E \cap M) = 00238$. and P(M) = 0.4259. Thus

$$P(E|F) = \frac{0.0238}{0.4259} = 0.0558.$$

As noted earlier, there is a 8.92% chance of selecting an education major. However, if we know that the person selected is a woman, then there is an 11.39% chance that we have selected an education major, and, if we know that the person selected is a man, then there is a 5.58% chance that we have selected an education major. As you probably expected, we see that the conditional probability of selecting an education major given that we have selected a woman, P(E|F) = 0.1139, is substantially larger than the conditional probability of selecting an education major given that we have selected a man, P(E|M) = 0.0558.

If we performed analogous computations for the other majors we could find the entire conditional distribution of majors among the 826,264 women and the entire conditional distribution of majors among the 613,000 men. All we need to do is extract the numbers (counts or proportions) in the "Female" column (respectively "Male" column) of Table 6.1 or 6.2 and normalize them by dividing each by their sum. These conditional distributions are given in Table 6.3.

Note in particular the probabilities for engineering majors. Let G denote the event that the person selected had an engineering major. The unconditional probability of selecting an

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engineering major is P(G) = 0.0802 and the conditional probabilities are P(G|F) = 0.0370and P(G|M) = 0.1385.

men		women		
major	probability	major	probability	
Business	0.2654	Business	0.2134	
SocialSciences	0.1493	SocialSciences	0.2668	
Humanities	0.1308	Humanities	0.1530	
Science/Math	0.1592	Science/Math	0.0936	
Art	0.1010	Art	0.1222	
Education	0.0558	Education	0.1139	
Engineering	0.1385	Engineering	0.0370	
total	1.0000	total	0.9999	

Table 6.3 The conditional distributions of majors for each sex.

Given events A and B, with P(A) > 0, the conditional probability of B given A is $P(B|A) = P(A \cap B)/P(A)$. Multiplying the conditional probability P(B|A) by P(A) yields the following **multiplication rule for probabilities**.

The multiplication rule. Given events A and B with P(A) > 0,

$$P(A \cap B) = P(A)P(B|A).$$

Note that if $P(A \cap B) > 0$, then P(A) > 0 and P(B) > 0 and we have $P(A \cap B) = P(A)P(B|A) = P(B)P(A|B)$.

The multiplication rule-three events. Given events A, B, C with $P(A \cap B \cap C) > 0$,

$$P(A \cap B \cap C) = P(A)P(B|A)P(C|A \cap B)$$

The multiplication rule-several events. Given events A_1, \ldots, A_n with $P(A_1 \cap A_2 \cap \cdots \cap A_n) > 0$,

$$P(A_1 \cap A_2 \cap \dots \cap A_n) = P(A_1)P(A_2|A_1)P(A_3|A_1 \cap A_2) \cdots P(A_n|A_1 \cap A_2 \cap \dots \cap A_{n-1})$$

The multiplication rule is especially useful for computing the probability of an intersection when the events involved occur sequentially in time.

6.2 Independence

Intuitively we say that the events A and B are **independent** (stochastically independent) when knowing that B has occurred has no effect on the probability of occurrence of A, *i.e.* when P(A) = P(A|B). For mathematical convenience the formal definition of independence is in terms of a product so that it does not depend on the existence of conditional probabilities.

Definition – **Independence.** The events A and B are said to be independent (stochastically independent) when $P(A \cap B) = P(A)P(B)$.

Example 6.1 Selecting one card, revisited Recall that we found that the probability of selecting a king and the conditional probability of selecting a king given that the card selected is a spade are both 1/13. Thus, the events K – the card is a king and S – the card is a spade are independent. Note also that $P(K \cap S) = 1/52 = (1/4)(1/13) = P(S)P(K)$.

Example 6.3 Tossing a die If a fair die is tossed once and we let $A = \{2, 4, 6\}$ denote the event that an even value occurs and $B = \{1, 2, 3, 4\}$ the event that the value is four or less, then $P(A) = \frac{1}{2}$, $P(B) = \frac{2}{3}$, and $P(A \cap B) = \frac{1}{3}$. Thus $P(A \cap B) = P(A)P(B)$ and A and B are independent. We can also verify the independence of A and B by noting that $P(A) = \frac{3}{6} = \frac{1}{2}$ and $P(A|B) = \frac{2}{4} = \frac{1}{2}$ so that P(A|B) = P(A).

If two events are disjoint (mutually exclusive), then they cannot occur at the same time; thus if A and B are mutually exclusive, then they cannot be independent unless at least one of them is the null event. That is, if A and B are mutually exclusive, with P(A) > 0and P(B) > 0, then $P(A \cap B) = 0$ cannot be equal to P(A)P(B).

In the next result we note some ways in which the independence of two events implies the independence of some associated pairs of events.

Some implications of independence. If the events A and B are independent, then the events A and B^c are independent, the events A^c and B are independent, and the events A^c and B^c are independent.

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An aside – independence in general and conditional independence

When more than two events are involved the definition of independence is somewhat more complicated. The reason for this complexity is that we need to be sure that all possible subsets of the events exhibit appropriate independence properties. We first define **independence** (**mutual independence**) for several events.

Definition – **Independence of several events.** The events A_1, \ldots, A_n are said to be independent (mutually independent) when $P(A_i \cap A_j) = P(A_i)P(A_j)$ for all pairs (i, j) with distinct elements $P(A_i \cap A_j \cap A_k) = P(A_i)P(A_j)P(A_k)$ for all triples (i, j, k) with distinct elements and so on for sets of four, five, ..., up to $P(A_i \cap \cdots \cap A_n) = P(A_1) \cdots P(A_n).$

This definition of mutual independence simply says that in order for the events A_1, \ldots, A_n to be independent, every possible subset of the A_i must satisfy the condition that the probability of the intersection of the events in the subset is equal to the product of the probabilities of the individual events which form the subset. In particular, this factorization of the probability of an intersection must hold for every possible pair of events, triple of events, and so on up to the collection of all n events.

There is a weaker type of independence, **pairwise independence**, which arises in some applications. As the name suggests pairwise independence only requires that the factorization property holds for pairs.

Definition – **Pairwise independence.** The events A_1, \ldots, A_n are said to be pairwise independent when

 $P(A_i \cap A_j) = P(A_i)P(A_j)$ for all pairs (i, j) with $i \neq j$.

Example 6.4 Independent events Let $P(\omega) = 1/8$ for $\omega \in \Omega = \{1, 2, 3, 4, 5, 6, 7, 8\}$, let $A = \{1, 2, 3, 4\}, B = \{1, 2, 5, 6\}$, and $C = \{1, 3, 5, 7\}$. Then $P(A) = P(B) = P(C) = \frac{1}{2}$, $P(A \cap B) = P(A \cap C) = P(B \cap C) = \frac{1}{4} = (\frac{1}{2})^2$, and $P(A \cap B \cap C) = \frac{1}{8} = (\frac{1}{2})^3$. Thus, in this example, the events A, B, and C are independent (mutually independent).

Example 6.5 Pairwise independent events that are not independent Let $P(\omega) = 1/8$ for $\omega \in \Omega = \{1, 2, 3, 4, 5, 6, 7, 8\}$, let $A = \{1, 2, 3, 4\}$, $B = \{1, 2, 5, 6\}$, and $C = \{1, 2, 7, 8\}$. Then $P(A) = P(B) = P(C) = \frac{1}{2}$ and $P(A \cap B) = P(A \cap C) = P(B \cap C) = \frac{1}{4} = (\frac{1}{2})^2$. But $P(A \cap B \cap C) = \frac{1}{4} \neq (\frac{1}{2})^3$. Thus, in this example, the events A, B, and C are pairwise independent but not mutually independent.

Definition – **Conditional independence.** Given events A, B, and C with

 $P(A \cap B \cap C) > 0$, the events A and B are said to be conditionally independent given the event C when $P(A \cap B|C) = P(A|C)P(B|C)$.

6.3 The law of total probability – Bayes' theorem toc

In some situations we may find it convenient to compute the probability of an event by first decomposing the event into disjoint subevents and then adding the probabilities of these subevents. For example, given events A and B, we can partition the sample space as $\Omega = B \cup B^c$ and the event A as $A = (A \cap B) \cup (A \cap B^c)$. Since B and B^c are disjoint, $A \cap B$ and $A \cap B^c$ are also disjoint. This gives the decomposition of the probability of Aas $P(A) = P(A \cap B) + P(A \cap B^c)$. This result is the simplest case of the following law of total probability.

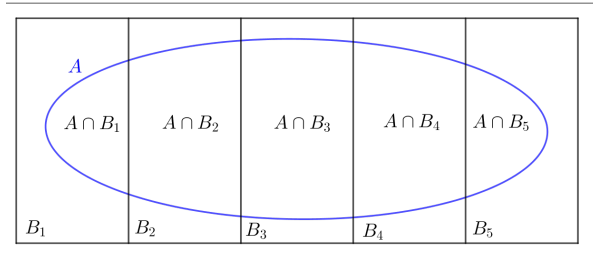
The law of total probability. Refer to Figure 6.1 for an illustration of this result. If the events B_1, \ldots, B_n form a partition of Ω , *i.e.* if B_1, \ldots, B_n are disjoint and $\Omega = B_1 \cup \cdots \cup B_n$, then, for any event A,

$$P(A) = P(A \cap B_1) + P(A \cap B_2) + \dots + P(A \cap B_n)$$
$$= \sum_{i=1}^n P(A \cap B_i).$$

Alternate statement of the law of total probability. If the events B_1, \ldots, B_n form a partition of Ω , and $P(B_i) > 0$ for $i = 1, \ldots, n$, then

$$P(A) = P(A|B_1)P(B_1) + P(A|B_2)P(B_2) + \dots + P(A|B_n)P(B_n)$$
$$= \sum_{i=1}^n P(A|B_i)P(B_i).$$

Figure 6.1 The law of total probability. The ellipse represents event A and the rectangular sections represent the events B_1, \ldots, B_5 of the partition.



Here we see that Ω is partitioned into 5 disjoint parts, A is similarly partitioned, and the probabilities of these part are added.

$$P(A) = P(A \cap B_1) + \dots + P(A \cap B_5)$$

 $P(A) = P(A|B_1)P(B_1) + \dots + P(A|B_5)P(B_5)$

We will now provide a variation of the law of total probability, called Bayes' theorem, which allows use to compute conditional probabilities of events at the first stage of an experiment given the outcome at the second stage.

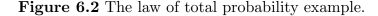
Bayes' theorem. If the events B_1, \ldots, B_n form a partition of Ω and $P(B_i) > 0$ for $i = 1, \ldots, n$, then for any event A with P(A) > 0, we have

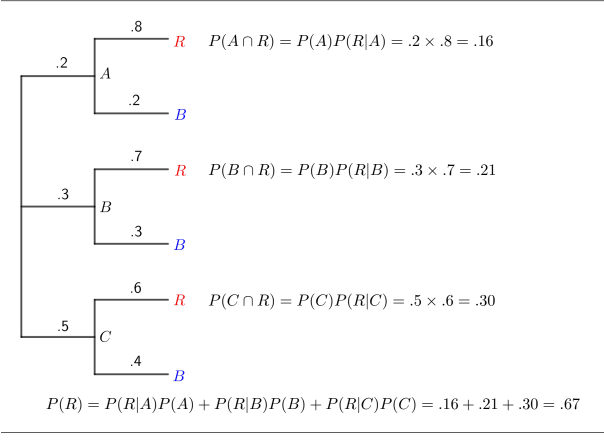
$$P(B_i|A) = \frac{P(A|B_i)P(B_i)}{P(A|B_1)P(B_1) + P(A|B_2)P(B_2) + \dots + P(A|B_n)P(B_n)}$$

Bayes' theorem is particularly useful for a situation where the occurrence of event A follows the occurrence of one of the events B_i in time and we are interested in the conditional probability that a particular B_i , say B_1 , has occurred given that event A has occurred. Note that if A and B are events and 0 < P(B) < 1, then the events B and B^c form a partition of Ω . Thus $P(A) = P(A \cap B) + P(A \cap B^c)$ and Bayes' theorem reduces to

$$P(B|A) = \frac{P(A \cap B)}{P(A \cap B) + P(A \cap B^c)} = \frac{P(A|B)P(B)}{P(A|B)P(B) + P(A|B^c)P(B^c)}$$

Example 6.6 Balls from a box Consider a box containing 100 balls of which 20 are labeled A, 30 are labeled B, and 50 are labeled C, and three other boxes labeled A, B, and C such that: box A contains 8 red and 2 blue balls; box B contains 7 red and 3 blue balls; and, box C contains 6 red and 4 blue balls. Now suppose that a ball is chosen at random from the box containing 100 balls, the letter (A, B, or C) on the ball is noted, and then a ball is chosen at random from the 10 balls in the box with the appropriate letter label. Clearly, the conditional probabilities of choosing a red ball given the letter label are: P(R|A) = .8, P(R|B) = .7, and P(R|C) = .6. It is also clear that the probabilities of selecting the label (A, B, or C) are: P(A) = .2, P(B) = .3, and P(C) = .5. Thus, $P(A \cap R) = P(R|A)P(A) = .16$, $P(B \cap R) = P(R|B)P(B) = .21$, and $P(C \cap R) = P(R|C)P(C) = .30$ are obtained by multiplication of the probabilities along the appropriate path in the tree diagram as indicated in Figure 6.2. The law of total probability says that the probability of choosing a red ball P(R) is obtained, as indicated at the bottom of the figure, by summing these three probabilities.





The values of conditional probabilities of the form P(A|R), the conditional probability that the ball was selected from box A given that it was red, are less obvious than the conditional probabilities of selecting red from a specified box. However, these conditional probabilities are readily computed using Bayes' Theorem. Here

$$P(R) = P(R|A)P(A) + P(R|B)P(B) + P(R|C)P(C) = .16 + .21 + .30 = .67$$

and we have

$$\begin{split} P(A|R) &= \frac{P(R \cap A)}{P(R)} = \frac{P(R|A)P(A)}{P(R)} = \frac{16}{67} \approx .24\\ P(B|R) &= \frac{P(R \cap B)}{P(R)} = \frac{P(R|B)P(B)}{P(R)} = \frac{21}{67} \approx .31\\ P(C|R) &= \frac{P(R \cap C)}{P(R)} = \frac{P(R|C)P(C)}{P(R)} = \frac{30}{67} \approx .45 \end{split}$$

Notice that each of these conditional probabilities is of the form $\frac{a}{a+b+c}$, where a, b, and c are the three probabilities summed to get P(R) in the tree diagram of Figure 6.2.

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It is interesting to compare the unconditional probabilities of drawing from boxes A, B, and C, P(A) = .2, P(B) = .3, and P(C) = .5, to the corresponding conditional probabilities given that the ball drawn is known to be red, $P(A|R) = \frac{16}{67} \approx .24$, $P(B|R) = \frac{21}{67} \approx .31$, and $P(C|R) = \frac{30}{67} \approx .45$. The initial probabilities (before we obtain the additional information that the ball drawn was red) are known as **prior probabilities** and the updated probabilities (conditional on the added information) are known as **posterior probabilities**. In this example, the conditional probability of drawing a red ball is highest when drawing from box A and lowest when drawing from box C. Thus, the added information that the ball was red increases the likelihood that the ball came from box A and decreases the likelihood that the ball came from box A and decreases the likelihood that the ball came from box C.

6.4 Problems

The problem sets and solutions can be found on the book web page at https://www.ucs.louisiana.edu/~jcb0773/.

$112\quad 6.3$ The law of total probability – Bayes' theorem

7 Discrete random variables

7.1 Random variables

In some of the examples we have considered the sample space is a set of integers, *e.g.*, $\Omega = \{1, 2, 3, 4, 5, 6\}$ corresponding to one toss of a die. In other examples we restricted our attention to events described in terms of a numerical value, *e.g.*, the number of heads in *n* tosses of a coin. We will now consider a more formal treatment of such assignments of numerical values to the outcomes of an experiment.

A function which assigns numerical values (real numbers) to the elements of a sample space is known as a **random variable** (denoted **r.v.**). Given an experiment with sample space Ω , a random variable associates a numerical value with each elementary outcome (element) ω of the sample space Ω . In the term random variable: the word variable indicates that the values of the function are numbers which are assigned to elementary outcomes; and, the adjective random is used as it is used in random experiment to indicate that the value of the random variable or outcome of the experiment is not known with certainty before the experiment is conducted at which time the value of the random variable or outcome of the random variable or outcome.

Given a random experiment with sample space Ω and a random variable X defined on Ω , the r.v. X defines a new sample space Ω_X comprised of all of the possible values of X. An event defined in terms of X (a subset of Ω_X) can be identified with the equivalent event from the underlying experiment (a subset of Ω). For example, the event X = 2corresponds to the event consisting of all elements ω of the underlying sample space Ω which are assigned the value X = 2. More formally, the event [X = x] (a subset of Ω_X) corresponds to the event { $\omega \in \Omega : X(\omega) = x$ } (a subset of Ω). In this last expression we are using the function notation $X(\omega)$ to indicate the value of X assigned to the elementary outcome ω .

Recall that a set is said to be discrete if it contains a finite or countably infinite number of elements. That is, the elements of a discrete set can be arranged in a list of the form x_1, x_2, \ldots, x_N , if the set is finite, or x_1, x_2, \ldots if the set is infinite. If Ω_X is discrete, then X is said to be a **discrete random variable**. In this chapter we will restrict our attention to discrete random variables.

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Probabilities of events defined in terms of the random variable X are determined by finding the probability of the equivalent event in terms of the underlying experiment. That is, given a random variable X and an event A (a subset of Ω_X), the probability that X takes on a value in A, $P(X \in A)$, is found by adding the probabilities of each of the elementary outcomes in Ω for which the corresponding X values are in A.

We will use a simple example to clarify the relationship between Ω and Ω_X , the relationship between events defined in terms of X and events of the original experiment, and the computation of probabilities of events defined in terms of X.

Example 7.1 Tossing a coin. Suppose we toss a coin three times. Letting H denote "heads", T denote "tails", and using 3-tuples of the letters H and T to represent the elementary outcomes, the sample space of the experiment is

$\Omega = \{HHH, HHT, HTH, HTT, THH, THT, TTH, TTT\}.$

Now let X denote the random variable "the number of heads in the three tosses". The sample space for this X is

$$\Omega_X = \{0, 1, 2, 3\}.$$

As noted above, an event defined in terms of X is a subset of Ω_X and each such event can be identified with an equivalent event from the underlying experiment. For example:

The event [X = 0] is equivalent to the event $\{TTT\}$;

The event [X = 1] is equivalent to the event $\{HTT, THT, TTH\};$

The event [X = 2] is equivalent to the event $\{HHT, HTH, THH\}$;

The event [X = 3] is equivalent to the event $\{HHH\}$; and,

The event $[X \leq 2]$ is equivalent to the event $\{TTT, HTT, THT, TTH, HHT, HTH, THH\}$.

If we assume that the coin we are tossing is fair so that each of the eight elementary outcomes in Ω has probability 1/8, then

$$\begin{split} P(X = 0) &= P(\{TTT\}) = 1/8; \\ P(X = 1) &= P(\{HTT, THT, TTH\}) = 3/8; \\ P(X = 2) &= P(\{HHT, HTH, THH\}) = 3/8; \\ P(X = 3) &= P(\{HHH\}) = 1/8; \text{ and,} \\ P(X \leq 2) &= P(\{TTT, HTT, THT, TTH, HHT, HTH, THH\}) = 7/8. \end{split}$$

We can characterize the distribution of the discrete r.v. X by specifying a **probability** mass function (denoted p.m.f.) p_X . The probability mass function p_X assigns a probability to each potential value of the random variable X, *i.e.*,

$$p_X(x) = P(X = x).$$

The p.m.f. assigns positive probabilities to the elements of Ω_X and zero probabilities to values not in Ω_X . Furthermore, since all of the values of X which have positive probability of occurrence belong to Ω_X , the probabilities of the elements of Ω_X must sum to one. The requisite properties of a p.m.f. are provided in the following definition.

Definition: probability mass function. Given a discrete sample space $\Omega_X = \{x_1, \ldots, x_N\}$ any function p_X with the properties that:

(1) $p_X(x) > 0$ for all $x \in \{x_1, \dots, x_N\};$

- (2) $p_X(x) = 0$ for all $x \notin \{x_1, \ldots, x_N\}$; and,
- (3) $p_X(x_1) + \dots + p_X(x_N) = 1$

can be viewed as the probability mass function (p.m.f.) of a random variable X with sample space Ω_X . The obvious modifications apply when $\Omega_X = \{x_1, x_2, \ldots\}$ is countably infinite.

For any event A defined in terms of the r.v. X, *i.e.*, for any $A \subset \Omega_X$, the probability of event A is equal to the sum of the probabilities of each of the distinct values of X which form the event (the sum of the $p_X(x)$ corresponding to each element $x \in A$). In symbols, the probability of A is

$$P(A) = \sum_{x \in A} p_X(x).$$

We will now return to the coin tossing example to clarify this definition of a probability mass function.

Example 7.1 Tossing a coin, revisited. As noted above, assuming that the coin we are tossing is fair so that each of the eight elementary outcomes in Ω has probability 1/8, P(X = 0) = 1/8, P(X = 1) = 3/8, P(X = 2) = 3/8, and P(X = 3) = 1/8. Hence, in function notation, X has probability mass function

$$p_X(x) = \begin{cases} 1/8 & \text{if } x = 0\\ 3/8 & \text{if } x = 1\\ 3/8 & \text{if } x = 2\\ 1/8 & \text{if } x = 3\\ 0 & \text{otherwise} \end{cases}$$

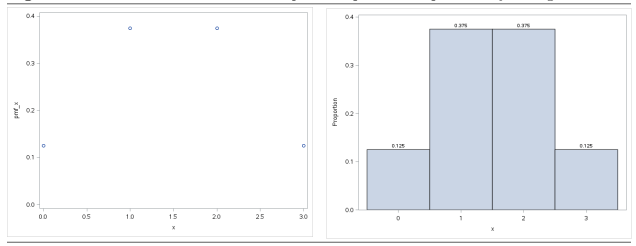
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We can also express this p.m.f. in tabular form as shown in Table 7.1.

Table	7.1 The p.m.f. of $X =$ the number of heads in 3 tosses of a fair coin.
x	$p_X(x)$
0	1/8
1	3/8
2	3/8
3	1/8

Note that observing the value of X is equivalent to selecting a ball at random from a box containing eight balls of which 1 is numbered zero, 3 are numbered one, 3 are numbered two, and 1 is numbered three. This is an example of a binomial distribution. This binomial p.m.f. is shown in the graph on the left in Figure 7.1; the graph on the right represents this distribution as a probability histogram. The probability histogram representation allows us to easily visualize the probability of an event as the sum of the areas of the rectangles over the values of X which comprise the event.

Figure 7.1 Binomial distribution n = 3 p = .5 - p.m.f. and probability histogram.



The distribution of X can also be characterized in terms of its **cumulative distribution** function (denoted c.d.f.) F_X , where

$$F_X(x) = P(X \le x).$$

For discrete random variables the p.m.f. usually gives the more convenient characterization of the distribution. The c.d.f. is useful for computing probabilities.

7.2 Some examples

Example 7.2 Tossing a die – uniform distribution. Suppose that a fair (balanced) die is tossed and let X denote the number on the upturned face. The sample space for X is $\Omega_X = \{1, 2, 3, 4, 5, 6\}$. Since the die is assumed to be fair we will assume that these 6 outcomes are equally probable. (Note that tossing a fair die once is equivalent to selecting a ball at random from a box containing six balls numbered from one to six.) Thus, the p.m.f. of X is given by

$$p_X(x) = \begin{cases} 1/6 & \text{if } x \in \{1, 2, 3, 4, 5, 6\} \\ 0 & \text{otherwise} \end{cases}$$

This is an example of a uniform distribution on a finite set of integers. This uniform p.m.f. is shown in the graph on the left in Figure 7.2; the graph on the right represents this distribution as a probability histogram.

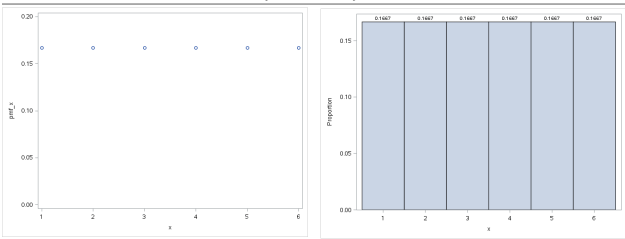


Figure 7.2 Uniform distribution on $\{1, 2, 3, 4, 5, 6\}$ – p.m.f. and probability histogram.

Example 7.3 The sum when a die is tossed twice – triangular distribution. Suppose that a fair (balanced) die is tossed twice and let X denote the sum of the numbers observed. The elementary outcomes for this experiment can be represented by the 36 ordered pairs of the form (a, b) where $a, b \in \{1, 2, 3, 4, 5, 6\}$. The sums and their relationship with these elementary outcomes are shown in Table 7.2. Here the row label represents the number on the first toss, the column label represents the number on the second toss, and the value in the body of the table is the corresponding sum. Because the sums range from 2 to 12 the sample space for X is $\Omega_X = \{2, 3, ..., 12\}$.

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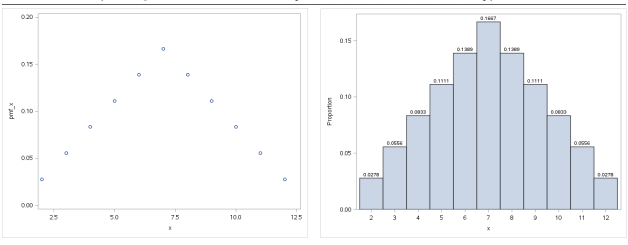
Tab	le 7.2	Sums	when a	a die is	tossed t	twice.
	1	2	3	4	5	6
1	2	3	4	5	6	7
2	3	4	5	6	7	8
3	4	5	6	7	8	9
4	5	6	7	8	9	10
5	6	7	8	9	10	11
6	7	8	9	10	11	12

Since the die is assumed to be fair we will assume that these 36 elementary outcomes are equally probable. Counting the number of favorable outcomes for each case we find that the p.m.f. of X is given by

$$p_X(x) = \begin{cases} 1/36 & \text{if } x = 2 \text{ or } x = 12\\ 2/36 & \text{if } x = 3 \text{ or } x = 11\\ 3/36 & \text{if } x = 4 \text{ or } x = 10\\ 4/36 & \text{if } x = 5 \text{ or } x = 9\\ 5/36 & \text{if } x = 6 \text{ or } x = 8\\ 6/36 & \text{if } x = 7\\ 0 & \text{otherwise} \end{cases}$$

In this example, we can think of selecting a ball at random from a box containing 36 balls of which 1 is numbered two, 2 are numbered three, 3 are numbered four, and so on. This p.m.f. is shown in the graph on the left in Figure 7.3; the graph on the right represents this distribution as a probability histogram. This is an example of a triangular distribution on a finite set of integers.

Figure 7.3 Distribution of sum of two fair dice – p.m.f. and probability histogram. (Triangular distribution on {2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12})



7.3 The binomial distribution

In Section 5.4 we introduced the binomial distribution as the distribution of the number of objects of one type in a sample selected at random with replacement. We also discussed two typical applications of the binomial distribution as the distribution of a random variable X denoting the number of objects of a specified type in a sample. In Example 5.8 X denoted the number of acces observed when a fair die was tossed 10 times. In Example 5.9 X denoted the number of red balls in a sample selected at random with replacement from a population containing 100 red balls and 500 blue balls.

We will now provide a slightly different description of the binomial distribution. Consider an experiment consisting of a sequence of n independent, dichotomous trials, where a trial is a process of observation or experimentation which results in the occurrence of one of two possible outcomes. The two possible outcomes are generically known as "success" and "failure". There is no connotation of "goodness" associated with the term "success", this term is simply used to indicate a success in the sense that the outcome of interest has occurred. In Example 5.8 a trial is a toss of the die and tossing an ace (one) is a success. In Example 5.9 a trial is the selection of a ball from the population and obtaining a red ball is a success. The success probability p is the probability of observing a success on a single trial and 1 - p is the probability of observing a failure on a single trial. The random variable of interest X is the number of successes observed in the sequence of n independent trials.

A sequence of independent Bernoulli trials

Given a positive integer n and a probability p (0), a sequence of independentBernoulli trials with success probability <math>p is a sequence of n dichotomous (success or failure) trials with the properties that:

- 1. On each trial the probability of success is p; and,
- 2. The outcomes of the trials are independent.

A binomial r.v. X denotes the number of successes in a sequence of n independent Bernoulli trials with success probability p.

toc

The binomial probability mass function

Given a positive integer n and a probability p (0), the binomial random variable <math>X denotes the number of successes in a sequence of n independent Bernoulli trials with success probability p. For x = 0, 1, ..., n, the **binomial probability mass function** is of the form

$$p_X(x) = \binom{n}{x} p^x (1-p)^{n-x}$$

Example 7.4 Number of red balls in a sample. Suppose we select a random sample of n = 4 balls with replacement from a box containing 60 balls of which 10 are red. We can think of the selection of a single ball from the box as a trial. Since the balls are being selected with replacement the outcomes of the four trials are independent and the probability of obtaining a red ball is 1/6 for each selection, we can view these four selections of a ball as forming a sequence of n = 4 Bernoulli trials with success probability p = 1/6. Hence, letting X denote the number of red balls in the four selections, X is the binomial random variable with p.m.f.

$$p_X(x) = \binom{4}{x} \left(\frac{1}{6}\right)^x \left(\frac{5}{6}\right)^{4-x}$$
 for $x = 0, 1, 2, 3, 4.$

Example 7.5 Number of aces when tossing a fair die. Suppose a fair (balanced) die is tossed n = 4 times and the number of aces (ones) is determined. Since the outcomes of these four tosses are independent and the probability of obtaining an ace is 1/6 on each toss, we can view these four tosses as forming a sequence of n = 4 Bernoulli trials with success probability p = 1/6. Hence, letting X denote the number of aces in the four tosses, X is the binomial random variable with p.m.f.

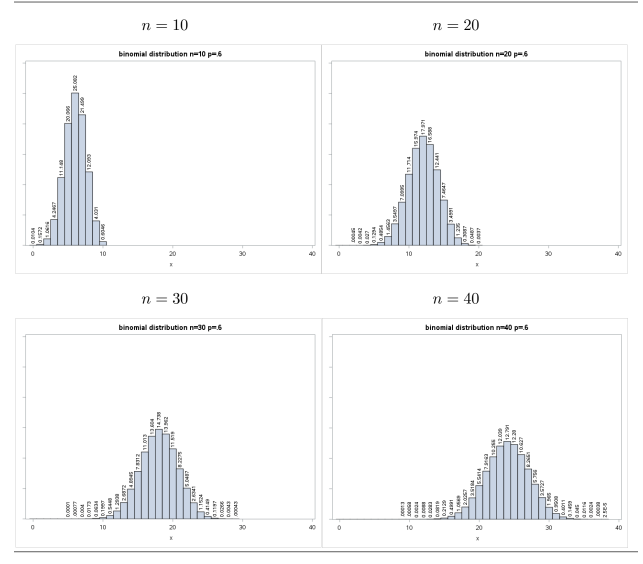
$$p_X(x) = \binom{4}{x} \left(\frac{1}{6}\right)^x \left(\frac{5}{6}\right)^{4-x}$$
 for $x = 0, 1, 2, 3, 4$.

Some properties of binomial distributions are illustrated in Figures 7.4 and 7.5. Figure 7.4 shows that the binomial distribution is centered at np and the variability in the distribution increases as the sample size n increases. In this figure the success probability is fixed at p = .6 and histograms are provided for several sample sizes (n = 10, 20, 30, and 40). Figure 7.5 shows how the shape of the binomial distribution and the variability in the distribution depend on the value of the success probability p. In this figure the sample size

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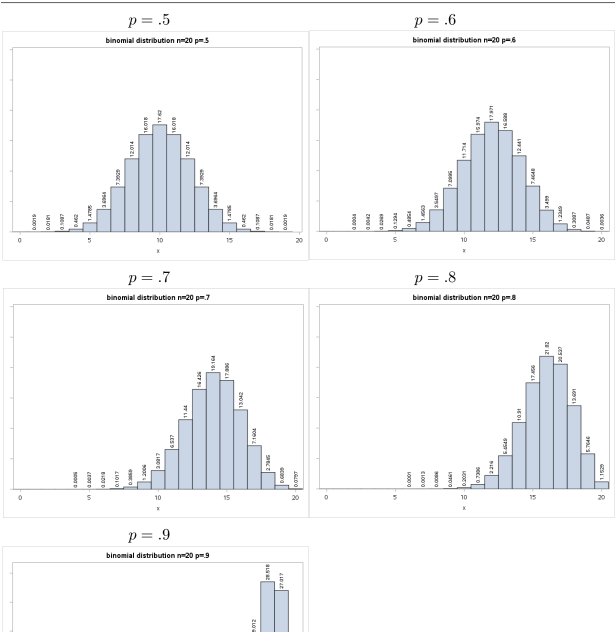
is fixed at n = 20 and histograms are provided for several values of the success probability (p = .5, .6, .7, .8, and .9). When p = .5, the distribution is exactly symmetric about np = 10. When $p \neq .5$ the binomial distribution is skewed and as the value of p moves away from .5 the distribution becomes more skewed. In this figure the values p = .6, .7, .8, and .9 are greater than .5 and the distributions are skewed left. if we had used values less than .5 the distributions would be skewed right. Notice that in addition to the increase in skewness there is a decrease in variability as p moves away from .5.

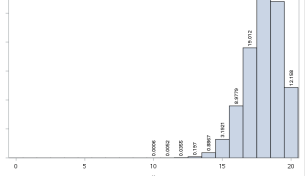
Figure 7.4 Binomial distributions for p = .6. Histograms are provided for n = 10, 20, 30, and 40. These distributions are centered at np for p = .6 and the appropriate value of n. Notice how the variability in the distribution increases as n increases.



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Figure 7.5 Binomial distributions for n = 20. Histograms for p = .5, .6, .7, .8, and .9. Each distribution is centered at 20p for the appropriate p. Notice how the variability in the distribution decreases as p moves away from .5. Notice also how the distribution becomes more skewed left as p moves away from .5.





When n = 1 our sequence of trials reduces to a single Bernoulli trial with X = 1 indicating that a success has occurred and X = 0 indicating a failure. The binomial distribution for the case when n = 1 is also know as a **Bernoulli distribution**. We will now consider an interesting application.

Example 7.6 Mendelian inheritance models. In his investigations, during the years 1856 to 1868, of the chromosomal theory of inheritance Gregor Mendel performed a series of experiments on ordinary garden peas. One characteristic of garden peas that Mendel studied was the color of the flowers (red or white). When Mendel crossed a plant with red flowers with a plant with white flowers, the resulting offspring all had red flowers. But when he crossed two of these first generation plants, he observed plants with white as well as red flowers.

The gene which determines the color of the flower occurs in two forms (alleles). Let R denote the allele for red flowers (which is dominant) and r denote the allele for white flowers (which is recessive). When two plants are crossed the offspring receives one allele from each parent, thus there are four possible genotypes (combinations) RR, Rr, rR, and rr. The three genotypes RR, Rr, and rR, which include the dominant R allele, will yield red flowers while the fourth genotype rr will yield white flowers. If a red flowered RR genotype parent is crossed with a white flowered rr genotype parent, then all of the offspring will have genotype Rr and will produce red flowers. The basic Mendelian inheritance model assumes that a pair of alleles is formed by randomly choosing one allele from each parent. Under this model, if two of these first generation Rr genotype plants are crossed, each of the four possible genotypes RR, Rr, rR, and rr is equally likely and plants with white as well as red flowers will occur. Under this simple model, with each of the four genotypes having the same probability of occurring, the probability that a plant will have red flowers is P(red) = 3/4 and the probability that a plant will have white flowers is P(white) = 1/4.

The distribution of the r.v. X denoting flower color, with X = 1 indicating the plant has red flowers and X = 0 indicating the plant has white flowers, follows the Bernoulli distribution with p = 3/4.

We could test the validity of this model, as Mendel did, by crossing n pairs of peas plants and determining the color of the flowers for these n crosses. If we let X denote the number of these n crosses which result in a red flowered offspring, then X will follow the binomial distribution with parameters n and p, where, according to Mendel's model p = 3/4.

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7.4 The hypergeometric distribution

In Section 5.5 we introduced the hypergeometric distribution distribution as the distribution of the number of objects of one type in a sample selected at random without replacement. We discussed a typical application of the hypergeometric distribution as the distribution of a random variable X denoting the number of objects of a specified type in a sample. In Example 5.10 X denoted the number of aces in a five card poker hand. For ease of reference the hypergeometric probability mass function is given below. Here we have simplified the notation compared to that of Section 5.5.

The hypergeometric probability mass function

Given positive integers n, A, and B with $n \leq A+B$, the hypergeometric random variable X denotes the number of successes in random sample of size n selected without replacement from a population containing A + B objects of which A are classified as successes and B are classified as failures. Subject to some restrictions noted below, for $x = 0, 1, \ldots, n$, the **hypergeometric probability mass function** is of the form

$$p_X(x) = \frac{\binom{A}{x}\binom{B}{n-x}}{\binom{A+B}{n}}.$$

Notice that the values of A and B may impose restrictions on the values of x for which this expression works. Specifically, we must have $x \leq A$ and $n - x \leq B$.

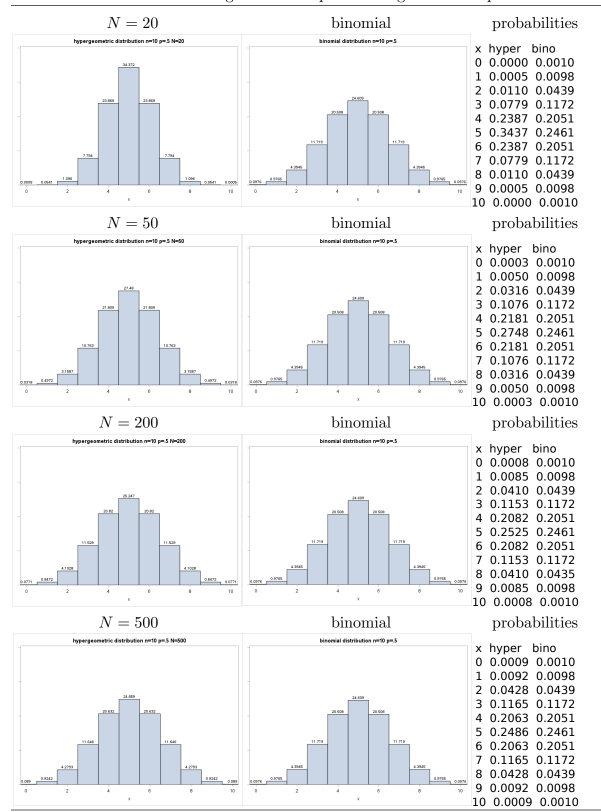
Example 7.7 The number of aces in a poker hand. Suppose that a five card poker hand is dealt from a well-shuffled deck. Dealing a five card hand in this way is equivalent to selecting five balls (cards) at random from a box containing 52 balls (cards) labeled so that they represent the cards in the deck. Let X denote the number of aces in the hand. In this application the 52 cards are partitioned into the set of A = 4 aces and the set of B = 48 non-aces. For x = 0, 1, 2, 3, 4, the p.m.f. for this X is

$$p_X(x) = \frac{\binom{4}{x}\binom{48}{5-x}}{\binom{52}{5}}.$$

For example, the probability of getting exactly 4 aces is

$$p_X(4) = \frac{\binom{4}{4}\binom{48}{1}}{\binom{52}{5}} = \frac{1 \cdot 48}{2598960} \approx .00002.$$

Figure 7.6 Hypergeometric and binomial distributions compared, with n = 10, p = .5. Left column: hypergeometric histograms for N = A + B = 20, 50, 200, and 500. Middle column: binomial histogram for comparison. Right column: probabilities.



126 7.5 The geometric distribution

For given values of the sample size n and the success proportion p, there is more variability in a hypergeometric distribution than a binomial distribution. The pattern of dependence of the basic shape of hypergeometric distributions on the values on n and p are similar to those for binomial distributions. In particular, when p = 1/2 the hypergeometric distribution is symmetric, when p < 1/2 it is skewed right, and when p > 1/2 it is skewed left. When the population size N = A + B is large, the hypergeometric and binomial distribution, for given values of n and p, are very similar. This relationship between the hypergeometric and binomial distribution is illustrated, graphically and numerically, for n = 10, p = 1/2 in Figure 7.6.

7.5 The geometric distribution

The binomial and hypergeometric distributions can be used to model the distribution of the number of red balls (successes) in a sample of fixed size. We will now explore a variation on this sampling procedure where balls are sampled, one at a time with replacement, until a ball of the specified color (a success) is obtained.

Consider a box containing 20 balls of which 5 are red and 15 are blue. First a single ball is selected at random from the 20 balls in the box and its color is determined. If the ball is red we stop sampling. On the other hand, if the ball is blue, it is returned to the box and this procedure is repeated until a red ball is obtained. Let X denote the number of the trial (draw) on which the red ball is obtained. For example, X = 1 means we got a red ball on the first draw and X = 10 means we got a blue ball on each of the first 9 draws and then a red ball on the tenth draw. Notice that there is no upper limit on how many draws might be needed to obtain a red ball so that, in this situation, the sample space for X is the positive integers (the counting numbers) 1, 2, 3, ...

We can view the results of this sampling process as forming a, potentially infinite, sequence of trials, where a trial is the selection of a ball from the box. In the present context, since we are sampling with replacement, the outcomes of the trials (draws) are independent and on every trial the probability of selecting a red ball is $p = \frac{5}{20} = \frac{1}{4}$. In other words, we are considering a potentially infinite sequence of independent Bernoulli trials with success probability $p = \frac{1}{4}$ which ends when the first success is obtained. In this context, using R to denote a red ball and G to denote a blue ball, the results of the sequence of trials (the elementary outcomes) can be represented by sequences of the form R, BR, BBR, Notice that, letting x denote the number of the trial on which the red ball was obtained,

 toc

these elementary outcome sequences contain x - 1 B's followed by a single R. Notice also that, since the outcomes of the trials are independent, the probability of observing x - 1B's (x - 1 blue balls) followed by 1 R (one red ball) is

$$p_X(x) = \left(\frac{15}{20}\right)^{x-1} \left(\frac{5}{20}\right).$$

The geometric probability mass function

Given a probability p, with 0 , consider a possibly infinite sequence of independentBernoulli trials with success probability <math>p. The geometric random variable X denotes the number of the trial on which the first success occurs. For x = 1, 2, 3..., the **geometric probability mass function** is of the form

$$p_X(x) = (1-p)^{x-1}p.$$

Example 7.8 Tossing a fair die until we get an ace. If a fair die is tossed repeatedly until an ace (a one) appears, then the probability of observing this initial ace on the toss x is $(\frac{5}{6})^{x-1}(\frac{1}{6})$. For example:

The probability that the first ace appears on the first toss is $(\frac{5}{6})^0(\frac{1}{6}) \approx .1667$; The probability that the first ace appears on the second toss is $(\frac{5}{6})^1(\frac{1}{6}) \approx .1389$; The probability that the first ace appears on the third toss is $(\frac{5}{6})^2(\frac{1}{6}) \approx .1157$; while, the probability that the first ace appears on the tenth toss is $(\frac{5}{6})^9(\frac{1}{6}) \approx .0323$; and, the probability that the first ace appears on the twentieth toss is $(\frac{5}{6})^{19}(\frac{1}{6}) \approx .0052$.

Example 7.9 Tossing a fair coin until we get a head. If a fair coin is tossed repeatedly until a head appears, then the probability of observing this initial head on the toss x is $(\frac{1}{2})^{x-1}(\frac{1}{2}) = (\frac{1}{2})^x$. For example:

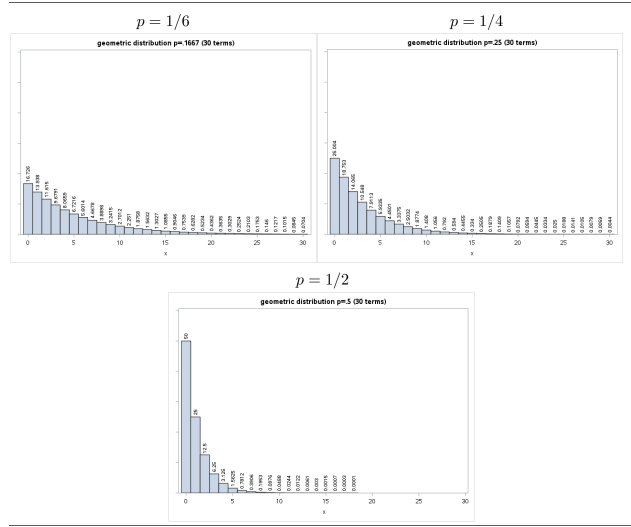
The probability that the first ace appears on the first toss is $(\frac{1}{2})^1 = .5$; The probability that the first ace appears on the second toss is $(\frac{1}{2})^2 = .25$; The probability that the first ace appears on the third toss is $(\frac{1}{2})^3 = .125$; while, the probability that the first ace appears on the tenth toss is $(\frac{1}{2})^{10} \approx .0010$; and, the probability that the first ace appears on the twentieth toss is $(\frac{1}{2})^{20} \approx .000001$.

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Graphical representations (with percentages) of the geometric distributions with p = 1/6 (the die tossing Example 7.8), p = 1/4 (the 5 red and 15 blue balls example), and p = 1/2 (the coin tossing Example 7.9) are provided in Figure 7.7. Note that in these plots the x-axis is "X - 1" the number of failures before first success.

Terminology note: Some authors define the geometric random variable as Y = X - 1, the number of failures before the first success.

Figure 7.7 Geometric distributions for p = 1/6, p = 1/4, p = 1/2 (first 30 terms) In these plots the *x*-axis is "X - 1" the number of failures before first success.



7.6 The Poisson distribution

In many settings, the Poisson distribution provides a realistic model for a random variable representing the number of occurrences of a "rare event". Consider a sequence of events occurring randomly in time or space and a count such as the number of radioactive particle emissions per unit time, the number of meteorites that collide with a satellite during a single orbit, the number of defects per unit length of some material, or the number of weed seeds per unit volume in a large batch of wheat seeds. We can picture the time (or location) of each occurrence as a point on the positive part of the number line. Consider the following assumptions about the times (locations) of these occurrences:

1. The probability of exactly one occurrence in a small interval of length t is approximately νt , where $\nu > 0$ is the mean rate at which events occur per unit time (the mean rate of occurrence).

2. The probability of more than one occurrence in a small interval of length t is negligible compared to the probability of exactly one occurrence in a small interval of length t.

3. The numbers of occurrences in non-overlapping intervals are independent in the sense that information concerning the number of events in one interval reveals nothing about the number of events in any other interval.

If we let X denote the number of occurrences in a period of length t, then these three assumptions imply that X follows the Poisson distribution with parameter $\lambda = \nu t$. The possible values of X are $0, 1, \ldots$, with no theoretical upper bound on the value.

The Poisson probability mass function

Given a constant $\lambda > 0$. For x = 1, 2, 3..., the **Poisson probability mass function** is of the form

$$P(X=x) = \frac{\lambda^x}{x!}e^{-\lambda}$$

where $e \approx 2.718$ is the base of the natural logarithm.

Two applications of the Poisson distribution are provided in the following examples.

Example 7.10 Example. Radioactive disintegrations. This example is taken from Feller (1957), p. 149 and Cramér (1946) p. 436. In a famous experiment by Rutherford, Chadwick, and Ellis (*Radiations from Radioactive Substances*, Cambridge, 1920) a radioactive

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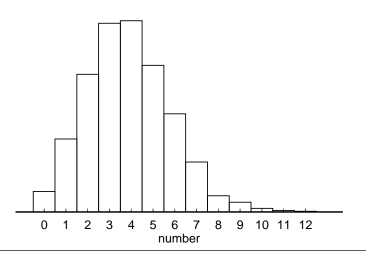
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substance was observed during 2608 consecutive time intervals of length t = 7.5 seconds each. The number of particles reaching a counter was recorded for each period. The results are summarized in Table 7.3 and Figure 7.8. (In this table the observations greater than or equal to 10 are grouped together. The data actually contain 10 tens, 4 elevens, and 2 twelves.) The last column of Table 7.3 contains expected relative frequencies (probabilities) computed using a Poisson model with λ estimated from these data. These Poisson probabilities appear to match the observed relative frequencies fairly well. A formal test of the goodness of fit of this Poisson model to these data indicates that the model does fit well ($\chi^2 = 12.885$, 9 d.f., *P*-value .17).

expected relative frequency
froquoney
nequency
.0209
.0807
.1562
.2015
.1949
.1509
.0973
.0538
.0260
.0112
.0065
.9999

 Table 7.3 Relative frequency distribution for radioactive disintegrations.

Figure 7.8	Histogram	for	radioactive	disintegrations	(with >	10	expanded)).
				0-0-0-0-0-0	(···=•==		/	/ -

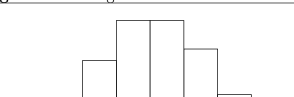


Example 7.11 Bacteria counts. This example is taken from Feller (1957), p.153. The original source is T. Matuszewsky, J. Supinska, and J. Neyman (1936), Zentralblatt für Bakteriologie, Parasitenkunde und Infektionskrankrankheiten, II Abt., 95. A Petri dish with bacteria colonies was examined under a microscope. The dish was divided into small squares and the number of bacteria colonies, visible as dark spots, was recorded for each square. In this example t is the area the square within which the count is determined and we will take this area to be one. If the bacteria colonies were randomly distributed over the Petri dish, without being clustered together, then the Poisson model should hold. The results for one of several experiments are summarized in Table 7.4 and Figure 7.9. The last column of Table 7.4 contains expected relative frequencies (probabilities) computed using a Poisson model, with λ estimated from these data. In this example the observed relative frequency in the " ≥ 6 " line is for "exactly 6", but, the expected relative frequency is for all values greater than or equal to 6. These Poisson probabilities appear to match the observed relative frequencies fairly well. Therefore, the evidence supports the contention that the bacteria colonies are randomly distributed over the Petri dish. A formal test of the goodness of fit of this Poisson model to these data indicates that the model does fit well ($\chi^2 = .8386, 5$ d.f., *P*-value .9745).

	observed	observed relative	expected relative
number	frequency	frequency	frequency
0	5	.0424	.0533
1	19	.1610	.1562
2	26	.2203	.2290
3	26	.2203	.2239
4	21	.1780	.1641
5	13	.1102	.0962
≥ 6	8	.0678	.0772
total	118	1.0000	.9999

 Table 7.4 Relative frequency distribution for bacteria counts.

132 7.7 Expected value





2

1

3

number

4

5

6

7.7 Expected value

0

 toc

There are many situations where we might wonder what value to expect when we perform an experiment and observe the value of a particular random variable. Before we can address this topic more formally we need to explain exactly what we mean when we say "what value to expect". The expected value of a random variable can be viewed as the long run average value of the random variable. That is, the average of the values of the random variable we would obtain if we conducted the experiment and observed the value of the random variable a large number of times. Before we introduce a formal definition it is helpful to consider a simple example of a real world application of an expected value.

Example 7.11 Expected winnings in roulette An American style roulette wheel, as shown in Figure 7.10, has 38 pockets of which 18 are red, 18 are black, and 2 are green. We will consider the expected winnings for a one dollar bet on red (a bet that the ball will land in one of the 18 red pockets).

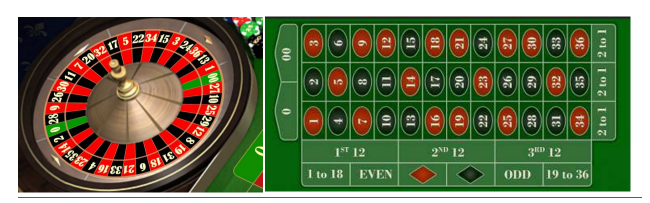


Figure 7.10 An American roulette wheel and betting table

Let the random variable W denote our winnings. This random variable assumes two values: W = 1 indicates that the ball lands in a red pocket and we win one dollar; while W = -1indicates that the ball lands in a black or green pocket and we lose one dollar. Since 18 of the 38 pockets are red, the probability mass function for W is given by

$$p_W(w) = \begin{cases} 18/38 & \text{if } w = 1\\ 20/38 & \text{if } w = -1\\ 0 & \text{otherwise} \end{cases}$$

If we imagine placing this bet a large number of times, then we see that in about 18 out of 38 tries (about 47.37% of the time) we will win a dollar (we will observe W = 1) and in about 20 out of 38 tries (about 52.63% of the time) we will lose a dollar (we will observe W = -1). Thus, on average in the long run, we expect to see an average winnings of (18 - 20)/38 = -2/38 dollars (about -.0526 dollars or -5.26 cents). That is, if we made a large number of bets, on average, we would lose 5.26 cents per bet. The long run average winnings we computed, (18 - 20)/38 = -2/38 dollars, is called the expected values of W. The notation for the expected value of W is E(W).

We will now provide a formal definition of the expected value of a discrete random variable.

If X is a discrete r.v. with finite sample space $\Omega_X = \{x_1, \ldots, x_N\}$ and p.m.f. p_X , then the **expected value** of X, denoted by E(X) or μ_X , is defined as the weighted average of the possible values of X (the elements of Ω_X) obtained using the associated probabilities (the corresponding values of $p_X(x)$) as weights. In symbols we have

$$E(X) = x_1 p_X(x_1) + x_2 p_X(x_2) + \dots + x_N p_X(x_N) = \sum_{i=1}^N x_i p_X(x_i).$$

Example 7.12 Two distributions to illustrate computations Let X and Y denote r.v.'s with the following p.m.f.'s

$$p_X(x) = \begin{cases} .1 & \text{if } x = 1 \\ .2 & \text{if } x = 2 \\ .4 & \text{if } x = 3 \\ .2 & \text{if } x = 4 \\ .1 & \text{if } x = 5 \\ 0 & \text{otherwise} \end{cases} p_Y(y) = \begin{cases} .1 & \text{if } y = 1 \\ .1 & \text{if } y = 2 \\ .2 & \text{if } y = 3 \\ .2 & \text{if } y = 4 \\ .4 & \text{if } y = 5 \\ 0 & \text{otherwise} \end{cases}$$

The expected values of X and Y are:

$$E(X) = .1(1) + .2(2) + .4(3) + .2(4) + .1(5) = .1 + .4 + 1.2 + .8 + .5 = 3$$

134 7.7 Expected value

$$E(Y) = .1(1) + .1(2) + .2(3) + .2(4) + .4(5) = .1 + .2 + .6 + .8 + 2.0 = 3.7.$$

A tabular representation of these computations may be helpful. In these tabular representations the expected value is the sum of the $xp_X(x)$ column.

x	$p_X(x)$	$xp_X(x)$	y	$p_Y(y)$	$yp_Y(y)$
1	.1	$1 \times .1 = .1$	1	.1	$1 \times .1 = .1$
2	.2	$2 \times .2 = .4$	2	.1	$2 \times .1 = .2$
3	.4	$3 \times .4 = 1.2$	3	.2	$3 \times .2 = .6$
4	.2	$4 \times .2 = .8$	4	.2	$4 \times .2 = .8$
5	.1	$5 \times .1 = .5$	5	.4	$5 \times .4 = 2.0$
		E(X) = 3			E(Y) = 3.7

If the sample space is countably infinite (as with the geometric or Poisson distribution), then $\Omega_X = \{x_1, x_2, \ldots\}$ is countably infinite and the sum in the definition of E(X) has an infinite number of terms (does not stop at a finite N).

An aside – existence of expected values

Some technicalities may arise when Ω_X is countably infinite. The problem is that for some choices of the p.m.f. p_X the series (infinite sum) in the definition of the expected value may not exist. (More formally, this series may not converge.) We do not need to worry about this technicality, since the expected value does exist for all of the discrete random variables with countably infinite sample spaces we will encounter.

Expected value as center of mass

If X is a discrete r.v. with finite sample space $\Omega_X = \{x_1, \ldots, x_N\}$ and p.m.f. p_X , then, as the name suggests, we can think of the probabilities $p_X(x_1), \ldots, p_X(x_N)$ as masses located at points x_1, \ldots, x_N on a segment of the number line. With this interpretation the center of mass of the distribution of X is located at $E(X) = x_1 p_X(x_1) + \cdots + x_N p_X(x_N)$. Similarly, if $\Omega_X = \{x_1, x_2, \ldots\}$, then the center of mass of the distribution of X is located at $E(X) = x_1 p_X(x_1) + x_2 p_X(x_2) + \cdots$.

Expected value as population mean

If an individual is selected at random from a population of N people and X represents the individual's height in inches so that Ω_X represents the collection of distinct heights for this population and $p_X(x_i)$ is the proportion of individuals in the population with height x_i , then E(X) is the population mean height (the average height of all N people in the population).

The expected value of a function of X

We will now show how the definition of the expected value of X can be extended to the expected value of a function of X. The expected value of a function of X may be of interest in some applications. Also, as we will see shortly, the variance of X is defined as the expected value of a specific function of X. Let g denote the function of interest and let Ω_Y denote the range of the function, that is, Ω_Y is the collection of values obtained when the function g is applied to the elements of Ω_X . If Ω_X and Ω_Y are discrete, then Y = g(X) is a discrete r.v. and there are two ways to compute the expected value of Y. Letting $\Omega_X = \{x_1, x_2, \ldots x_N\}$ and $\Omega_Y = \{y_1, y_2, \ldots y_M\}$:

(1) We can find the p.m.f. p_Y of Y and compute the expected value of Y as

$$E(Y) = y_1 p_Y(y_1) + y_2 p_Y(y_2) + \dots + y_M p_Y(y_M) = \sum_{i=1}^M y_i p_Y(y_i)$$

(2) We can use the p.m.f. p_X of X to compute the expected value of Y as

$$E(Y) = E(g(X)) = g(x_1)p_X(x_1) + g(x_2)p_X(x_2) + \dots + g(x_N)p_X(x_N) = \sum_{i=1}^N g(x_i)p_X(x_i).$$

The analogous expressions with infinite sums apply in the countably infinite case. We will demonstrate these computations in the context of finding the expected winnings of a bet on red in the roulette example.

Example 7.11 Expected winnings in roulette (revisited) The 36 pockets on the roulette wheel shown in Figure 7.10, are numbered as follows: pockets 0 and 00 are green; pockets 1,3,5,7,9,12,14,16,18,19,21,23,25,27,30,32,34, and 36 are red; and,

pockets 2,4,6,8,10,11,13,15,17,20,22,24,26,28,29,31,33, and 35 are black. If we let X denote the number on the pocket the ball lands in, then the r.v. X follows the uniform distribution with $\Omega_X = \{0, 00, 1, 2, \dots, 36\}$. In terms of this X the function g which defines the

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winnings for a bet on red assigns the value 1 to the numbers associated with the red pockets and assigns the value -1 to the numbers associated with the green and black pockets so that the r.v. W = g(X) is the winnings for a bet on red. As noted earlier, $E(W) = (1)\frac{18}{38} + (-1)\frac{20}{38} = -\frac{2}{38}$. Using the relationship W = g(X) and the uniform distribution of X, with some regrouping, we have

$$E(W) = (1) [p_X(1) + p_X(3) + \dots + p_X(34) + p_X(36)] + (-1) [p_X(2) + p_X(4) + \dots + p_X(33) + p_X(35) + p_X(0) + p_X(00)] = (1) \frac{18}{38} + (-1) \frac{20}{38} = -\frac{2}{38},$$

since $p_X(x) = \frac{1}{38}$ for the 38 values in Ω_X and there are 18 values with g(x) = 1 and 20 values with g(x) = -1.

Example 7.12 Two distributions to illustrate computations, revisited Recall that X and Y denote r.v.'s with the following p.m.f.'s

$$p_X(x) = \begin{cases} .1 & \text{if } x = 1 \\ .2 & \text{if } x = 2 \\ .4 & \text{if } x = 3 \\ .2 & \text{if } x = 4 \\ .1 & \text{if } x = 5 \\ 0 & \text{otherwise} \end{cases} p_Y(y) = \begin{cases} .1 & \text{if } y = 1 \\ .1 & \text{if } y = 2 \\ .2 & \text{if } y = 3 \\ .2 & \text{if } y = 4 \\ .4 & \text{if } y = 5 \\ 0 & \text{otherwise} \end{cases}$$

We will now find the expected values of X^2 and Y^2 . The reason that these expected values are of interest will become clear shortly.

$$E(X^{2}) = .1(1^{2}) + .2(2^{2}) + .4(3^{2}) + .2(4^{2}) + .1(5^{2})$$

= .1(1) + .2(4) + .4(9) + .2(16) + .1(25)
= .1 + .8 + 3.6 + 3.2 + 2.5 = 10.2
$$E(Y^{2}) = .1(1^{2}) + .1(2^{2}) + .2(3^{2}) + .2(4^{2}) + .4(5^{2})$$

= .1(1) + .1(4) + .2(9) + .2(16) + .4(25)
= .1 + .4 + 1.8 + 3.2 + 10.0 = 15.5.

x	$p_X(x)$	$x^2 p_X(x)$	y	$p_Y(y)$	$y^2 p_Y(y)$
1	.1	$1 \times .1 = .1$	1	.1	$1 \times .1 = .1$
2	.2	$4 \times .2 = .8$	2	.1	$4 \times .1 = .4$
3	.4	$9 \times .4 = 3.6$	3	.2	$9 \times .2 = 1.8$
4	.2	$16 \times .2 = 3.2$	4	.2	$16 \times .2 = 3.2$
5	.1	$25 \times .1 = 2.5$	5	.4	$25 \times .4 = 10.0$
		$\overline{E(X^2)} = 10.2$			$E(Y^2) = 15.5$

As before, a tabular representation of these computations may be helpful. In these tabular representations the expected value of X^2 is the sum of the $x^2 p_X(x)$ column.

7.8 Variance

The expected value or mean of the distribution of X, $E(X) = \mu_X$, provides an indication of where the "center" of the distribution is located on the number line. It would also be useful to have a measure of the amount of variability in the distribution of X. The variance is one such measure of variability. The variance of the distribution of X is the long run average value of the square of the distance between the points in the sample space Ω_X and the mean μ_X of the distribution. More formally, letting $E(X) = \mu_X$ denote the mean of the distribution of X, and assuming that this mean exists, the function $g(x) = (x - \mu_X)^2$ (the squared deviation of x from the mean of the distribution of X) can be used to define a measure of the variability in the distribution of X. The expected value of the r.v. $g(X) = (X - \mu_X)^2$ is the variance of the distribution of X.

If X is a discrete r.v. with sample space $\Omega_X = \{x_1, \ldots, x_N\}$, p.m.f. p_X , and mean $\mu_X = E(X)$, then the **variance** of X, denoted by var(X) or σ_X^2 , is defined by

$$\operatorname{var}(X) = (x_1 - \mu_X)^2 p_X(x_1) + \dots + (x_N - \mu_X)^2 p_X(x_N) = \sum_{i=1}^N (x_i - \mu_X)^2 p_X(x_i).$$

Similarly, if $\Omega_X = \{x_1, x_2, \ldots\}$, then the variance is

$$\operatorname{var}(X) = (x_1 - \mu_X)^2 p_X(x_1) + (x_2 - \mu_X)^2 p_X(x_2) + \dots = \sum_{i=1}^{\infty} (x_i - \mu_X)^2 p_X(x_i).$$

Since squared distances are used in its definition the scale of measurement for the variance of X is the square of the scale of measurement for X itself. For example, if X denotes

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the height of a person in inches, then the variance of X, which provides a measure of variability among the heights of different individuals, is measured in inches squared. This change of scale is undesirable in practical applications where we really need a measure of variability on the original scale of measurement. To obtain a measure of variability on the original scale of measurement. To obtain a measure of variability on the original scale of measurement. To obtain a measure of variability on the standard deviation. The principal (positive) square root of the variance of X, $SD(X) = \sigma_X = \sqrt{var(X)}$, is know as the **standard deviation** of X.

Example 7.12 Two distributions to illustrate computations, revisited Recall that X and Y denote r.v.'s with the following p.m.f.'s

$$p_X(x) = \begin{cases} .1 & \text{if } x = 1 \\ .2 & \text{if } x = 2 \\ .4 & \text{if } x = 3 \\ .2 & \text{if } x = 4 \\ .1 & \text{if } x = 5 \\ 0 & \text{otherwise} \end{cases} p_Y(y) = \begin{cases} .1 & \text{if } y = 1 \\ .1 & \text{if } y = 2 \\ .2 & \text{if } y = 3 \\ .2 & \text{if } y = 4 \\ .4 & \text{if } y = 5 \\ 0 & \text{otherwise} \end{cases}$$

and that E(X) = 3 and E(Y) = 3.7. We will now find the variances and standard deviations of X and Y.

$$\operatorname{var}(X) = .1(1-3)^2 + .2(2-3)^2 + .4(3-3)^2 + .2(4-3)^2 + .1(5-3)^2$$

= $.1(-2)^2 + .2(-1)^2 + .4(0)^2 + .2(1)^2 + .1(2)^2$
= $.4 + .2 + 0 + .2 + .4 = 1.2$
$$\operatorname{var}(Y) = .1(1-3.7)^2) + .1(2-3.7)^2 + .2(3-3.7)^2 + .2(4-3.7)^2 + .4(5-3.7)^2$$

= $.1(-2.7)^2 + .1(-1.7)^2 + .2(-0.7)^2 + .2(0.3)^2 + .4(1.3)^2$
= $.729 + .289 + .098 + .018.676 = 1.81.$

The standard deviations are $SD(X) = \sigma_X = \sqrt{1.2} \approx 1.0954$ and $SD(Y) = \sigma_Y = \sqrt{1.81} \approx 1.3454$.

Again, a tabular representation of these computations may be helpful. In these tabular representations the variance of X is the sum of the $(x - E(X))^2 p_X(x)$ column.

x	$p_X(x)$	$(x - E(X))^2 p_X(x)$	y	$p_Y(y)$	$(y - E(Y))^2 p_Y(y)$
1	.1	$(-2)^2 \times .1 = .4$	1	.1	$(-2.7)^2 \times .1 = .729$
2	.2	$(-1)^2 \times .2 = .2$	2	.1	$(-1.7)^2 \times .1 = .289$
3	.4	$(0)^2 \times .4 = 0$	3	.2	$(7)^2 \times .2 = .098$
4	.2	$(1)^2 \times .2 = .2$	4	.2	$(.3)^2 \times .2 = .018$
5	.1	$(2)^2 \times .1 = .4$	5	.4	$(1.3)^2 \times .4 = .676$
		$\operatorname{var}(X) = 1.2$			$\overline{\operatorname{var}(Y) = 1.81}$

A computational formula for the variance

The variance of X can be computed by subtracting the square of the expected value of X from the expected value of the square of X. In symbols,

$$\operatorname{var}(X) = E(X^2) - [E(X)]^2.$$

Example 7.12 Two distributions to illustrate computations, revisited For the random variables X and Y of this example. We have E(X) = 3, E(Y) = 3.7, $E(X^2) = 10.2$, and $E(Y^2) = 10.5$. Thus $var(X) = 10.2 - (3)^2 = 1.2$ and $var(Y) = 15.5 - (3.7)^2 = 15.5 - 13.69 = 1.81$.

Some useful properties of expected values and variances

For these properties a and b denote constants, X and Y are discrete r.v.'s, and E(X), E(Y), and var(X) are assumed to exist.

(1) The expected value of a constant is that constant. In symbols,

$$E(a) = a.$$

(2) If X is bounded below by a (X is always greater than or equal to a or more formally X is greater than or equal to a with probability one), then the expected value of X is greater than or equal to a. In symbols,

if
$$P(X \ge a) = 1$$
, then $E(X) \ge a$.

(3) Similarly, if X is bounded above by b (X is always less than or equal to b or more formally X is less than or equal to b with probability one), then the expected value of X is less than or equal to b. In symbols, if

if
$$P(X \le b) = 1$$
, then $E(X) \le b$.

(4) Combining (2) and (3) we see that if X is always between a and b or more formally X takes on a value between a and b, inclusive, with probability one, then the expected value of X is between a and b, inclusive. In symbols,

if
$$P(a \le X \le b) = 1$$
, then $a \le E(X) \le b$.

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(5) If we transform the value of X by first multiplying X by a constant b and then adding a constant a, then the expected value of X is transformed in the same way. In symbols,

$$E(a+bX) = a+bE(X).$$

(6) The expected value of the sum of two (or more) random variables is equal to the sum of their expected values. In symbols,

$$E(X+Y) = E(X) + E(Y).$$

(7) We will now note what happens to the variance of X when we transform the value of X by first multiplying X by a constant b and then adding a constant a. Since variance is a measure of variability adding the constant a has no effect on the variance. Multiplying X by the constant b changes the scaling of the values of X, if |b| > 1 the values get more spread out on the number line, if |b| < 1 the values get closer together (less spread out) on the number line, and if b < 0 then the ordering of the values is reversed. Therefore, the transformation of X to a+bX has a multiplicative effect on the variance of X corresponding to a multiplication by the square of the constant b. In symbols,

$$\operatorname{var}(a+bX) = b^2 \operatorname{var}(X).$$

(8) The variance of a constant is zero,

$$\operatorname{var}(a) = 0.$$

7.9 Means, variances, and pmf's for several families of distributions toc

A family of distributions is a collection of distributions of a specified form which is known up to the values of one or more parameters. For the families summarized below, the p.m.f. is expressed as a function of one or more parameters and the family is obtained by varying the parameter or parameters over all suitable values.

Binomial distribution

The binomial distribution with parameters n (number of trials) and p (success probability).

Restrictions on the parameters: n is a positive integer (1, 2, 3, ...) and 0 .Probability mass function:

$$p_X(x) = \binom{n}{x} p^x (1-p)^{n-x}$$
 for $x = 0, 1, \dots, n$

Mean and variance:

$$E(X) = np$$
 and $var(X) = np(1-p)$

Hypergeometric distribution

The hypergeometric distribution with parameters A (number of units of the first type in the population), B (number of units of the second type in the population), and n (sample size).

Restrictions on the parameters: A, B, and n are positive integers (1, 2, 3, ...) with $n \le A + B$.

Probability mass function:

$$p_X(x) = \frac{\binom{A}{x}\binom{B}{n-x}}{\binom{A+B}{n}}$$
 for $x = 0, 1, \dots, n$ subject to the restrictions below.

Notice that the values of A and B may impose restrictions on the values of x for which this expression works. Specifically, we must have $x \leq A$ and $n - x \leq B$.

Mean and variance:

$$E(X) = n\left(\frac{A}{A+B}\right)$$
 and $\operatorname{var}(X) = n\left(\frac{A}{A+B}\right)\left(\frac{B}{A+B}\right)\left(\frac{A+B-n}{A+B-1}\right)$

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Geometric distribution

The geometric distribution with parameter p (success probability).

Restrictions on the parameter: 0 .

Probability mass function:

$$p_X(x) = (1-p)^{x-1}p$$
 for $x = 1, 2, 3, \dots$

Mean and variance:

$$E(X) = \frac{1}{p}$$
 and $\operatorname{var}(X) = \frac{1-p}{p^2}$

Uniform distribution

The discrete uniform distribution on the integers $1, 2, \ldots, N$.

Restrictions on the parameter: The parameter N is a positive integer (1, 2, 3, ...). **Probability mass function:**

$$p_X(x) = \frac{1}{N}$$
 for $x = 1, 2, \dots, N$

Mean and variance:

$$E(X) = \frac{N+1}{2}$$
 and $var(X) = \frac{N^2 - 1}{12}$

Poisson distribution

The Poisson distribution with parameter λ .

Restrictions on the parameter: $\lambda > 0$.

Probability mass function:

$$p_X(x) = \frac{\lambda^x}{x!} e^{-\lambda}$$
 for $x = 0, 1, 2, 3, \dots$

Mean and variance:

 $E(X) = \lambda$ and $var(X) = \lambda$

7.10 Problems

The problem sets and solutions can be found on the book web page at https://www.ucs.louisiana.edu/~jcb0773/.

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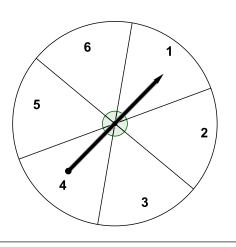
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8 Continuous random variables

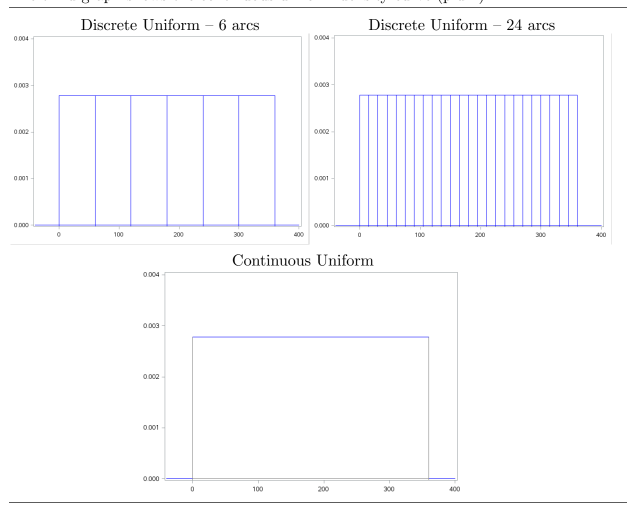
8.1 Moving from a discrete to a continuous random variable toc

A continuous random variable is a random variable X for which the sample space Ω_X is a (bounded or unbounded) interval of values on the number line. The term continuous here refers to the fact that when Ω_X is an interval the possible values of the random variable form a continuum. That is, there is a continuous transition from one possible value to the next in contrast to the jumps between values with a discrete transition. We will begin with a simple example showing how we can move from a discrete uniform distribution on six integers to a continuous uniform distribution on an interval.

Figure 8.1 A spinner on a disk with six equiangular regions



Consider a spinner atop a disk and an experiment consisting of spinning the pointer and noting its location on the circumference of the disk. In particular consider the spinner in Figure 8.1 where the circumference of the disk is divided into six equal length arcs. If we assume that the spinner is very well made and balanced, then it is reasonable to claim that the six values on the disk are equally probable. (If the pointer lands exactly on one of the boundaries, then we can simply spin again until the pointer lands inside a region.) Hence, under these assumptions, the discrete random variable X, denoting the number on the region where the pointer lands, follows the uniform distribution on the integers in the sample space $\Omega_X = \{1, 2, 3, 4, 5, 6\}$. **Figure 8.2** Uniform distributions on the interval from 0 to 360. The first two graphs show probability histograms for discrete uniform distributions on the midpoints of 6 and 24 equal length subintervals (arcs). The third graph shows the continuous uniform density curve (p.d.f.).



On the other hand, if we think of the circumference of the disk as a continuum, then we can represent an elementary outcome as a real number representing the angle of the radius to the point where the pointer stops. Let the points on the circumference of the disk be labeled in degrees starting with zero on the radius between 6 and 1 and moving clockwise and let Y denote the angle corresponding to the point where the pointer lands. With this convention the sample space is the interval from 0 to 360 degrees, *i.e.*, $\Omega_Y = [0, 360)$, with 0 < Y < 60 corresponding to X = 30, 60 < Y < 120 corresponding to X = 90, 120 < Y < 180 corresponding to X = 150, 180 < Y < 240 corresponding to X = 210, 240 < Y < 300 corresponding to X = 270, and 300 < Y < 360 corresponding to X = 330.

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If we think of these six arcs as six discrete outcomes corresponding to these six values of X, then we can label and scale the probability histogram of X so that it lies on the interval from 0 to 360 on the number line. As noted above, with this scaling the midpoints of the arcs 30, 90, 150, 210, 270, and 330 are the values of X. This probability histogram is shown in the first graph of Figure 8.2. Each of the rectangles in the probability histogram has height 1/360.

It seems natural to argue that the continuous random variable Y should be uniformly distributed on the sample space $\Omega_Y = [0, 360)$. More formally, it seems natural to require that the probability that Y takes on a value within any arc of length θ ($0 < \theta < 360$) is equal to the length of the arc divided by 360, *i.e.*, for $0 < \theta < 360$ and any arc of length θ , the probability that Y belongs to this arc (that the pointer stops within this arc) is $\frac{\theta}{360}$. As shown in Figure 8.2, the probability histogram for this continuous uniform distribution on $\Omega_Y = [0, 360)$ is a single rectangle of height 1/360 located over the interval from 0 to 360.

The probability histogram for the discrete uniform distribution on 24 arcs in Figure 8.2 is provided to indicate how we can think of the continuous uniform probability histogram as the limiting version of the discrete uniform probability histogram (on this same interval) which we would obtain if we made the rectangles narrower and narrower. That is, to indicate that by increasing the number of arcs that the circle (interval) is partitioned into the histogram of the discrete uniform distribution would approach the histogram of the continuous uniform distribution.

Notice that with this continuous uniform distribution on the circumference of the disk the probability that the pointer lands at a specified point is zero but the probability that it lands in a specified arc, of length θ , which contains the point is $\frac{\theta}{360} > 0$. Notice also that that if the length of the arc containing the point is made shorter and shorter, then the arc degenerates to the point and the probability $\frac{\theta}{360}$ approaches zero; thus, the notion of a point having probability zero is consistent with this assignment of probability to an arc.

8.2 Continuous random variables

We will now consider probability models for the distribution of a continuous random variable. Recall that a continuous random variable is a random variable X for which the sample space Ω_X is a (bounded or unbounded) interval of values on the number line. In

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Section 8.1 we showed how a continuous uniform distribution on an interval arises as a sort of limiting version of a discrete uniform distribution on the same interval obtained by letting the number of possible values increase without bound. If we modified our division of the disk by allowing the lengths of the arcs to vary, our limiting argument would lead to a non–uniform continuous distribution. We will now consider continuous random variables more generally.

As noted above, the sample space of a continuous random variable is a (bounded or unbounded) interval of values on the number line. For simplicity, the sample space is often the entire number line or the nonnegative part of the number line. We can characterize the distribution of the continuous r.v. X by specifying a **probability density function** (denoted p.d.f.) f_X . We will emphasize the graph of the p.d.f. and we will also refer to the p.d.f. as a density curve. A probability density function f_X is a nonnegative valued function with the property that the area under the graph of the function (the density curve) is one. That is, the area between the x-axis and the density curve is one. We can think of the density curve as a smooth version of a probability histogram with the rectangles of the histogram replaced by a smooth curve indicating where the tops of the rectangles would be.

As noted above, the area under a density curve (p.d.f.) is one. The probability of an event is equal to the area under the density curve over the interval corresponding to the event. Thus, for any event A defined in terms of the continuous r.v. X with p.d.f f_X , the probability of event A is equal to the area under the graph of f_X over the region A. For example, if the event A is an interval, say A = (a, b), then the probability that X belongs to A, P(a < X < b) is equal to the area under the p.d.f. f_X (under the density curve) over the interval (a, b). We can use a computer or a calculator to find such probabilities (areas).

This identification of the probability of an event with an area under a density curve is illustrated in Figures 8.3 and 8.4. In these figures, X_1 is a continuous random variable with a symmetric distribution and X_2 is a continuous random variable with a skewed right distribution. In Figure 8.3 the probabilities are of the form $P(X \le a)$ and in Figure 8.4 the probabilities are of the form $P(a \le X \le b)$.

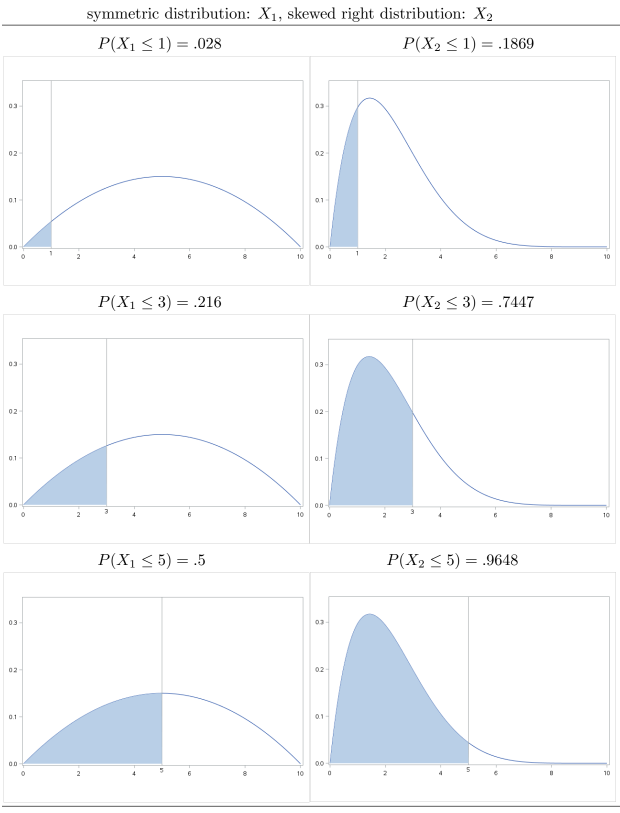


Figure 8.3 Continuous distributions – probability as area

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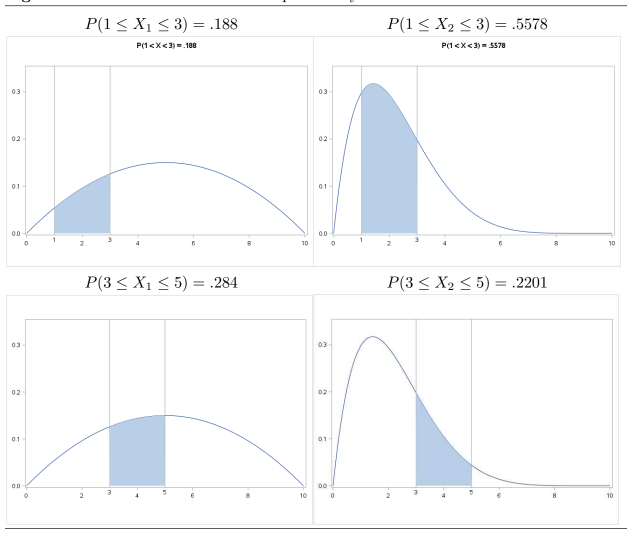


Figure 8.4 Continuous distributions – probability as area

Notice that $f_X(x)$ is not a probability! With a continuous random variable only events which can be represented as non-degenerate intervals (or unions of non-degenerate intervals) can have positive probability, since probability is defined as the area under the density curve and we need at least one non-degenerate interval to obtain a positive area.

Given a probability model for the distribution of a continuous random variable X, *i.e.*, given a density curve for the distribution of X, we can define population parameters which characterize relevant aspects of the distribution. For example, we can define the population mean μ as the balance point of the unit mass bounded by the density curve and the number line. We can also think of the population mean as the weighted average of the infinite collection of possible values of X with weights determined by the density

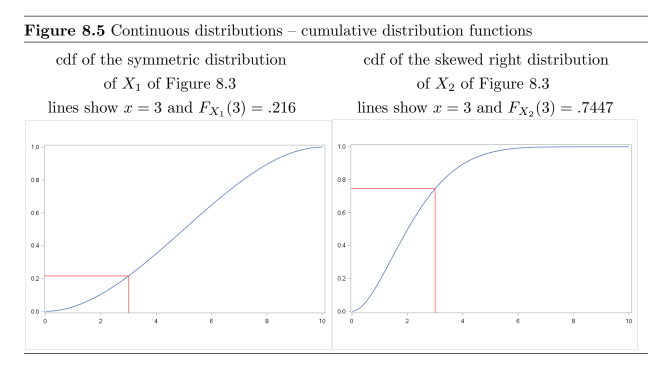
curve. We can similarly define the population median M as the point on the number line where a vertical line would divide the area under the density curve into two equal areas (each of size one-half).

The distribution of X can also be characterized in terms of its **cumulative distribution** function (denoted c.d.f.) F_X , the c.d.f. gives the probability of events of the form $[X \leq x]$, thus

$$F_X(x) = P(X \le x).$$

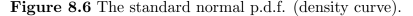
For a continuous random variable the value of the c.d.f. $F_X(x)$ is the area under the density curve over the interval which ranges from minus infinity to x. As noted earlier, we can use a computer or a calculator to determine such areas (integrals).

The cumulative distribution functions corresponding to the probability density functions of Figure 8.3 are provided in Figure 8.5. In this figure lines at x = 3 and $F_X(3)$ are included to show how the c.d.f yields the value of the corresponding area under the p.d.f over the interval (0,3) as shown in Figure 8.3.



8.3 The normal distribution

The most widely used continuous probability model is the normal probability model or normal distribution. The normal distribution with mean μ and standard deviation σ can be characterized by its density curve. The density curve for the normal distribution with mean μ and standard deviation σ is the familiar bell shaped curve. The standard normal density curve, which has mean $\mu = 0$ and standard deviation $\sigma = 1$, is shown in Figure 8.6. The standard normal cumulative distribution function is shown in Figure 8.7.



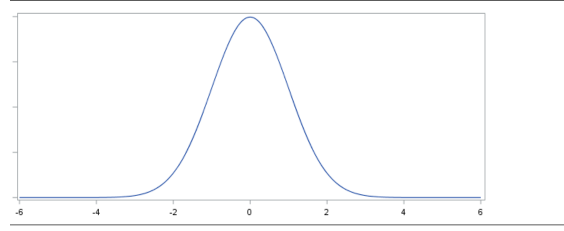
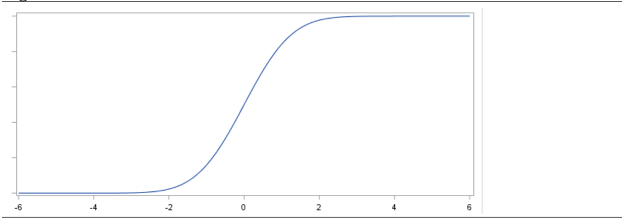


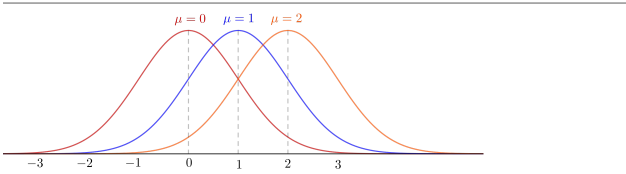
Figure 8.7 The standard normal c.d.f.



The normal distribution with mean μ and its density curve are symmetric around μ , *i.e.*, if we draw a vertical line through μ , then the two sides of the density curve are mirror images of each other. Therefore the mean of a normal distribution μ is also the median of the normal distribution. The mean μ locates the normal distribution on the number line so that if we hold σ constant and change the mean μ , the normal distribution is simply shifted

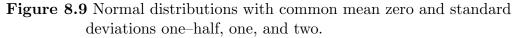
along the number line until it is centered at the new mean. In other words, holding σ fixed and changing μ simply relocates the density curve on the number line; it has no effect on the shape of the curve. Figure 8.8 provides the density curves for normal distributions with respective means $\mu = 0$, $\mu = 1$, and $\mu = 2$ and common standard deviation $\sigma = 1$.

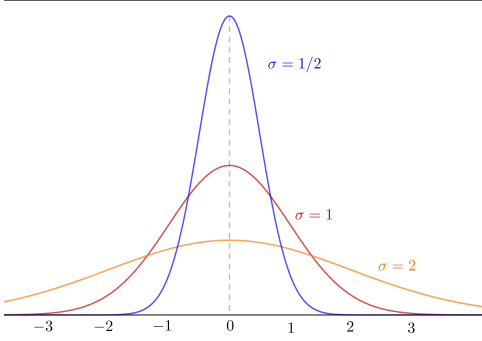
Figure 8.8 Normal distributions with common standard deviation one and means of zero, one, and two.



The standard deviation σ indicates the amount of variability in the normal distribution. If we hold μ fixed and increase the value of σ , then the normal density curve becomes flatter, while retaining its bell–shape, indicating that there is more variability in the distribution. Similarly, if we hold μ fixed and decrease the value of σ , then the normal density curve becomes more peaked around the mean μ , while retaining its bell–shape, indicating that there is less variability in the distribution. Normal distributions with mean $\mu = 0$ and respective standard deviations $\sigma = 1/2$, $\sigma = 1$, and $\sigma = 2$ are plotted in Figure 8.9.

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Computer programs and calculators can be used to compute normal probabilities or equivalently to compute areas under the normal density curve. These probabilities can also be calculated using tables of standard normal distribution probabilities such as Table 8.12. To use this table or a calculator which only handles the standard normal distribution, you need re-express probability statements in terms of a standard normal random variable. The relationship between the standard normal random variable Z and the general normal random variable X, when X has mean $E(X) = \mu$ and standard deviation σ , is

$$Z = \frac{X - \mu}{\sigma}$$
 or equivalently $X = \mu + Z\sigma$

This relationship implies that a probability statement about the normal variable X can be re-expressed as a probability statement about the standard normal variable Z by reexpressing the statement in terms of standard deviation units from the mean. Given two constants a < b, observing a value of X between a and b (observing $a \le X \le b$) is equivalent to observing a value of $Z = (X - \mu)/\sigma$ between $(a - \mu)/\sigma$ and $(b - \mu)/\sigma$ (observing $(a - \mu)/\sigma \le (X - \mu)/\sigma \le (b - \mu)/\sigma$). Furthermore, $Z = (X - \mu)/\sigma$ behaves in accordance with the standard normal distribution so that the probability of observing a value of X between a and b, denoted by $P(a \le X \le b)$, is equal to the probability that the standard normal variable Z takes on a value between $(a - \mu)/\sigma$ and $(b - \mu)/\sigma$, *i.e.*,

$$P(a \le X \le b) = P\left(\frac{a-\mu}{\sigma} \le Z \le \frac{b-\mu}{\sigma}\right).$$

In terms of areas this probability equality says that the area under the normal density curve with mean μ and standard deviation σ over the interval from a to b is equal to the area under the standard normal density curve over the interval from $(a-\mu)/\sigma$ to $(b-\mu)/\sigma$. Similarly, given constants c < d, we have the analogous result that

$$P(c \le Z \le d) = P(\mu + c\sigma \le X \le \mu + d\sigma).$$

Table 8.12 provides cumulative standard normal probabilities of the form $P(Z \le a)$ for values of a (Z in the table) between 0 and 3.69. Computer programs usually produce cumulative probabilities like these. To use these cumulative probabilities to compute a probability of the form $P(a \le Z \le b)$ note that

$$P(a \le Z \le b) = P(Z \le b) - P(Z \le a)$$

and note that the symmetry of the normal distribution implies that

$$P(Z \le -a) = P(Z \ge a) = 1 - P(Z \le a).$$

Calculators often provide probabilities of the form $P(a \le Z \le b)$ directly.

Example 8.1 NHANES The National Health and Nutrition Examination Survey (NHANES) We will use some data from the 2013–2014 NHANES (see Example 1.1) to illustrate some cases where the normal distribution does and does not provide a suitable model for the distribution of a variable. We will consider the distributions of three variables: age, height, and weight. All available data for the 5588 adults (age 20 and over) in the 2013–2014 NHANES are used. For females the samples sizes are: n = 2919 for age and n = 2888 for height and weight. For males we have: n = 2669 for age, n = 2642 for height, and n = 2645 for weight. The data for the females is summarized graphically in Figure 8.10 and that for the males in Figure 8.11. Two density curves are superimposed on the histograms in the left column of these figures. There is a normal density curve, with μ and σ

154 8.3 The normal distribution

chosen to match the data, and there is a "kernel" density curve which is basically a smooth version of the histogram. It is somewhat tricky to properly compare distributions based on histograms or density curves. Cumulative distribution functions, and associated graphs and statistics, form a better basis for such comparisons. The graphs in the right column contain an empirical (sample) cumulative distribution function and the cumulative distribution function for the fitted normal distribution. The empirical c.d.f is a "step function" based on the observed data values. The "steps" in this function are most pronounced in the graphs for the age distributions. The fact that the empirical (sample) and theoretical (normal) c.d.f.'s for the height distributions (females and males) are essentially indistinguishable indicates that the normal distribution provides a reasonable model for the height distribution. The weight distributions (females and males) are not modeled very well by a normal distribution. For both sexes the weight distributions are too strongly skewed right for the normal model to fit well. The age distributions are clearly non-normal. Actually, the reason that I included these age distributions is so that you could more easily see the appearance (steps) of an empirical c.d.f.

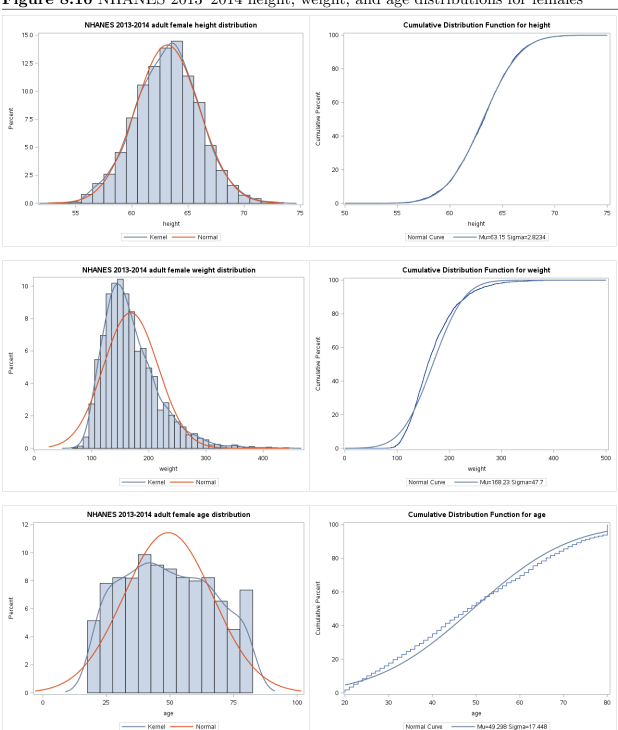


Figure 8.10 NHANES 2013–2014 height, weight, and age distributions for females

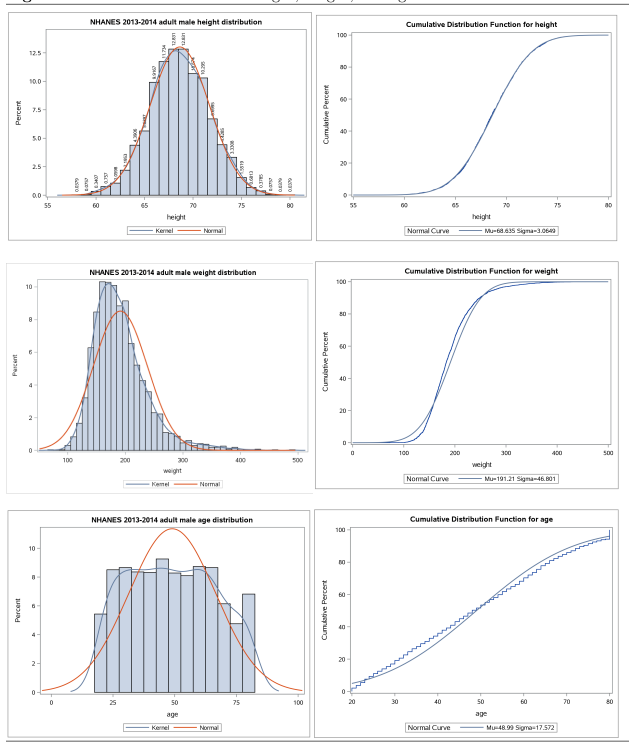


Figure 8.11 NHANES 2013–2014 height, weight, and age distributions for males

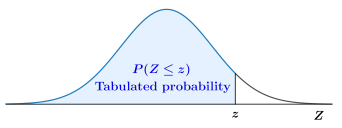


Table 8.12Cumulative normal probabilities.	
(Areas under the standard normal curve to the left of Z .)	

										/
Z	Second decimal place in Z									
	.00	.01	.02	.03	.04	.05	.06	.07	.08	.09
0.0	.5000	.5040	.5080	.5120	.5160	.5199	.5239	.5279	.5319	.5359
0.1	.5398	.5438	.5478	.5517	.5557	.5596	.5636	.5675	.5714	.5753
0.2	.5793	.5832	.5871	.5910	.5948	.5987	.6026	.6064	.6103	.6141
0.3	.6179	.6217	.6255	.6293	.6331	.6368	.6406	.6443	.6480	.6517
0.4	.6554	.6591	.6628	.6664	.6700	.6736	.6772	.6808	.6844	.6879
0.5	.6915	.6950	.6985	.7019	.7054	.7088	.7123	.7157	.7190	.7224
0.6	.7257	.7291	.7324	.7357	.7389	.7422	.7454	.7486	.7517	.7549
0.7	.7580	.7611	.7642	.7673	.7704	.7734	.7764	.7794	.7823	.7852
0.8	.7881	.7910	.7939	.7967	.7995	.8023	.8051	.8078	.8106	.8133
0.9	.8159	.8186	.8212	.8238	.8264	.8289	.8315	.8340	.8365	.8389
1.0	.8413	.8438	.8461	.8485	.8508	.8531	.8554	.8577	.8599	.8621
1.1	.8643	.8665	.8686	.8708	.8729	.8749	.8770	.8790	.8810	.8830
1.2	.8849	.8869	.8888	.8907	.8925	.8944	.8962	.8980	.8997	.9015
1.3	.9032	.9049	.9066	.9082	.9099	.9115	.9131	.9147	.9162	.9177
1.4	.9192	.9207	.9222	.9236	.9251	.9265	.9279	.9292	.9306	.9319
1.5	.9332	.9345	.9357	.9370	.9382	.9394	.9406	.9418	.9429	.9441
1.6	.9452	.9463	.9474	.9484	.9495	.9505	.9515	.9525	.9535	.9545
1.7	.9554	.9564	.9573	.9582	.9591	.9599	.9608	.9616	.9625	.9633
1.8	.9641	.9649	.9656	.9664	.9671	.9678	.9686	.9693	.9699	.9706
1.9	.9713	.9719	.9726	.9732	.9738	.9744	.9750	.9756	.9761	.9767
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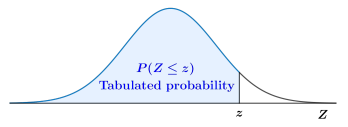


Table 8.12 (continuation)Cumulative normal probabilities.(Areas under the standard normal curve to the left of Z.)

-										,
Z	Second decimal place in Z									
	.00	.01	.02	.03	.04	.05	.06	.07	.08	.09
2.0	.9772	.9778	.9783	.9788	.9793	.9798	.9803	.9808	.9812	.9817
2.1	.9821	.9826	.9830	.9834	.9838	.9842	.9846	.9850	.9854	.9857
2.2	.9861	.9864	.9868	.9871	.9875	.9878	.9881	.9884	.9887	.9890
2.3	.9893	.9896	.9898	.9901	.9904	.9906	.9909	.9911	.9913	.9916
2.4	.9918	.9920	.9922	.9925	.9927	.9929	.9931	.9932	.9934	.9936
2.5	.9938	.9940	.9941	.9943	.9945	.9946	.9948	.9949	.9951	.9952
2.6	.9953	.9955	.9956	.9957	.9959	.9960	.9961	.9962	.9963	.9964
2.7	.9965	.9966	.9967	.9968	.9969	.9970	.9971	.9972	.9973	.9974
2.8	.9974	.9975	.9976	.9977	.9977	.9978	.9979	.9979	.9980	.9981
2.9	.9981	.9982	.9982	.9983	.9984	.9984	.9985	.9985	.9986	.9986
3.0	.9987	.9987	.9987	.9988	.9988	.9989	.9989	.9989	.9990	.9990
3.1	.9990	.9991	.9991	.9991	.9992	.9992	.9992	.9992	.9993	.9993
3.2	.9993	.9993	.9994	.9994	.9994	.9994	.9994	.9995	.9995	.9995
3.3	.9995	.9995	.9995	.9996	.9996	.9996	.9996	.9996	.9996	.9997
3.4	.9997	.9997	.9997	.9997	.9997	.9997	.9997	.9997	.9997	.9998
3.5	.9998	.9998	.9998	.9998	.9998	.9998	.9998	.9998	.9998	.9998
3.6	.9998	.9998	.9999	.9999	.9999	.9999	.9999	.9999	.9999	.9999

8.4 The exponential distribution

Exponential distributions are often used to model the time until the occurrence of an event. For example, we might use an exponential distribution to model the amount of time (starting from now) until an earthquake occurs or until a machine malfunctions.

The exponential distribution is defined for nonnegative values, that is, the sample space for an exponential random variable is the nonnegative part of the *x*-axis. An exponential distribution is determined by the value of a single parameter $\lambda > 0$. The p.d.f. of the exponential distribution with parameter $\lambda > 0$ is given by

$$f_X(x) = \lambda e^{-\lambda x}$$

for $x \ge 0$. If X denotes a random variable which follows this exponential distribution with parameter λ , then $E(X) = 1/\lambda$ and $\operatorname{var}(X) = 1/\lambda^2$. The p.d.f's of exponential distributions with respective means 1/2, 1, and 2 (respective parameters $\lambda = 2$, $\lambda = 1$, and $\lambda = 1/2$) are graphed in Figure 8.13. Figure 8.14 shows the p.d.f and c.d.f of the exponential distribution with mean one.

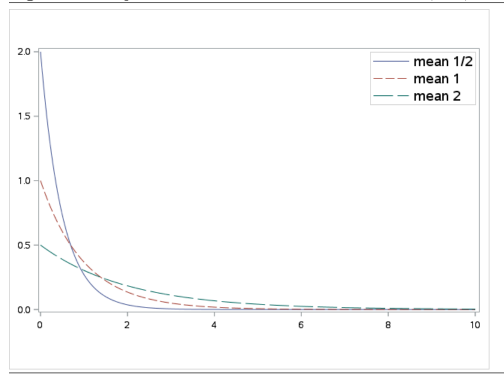


Figure 8.13 Exponential distributions with means one-half, one, and two.

160 8.4 The exponential distribution

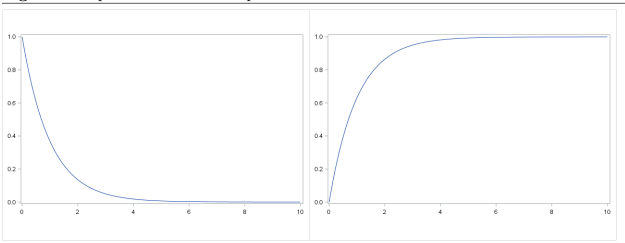


Figure 8.14 p.d.f. and c.d.f. of exponential distribution with mean one

There is an important connect between the Poisson distribution and the exponential distribution. As noted earlier, exponential distributions are often used to model the time until the occurrence of an event. In Section 7.6 we listed three assumptions about the times of occurrence of events which justify the use of the Poisson distribution to model a count of the number of occurrences of an event. If the events of interest behave in accordance with these assumptions, then the time until the occurrence of an event and the time between any two successive events will follow an exponential distribution.

8.5 Problems

toc

The problem sets and solutions can be found on the book web page at https://www.ucs.louisiana.edu/~jcb0773/.

9 Inference for a Proportion

9.1 Introduction

A dichotomous population is a collection of units which can be divided into two nonoverlapping subcollections corresponding to the two possible values of a dichotomous variable, e.q. male or female, dead or alive, pass or fail. It is conventional to refer to one of the two possible values which dichotomize the population as "success" and the other as "failure." These generic labels are not meant to imply that success is good. Rather, we can think of choosing one of the two possible classifications and asking "does the unit belong to the subcollection of units with this classification?" with the two possibilities being yes (a success) and no (a failure). When a unit is selected from the population and the unit is found to belong to the success subgroup we say that a **success** has occurred. Similarly, when a member of the failure subgroup is selected we say that a **failure** has occurred. The proportion of units in the population that belong to the success subgroup (the units classified as successes) is the **population success proportion**. This population success proportion is denoted by the lower case letter p. The population success proportion p is a parameter, since it is a numerical characteristic of the population. The sample success **proportion** or observed proportion of successes in a sample from a dichotomous population is denoted by \hat{p} (read this as p hat). The observed success proportion \hat{p} is a statistic, since it is a numerical characteristic of the sample.

We can use the selection of balls from a box of balls as a model for sampling from a dichotomous population. Consider a box containing balls of which some are red (successes) and the rest are blue (failures). The population success proportion, p, is the proportion of red balls in the box. If we select a random sample of n balls from this box, then the sample success proportion, \hat{p} , is the proportion of red balls in the sample. Thus, in this model, a ball is a unit, the box of balls is the population, selecting a red ball is a success, the proportion of red balls in the box p is the parameter (the population success proportion), and the proportion of red balls in the sample \hat{p} is the statistic (the sample success proportion).

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9.2 The sampling distribution and the normal approximation

We will now discuss how the sample proportion \hat{p} can be used to make inferences about the population proportion p. Assuming that the sample size n is reasonably large, it seems reasonable to view the sample proportion \hat{p} as an estimate of the population proportion p. Clearly there will be some variability from sample to sample in the computed values of the statistic \hat{p} . That is, if we took several random samples from the population, we would not expect the observed sample success proportions, the \hat{p} 's, to be exactly the same. In the box of balls example, if we took several samples from the box we would expect the proportion of red balls in the sample to vary from sample to sample.

Two questions we might ask about the sample proportion \hat{p} as an estimator of the population proportion p are:

- (1) Can we expect the sample proportion \hat{p} to be close to the population proportion p?
- (2) Can we quantify how close \hat{p} will be to p?

The sampling distribution of \hat{p} , which describes the sample to sample variability in \hat{p} , can be used to address these questions.

In general, the **sampling distribution of a statistic** is the distribution of the possible values of the statistic that could be obtained from random samples. We can think of the sampling distribution of a statistic as a theoretical relative frequency distribution for the possible values of the statistic which describes the sample to sample variability in the statistic. The form of the sampling distribution of a statistic depends on the nature of the population the sample is taken from, the size of the sample, and the method used to select the sample.

For example, suppose that we select a simple random sample of n = 10 balls, with replacement, from a box containing six red balls and four blue balls. If we identify red as a success, then, in this example, the population success proportion is p = .6, since 60% of the balls in the box are red. The possible values of the sample success proportion \hat{p} when n = 10 are: 0, .1, .2, ..., 1. The sampling distribution of \hat{p} summarized in Figure 9.1 gives the probabilities corresponding to these possible values. As you would expect, the probabilities are highest for values near p = .6 and very small for values far away from p = .6. In particular, when we select a simple random sample of size n = 10 from this box, 25.08% of the time we would observe $\hat{p} = .6$, 20.07% of the time we would observe $\hat{p} = .5$, and 21.50% of the time we would observe $\hat{p} = .7$. On the other hand, it would be possible but very unlikely to observe $\hat{p} = .2$ (only 1.06% of the time) or $\hat{p} = .9$ (only 4.03% of the time).

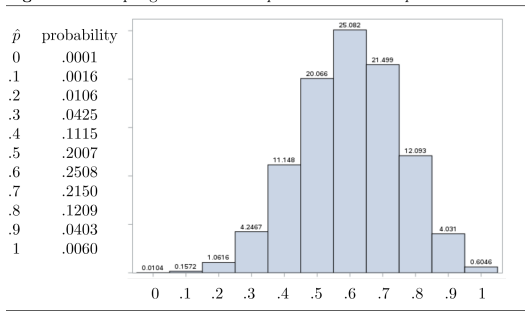


Figure 9.1 Sampling distribution of \hat{p} when n = 10 and p = .6

The mean and the standard deviation of the sampling distribution are of particular interest. The mean of the sampling distribution indicates whether the statistic is biased as an estimator of the parameter of interest. If the mean of the sampling distribution is equal to the parameter of interest, then the statistic is said to be **unbiased** as an estimator of the parameter. Otherwise, the statistic is said to be **biased** as an estimator of the parameter. To say that a statistic is **unbiased** means that, even though the statistic will overestimate the parameter for some samples and will underestimate the parameter for other samples, it will do so in such a way that, in the long run, the values of the statistic will average to give the correct value of the parameter. When the statistic is **biased** the statistic will tend to consistently overestimate or consistently underestimate the parameter; therefore, in the long run, the values of a biased statistic will not average to give the correct value of the parameter. The standard deviation of the sampling distribution is known as the standard error of the statistic. The standard error of the statistic provides a measure of the sample to sample variability in the values of the statistic. The standard error of the statistic can be used to quantify how close we can expect the value of the statistic to be to the value of the parameter.

164 9.2 The sampling distribution and the normal approximation

Returning to our discussion of the sampling distribution of \hat{p} we first present two important properties of this sampling distribution. The observed proportion \hat{p} in a simple random sample selected with replacement from a population with population proportion p has a sampling distribution with the following properties.

1. The mean of the sampling distribution of \hat{p} is the population probability p. Therefore, \hat{p} is unbiased as an estimator of p.

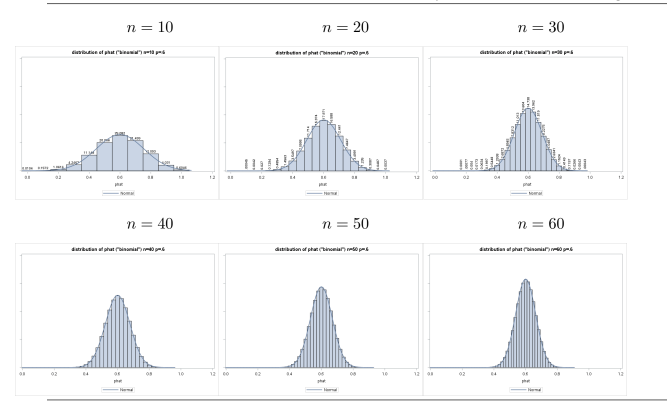
2. The population standard error of \hat{p} , denoted by $SE(\hat{p})$, is

$$\operatorname{SE}(\hat{p}) = \sqrt{\frac{p(1-p)}{n}}.$$

The inferential methods we will consider are based on a large sample size normal approximation to the sampling distribution of \hat{p} . Before we discuss the normal approximation to the sampling distribution of \hat{p} , we will provide an indication of some basic properties of the sampling distribution of \hat{p} in terms of some graphical representations of the probability histogram of the distribution of \hat{p} with superimposed fitted normal density curves.

First consider how the sampling distribution of \hat{p} depends on the sample size n. From the expression given above for the population standard error of \hat{p} we can see that, as you would expect, the variability in \hat{p} as an estimator of p decreases as the sample size increases. This behavior of the sampling distribution is illustrated in Figure 9.2.

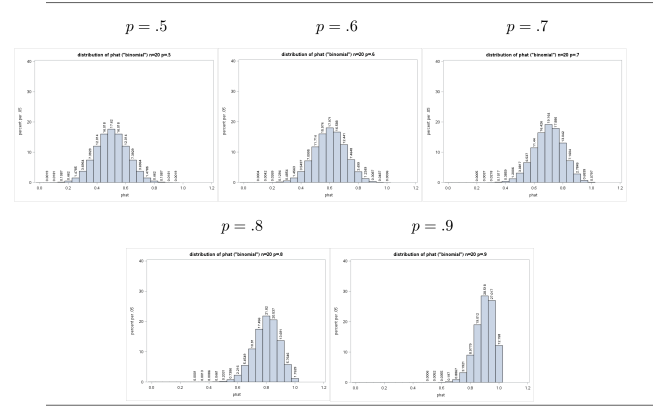
Figure 9.2 The sampling distribution of \hat{p} with normal approximation for p = .6. Histograms are provided for n = 10, 20, 30, 40, 50, and 60. All of these distributions are centered at p = .6. Notice how the variability in the distribution decreases as the sample size n increases and how much better the normal density curve matches the histogram.



Next consider how the sampling distribution of \hat{p} depends on the value of the population success proportion p. The sampling distribution of \hat{p} is unimodal (single peaked) with its peak at p. If p = .5, then the sampling distribution of \hat{p} is symmetric. If p < .5, then sampling distribution of \hat{p} is skewed right. If p > .5, then sampling distribution of \hat{p} is skewed left. Since \hat{p} is unbiased as an estimator of p, the mean of the distribution of \hat{p} is p. Thus, the probability histogram of the distribution of \hat{p} has its balance point at p. From the expression given above for the population standard error of \hat{p} we can see that, for fixed n, the variability in \hat{p} as an estimator of p is highest when p = .5 and decreases as p moves away from .5. These properties of the sampling distribution are illustrated in Figure 9.3.

166 9.2 The sampling distribution and the normal approximation

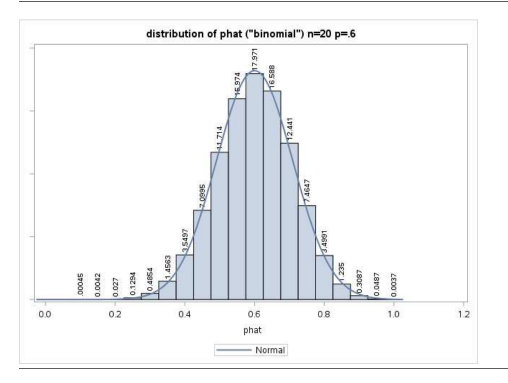
Figure 9.3 The sampling distribution of \hat{p} with normal approximation for n = 20. Histograms are provided for p = .5, .6, .7, .8, and .9. Each distribution is centered at its p. Notice how the variability in the distribution decreases as p moves away from .5. Notice also how the distribution becomes more skewed left as p moves away from .5.



The normal approximation to the sampling distribution of \hat{p} is illustrated graphically in Figure 9.4. In this figure, the probability histogram of the sampling distribution of \hat{p} for the situation when a simple random sample of size n = 20 is selected with replacement from a dichotomous population with p = .6 is given along with the approximating normal density curve. The exact probabilities which correspond to the areas of the rectangles of the histogram are given as percentages, *e.g.*, the probability of observing $\hat{p} = .06$ is .17971 (17.97%). The normal approximation, explained more fully below, basically says that, over a specified range of \hat{p} values, the area under the normal density curve and the area in the rectangles of the histogram are similar. You can see that the curve matches the histogram reasonably well in Figure 9.4.

toc

Figure 9.4 The sampling distribution of \hat{p} with normal approximation for n = 20 and p = .6. This assumes a simple random sample selected with replacement. If the \hat{p} values were converted to counts, $X = n\hat{p}$, then this would be a binomial distribution.



The normal approximation to the sampling distribution of \hat{p} says that, for large values of n, the standardized value of \hat{p} obtained by subtracting the population proportion pfrom \hat{p} and dividing this difference by the population standard error of \hat{p} , behaves in approximate accordance with the standard normal distribution. That is, for large values of n the quantity $Z = (\hat{p} - p)/\text{SE}(\hat{p})$ behaves like a standard normal variable. This means, as will be shown below, that we can use an area under the standard normal density curve (a probability in terms of Z) to approximate an area in the probability histogram of the sampling distribution of \hat{p} (a probability in terms of \hat{p}). The relationship between \hat{p} and pindicated by this expression for Z and the normal distribution itself allow us to use \hat{p} to make formal, quantifiable inferences about p.

9.3 Estimation of p

As noted above, the sample proportion \hat{p} is an unbiased estimator of the population proportion p. We can think of \hat{p} as our "best guess" of the value of p. To allow for sampling variability it would be more useful to report a range or interval of plausible values for p.

168 9.3 Estimation of p

In particular, given the data we would like to be able to say, with a reasonable level of confidence, that the true value of p is between two particular limiting values. We will now use the normal approximation to the sampling distribution of \hat{p} to develop such an interval estimate of p.

The probability that a standard normal variable Z takes on a value between -1.96 and 1.96 is equal to .95, *i.e.*, $P(-1.96 \le Z \le 1.96) = .95$. Thus, when we observe the value of a standard normal variable Z, 95% of the time we will find that $-1.96 \le Z \le 1.96$. Graphically this means that the area under the standard normal density curve over the interval from -1.96 to 1.96 is .95. Thus, for sufficiently large values of n we have the approximation,

$$P\left[-1.96 \le \frac{\hat{p} - p}{\mathrm{SE}(\hat{p})} \le 1.96\right] = .95$$

or equivalently

 $P[p - 1.96 \cdot \text{SE}(\hat{p}) \le \hat{p} \le p + 1.96 \cdot \text{SE}(\hat{p})] = .95.$

Note that this indicates that 95% of the time when a simple random sample is selected and \hat{p} is computed the observed value of \hat{p} will be between $p - 1.96 \cdot \text{SE}(\hat{p})$ and $p + 1.96 \cdot \text{SE}(\hat{p})$, *i.e.*, \hat{p} will be within 1.96 population standard error units of p. We will refer to the interval from $p - 1.96 \cdot \text{SE}(\hat{p})$ to $p + 1.96 \cdot \text{SE}(\hat{p})$ as the central 95% interval of the distribution of \hat{p} , since it is centered at p and it will contain the observed value of \hat{p} 95% of the time.

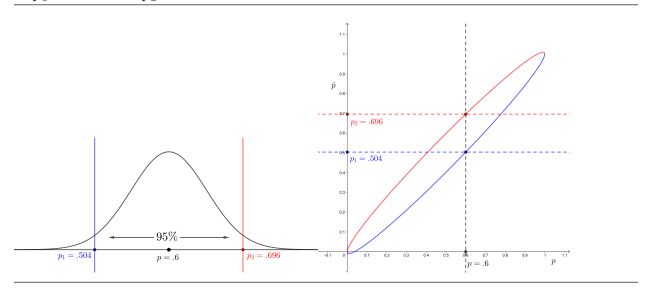
The relationship between \hat{p} and p is illustrated, for n = 100 and p = .6, in Figure 9.5. In this case, p = .6, $SE(\hat{p}) = \sqrt{.6(.4)/100} = .0490$, and $p_1 = .6 - 1.96 \cdot SE(\hat{p}) = .5040$ and $p_2 = .6 + 1.96 \cdot SE(\hat{p}) = .6960$ are the limits of the central 95% interval of the distribution of \hat{p} . The shaded region with area .95 in Figure 9.5 indicates that 95% of all samples will yield a sample proportion \hat{p} which is between p_1 and p_2 . That is, when p = .6 and n = 100, 95% of all samples will yield a sample proportion \hat{p} which is between p_1 and p_2 . That is, between $p_1 = .5040$ and $p_2 = .6960$. In terms of a box of balls this means that, if exactly 60% of the balls in the box were red, then 95% of the time when we selected n = 100 balls from the box, at random and with replacement, we would find between 51 and 69 red balls (between 50.40 and 69.60) among the 100 balls in the sample.

The plot on the right in Figure 9.5 shows how the endpoints of the central 95% interval of the distribution of \hat{p} depend on p. In this plot the sample size is n = 100. (The pattern is similar for other values of n.) The upper (red) curve gives values of $p + 1.96 \cdot \text{SE}(\hat{p}) = p + 1.96 \sqrt{p(1-p)/n}$ as a function of p and the lower (blue) curve gives values of

 $p - 1.96 \cdot \text{SE}(\hat{p}) = p - 1.96 \sqrt{p(1-p)/n}$ as a function of p. The intersections of a vertical line draw at a particular value of p with these curves are the endpoints of the central 95% interval of the distribution of \hat{p} for that value of p. The lines drawn in the figure demonstrate this for the case p = .6. Note that, as mentioned above, $p_1 = .5040$ and $p_2 = .6960$, and these are the endpoints of the interval in the plot on the left in Figure 9.5.

Figure 9.5 The plot on the left shows the central 95% interval of the distribution of \hat{p} for n = 100 and p = .6.

The curves in the plot on the right show the endpoints, $p \pm 1.96\sqrt{p(1-p)/n}$, of the central 95% interval of the distribution of \hat{p} as a function of p for n = 100. The endpoints for the case p = .6 are indicated by the lines marking the intersections at $p_1 = .5040$ and $p_2 = .6960$.



A confidence interval for p

As noted above, for a particular value of p, 95% of all samples will yield a value of \hat{p} within the corresponding central 95% interval from $p - 1.96\text{SE}(\hat{p})$ to $p + 1.96\text{SE}(\hat{p})$. Thus for each possible value of p we can find an interval of likely values for \hat{p} . But we need an interval of plausible values for p not for \hat{p} ! We will now show how the central 95% intervals of the distribution of \hat{p} can be used to form a 95% confidence interval estimate of p. We will provide a formal definition of a 95% confidence interval estimate later.

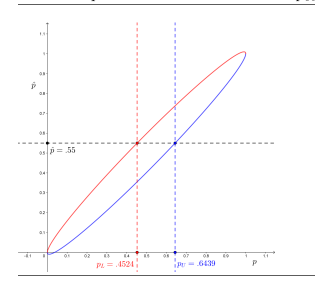
Suppose that a simple random sample has been selected and let \hat{p}_{obs} denote the observed value of \hat{p} . Given this \hat{p}_{obs} , we want to know which values of p are plausible. More precisely, we want to know which values of p determine sampling distributions for \hat{p} under which

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seeing $\hat{p} = \hat{p}_{obs}$ would not be surprising. We can formalize this goal by saying that we want to know which values of p determine a sampling distributions for \hat{p} for which the central 95% interval of the distribution of \hat{p} contains the observed value \hat{p}_{obs} .

The plot on the right in Figure 9.5 shows how the central 95% interval of the distribution of \hat{p} depends on the value of p. We want to determine which values of p yield central 95% intervals which contain \hat{p}_{obs} . This requires using the graph of Figure 9.5 in the other direction. In Figure 9.6 a horizontal line is drawn at \hat{p}_{obs} and its intersections, p_L and p_U , with the two curves are indicated. Notice that p_L is the smallest value of p for which the central 95% interval of the distribution of \hat{p} contains \hat{p}_{obs} and p_U is the largest value of p for which the central 95% interval of the distribution of \hat{p} contains \hat{p}_{obs} . (Figure 9.6 is drawn for n = 100 and $\hat{p}_{obs} = .55$. In this case, $p_L = .4524$ and $p_U = .6439$.) Thus, if we draw a vertical line, as in Figure 9.5, at any value of p between p_L and p_U , then the corresponding central 95% interval of the distribution of \hat{p} contains \hat{p}_{obs} .

Figure 9.6 The smallest and largest values of p, p_L and p_U , for which the central 95% interval of the distribution of \hat{p} contains \hat{p}_{obs} . This example is drawn for n = 100 and $\hat{p}_{obs} = .55$.



In order to determine the value of p_L we simply set $p + 1.96\text{SE}(\hat{p})$ equal to \hat{p}_{obs} and solve for p (draw the horizontal line as in Figure 6 and project down at the intersection with the upper (red) curve). To determine the value of p_U we set $p - 1.96\text{SE}(\hat{p})$ equal to \hat{p}_{obs} and solve for p (draw the horizontal line as in Figure 9.6 and project down at the intersection with the lower (blue) curve).

	control (507 interval
~		05% interval
$p \\ .10$	$p_1 \\ .041$	p_2 .159
.20	.122	.278
.30	.210	.390
.40	.304	.496 548 ***
.45	.352	.040
.46	.362	.558
.47	.372	.568
.48	.382	.578
.49	.392	.588
.50	.402	.598
.51	.412	.608
.52	.422	.618
.53	.432	.628
.54	.442	.638
.55	.452	.648
.56	.463	.657
.57	.473	.667
.58	.483	.677
.59	.494	.686
.60	.504	.696
.61	.514	.706
.62	.525	.715
.63	.535	.725
.64	.546	.734
.65	.556	.743 ***
.70	.610	.790
.80	.722	.878
.90	.841	.959

Table 9.1 Central 95% intervals $[p_1, p_2]$ of the distribution of \hat{p} for selected values of p when n = 100. The intervals for p = .46 through p = .64 contain $\hat{p}_{obs} = .55$.

A simple example will help to clarify this discussion. Consider a box containing a large number of balls. Suppose that some of the balls in the box are red. Let p denote the proportion of red balls in the box. Now suppose that a simple random sample of n = 100 balls has been selected and 55 of the 100 balls found to be red. In this example 55% of the balls in the sample are red, *i.e.*, $\hat{p}_{obs} = .55$. Table 9.1 contains central 95% intervals of the distribution of \hat{p} for n = 100 and selected values of p. Notice that when p is between .46 and .64 the intervals contain $\hat{p}_{obs} = .55$.

For example, when p = .50, 95% of all samples will yield a \hat{p} value between .402 and .598. In a case like this, where the central interval contains .55, it would not be surprising to see $\hat{p}_{obs} = .55$ and the correspond value of p would be deemed plausible. That is, if exactly 50% of the balls in the box were red, then 95% of the time when we selected a random sample of n = 100 balls we would find that between 40.2% and 59.8% of the 100 balls in the sample were red. Since 55% belongs to this range it is plausible that exactly 50% of all the balls in the box are red (p = .50).

On the other hand, if p = .30, 95% of all samples will yield a \hat{p} value between .210 and .390. Thus, if exactly 30% of the balls in the box were red, then 95% of the time when we selected a random sample of n = 100 balls we would find that between 21.0% and 39.0% of the 100 balls in the sample were red. Since this entire interval is less than 55%, it would be surprising to see \hat{p} as large as .55 when p was .30. Therefore, when we see $\hat{p}_{obs} = .55$ it is reasonable to conclude that the percentage of red balls in the box is not 30% ($p \neq .30$).

Similarly, if p = .70, 95% of all samples will yield a \hat{p} value between .610 and .790. Thus, if exactly 70% of the balls in the box were red, then 95% of the time when we selected a random sample of n = 100 balls we would find that between 61.0% and 79.0% of the 100 balls in the sample were red. Since this entire interval is greater than 55%, it would be surprising to see \hat{p} as small as .55 when p was .70. Therefore, when we see $\hat{p}_{obs} = .55$ it is reasonable to conclude that the percentage of red balls in the box is not 70% ($p \neq .70$).

In cases like the last two, where the entire interval is less than .55 or the entire interval is greater than .55, it would be surprising to see $\hat{p}_{obs} = .55$ and the correspond value of pwould not be deemed plausible. From the table we can see that $p_L \approx .45$, since the upper endpoint of the central 95% interval p_2 is approximately .55 when p = .45, and $p_U \approx .64$, since the lower endpoint p_1 of the central 95% interval is approximately .55 when p = .64. More precise computation indicates that $p_L = .4524$ and $p_U = .6439$. Using this reasoning, we see that when n = 100 and we observe $\hat{p}_{obs} = .55$ we can argue that values of p between $p_L = .4524$ and $p_U = .6439$ are plausible. In other words, if we selected a simple random sample of n = 100 balls from the box and found that 55% of the balls in the sample were red, then we would conclude that somewhere between 45.24% and 64.39% of all the balls in the box are red.

The interval of plausible values for p with endpoints p_L and p_U discussed above is a 95% confidence interval estimate of p. In the preceding example, the more formal way of stating

the conclusion is as follows. If we selected a simple random sample of n = 100 balls from the box and found that 55% of the balls in the sample were red, then we would conclude that we were 95% confident that the percentage of red balls in the box was somewhere between 45.24% and 64.39% (in terms of p, .4524 $\leq p \leq .6439$).

A confidence interval for p – more formally

We will now give a more formal definition of a confidence interval estimate of p. The purpose of a confidence interval estimate is to provide a range or interval of plausible values for p. In particular, given the data we would like to be able to say, with a reasonable level of confidence (95% in the example above), that the true value of p is between two particular values (p_L and p_U in the example above). A confidence interval estimate of pconsists of two parts. There is an interval of plausible values for p and a corresponding level of confidence. We will adopt the usual convention of using a confidence level of 95%. The confidence level indicates our confidence that the unknown p actually belongs to the corresponding interval.

There is some chance for confusion about what it means to say we are 95% confident that p is between p_L and p_U . The important thing to remember is that it is the endpoints of the interval p_L and p_U that vary from sample to sample. The population proportion p is a fixed, unknown parameter which does not vary. Therefore, the 95% confidence level applies to the method used to generate the confidence interval estimate. That is, the method (obtain a simple random sample and compute the numbers p_L and p_U forming the confidence interval is such that 95% of the time it will yield a pair of confidence interval limits which bracket the population success proportion p. Therefore, when we obtain a sample, compute the confidence interval, and say that we are 95% confident that this interval contains p what we mean is that we feel "pretty good" about claiming that p is in this interval, since the method used to construct the interval works for 95% of all possible samples and so it probably worked for our sample.

A 95% confidence interval for p – summary and computation.

As noted above, a 95% confidence interval estimate of p is an interval of plausible values for p constructed using a method of generating such intervals with the property that this method will work, in the sense of generating an interval that contains p, for 95% of all possible samples.

The Wilson (score) 95% confidence interval for p.

The 95% confidence interval estimate of the population proportion p discussed above is usually called the Wilson interval or the score interval. The graphical derivation above is readily modified for a confidence level other that 95%. The requisite modification is to change the 95% confidence level multiplier 1.96 to the value appropriate for the desired confidence level, *e.g.*, the multiplier 1.645 leads to a 90% confidence interval.

Computation of the Wilson (score) 95% confidence interval for p.

Computations of the Wilson 95% confidence interval estimate of p for the box of balls example with n = 100 and 55 red balls are illustrated in Figures 9.7–9.10. Recall that in this example, we selected a simple random sample of n = 100 balls from the box and found that 55% of the balls in the sample were red, then we concluded that we were 95% confident that the percentage of red balls in the box was somewhere between 45.24% and 64.39% (in terms of p, $.4524 \le p \le .6439$). Figure 9.7 SAS output giving the Wilson 95% confidence interval for the box of balls example with n = 100 and $\hat{p}_{obs} = .55$ with SAS command file.

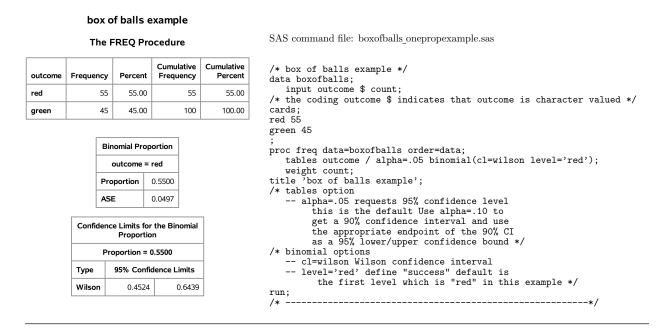
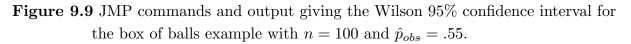


Figure 9.8 R commands and output giving the Wilson 95% confidence interval for the box of balls example with n = 100 and $\hat{p}_{obs} = .55$.

R commands and output for the box of balls example

```
R Commands:
# box of balls example
prop.test(55,100,correct=0)
R Output:
1-sample proportions test without continuity correction
data: 55 out of 100, null probability 0.5
X-squared = 1, df = 1, p-value = 0.3173
alternative hypothesis: true p is not equal to 0.5
95 percent confidence interval:
    0.4524460 0.6438546
sample estimates:
    p
0.55
```

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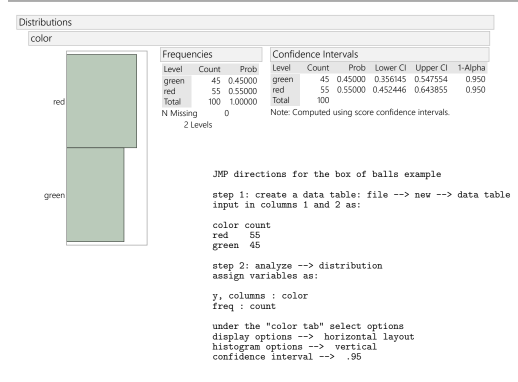
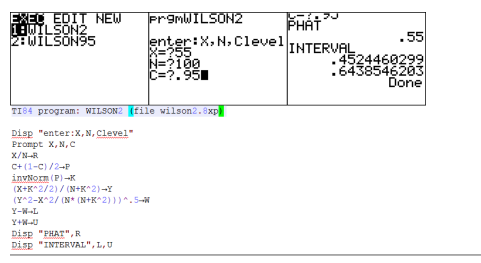


Figure 9.10 TI84 program "WILSON2" screen captures followed by command listing. This output shows how to get the Wilson 95% confidence interval for the box of balls example with n = 100 and $\hat{p}_{obs} = .55$.



Example 9.1 Insects in an apple orchard. The manager of a large apple orchard is concerned with the presence of a particular insect pest in the apple trees in the orchard. An insecticide

that controls this particular insect pest is available. However, application of this insecticide is rather expensive. It has been determined that the cost of applying the insecticide is not economically justifiable unless more than 20% of the apple trees in the orchard are infested. The manager has decided to assess the extent of infestation in the orchard by examining a simple random sample of 200 apple trees. In this example a unit is an apple tree and the target population is all of the apple trees in this orchard. We will assume that the simple random sample is selected from all of the apple trees in the orchard so that the sampled population is the same as the target population. We will also assume that the 200 trees in the sample form a small proportion of all of the trees in the entire orchard so that we do not need to worry about whether the sample is chosen with or without replacement. An appropriate dichotomous variable is whether an apple tree is infested with possible values of yes (the tree is infested) and no (the tree is not infested). Since we are interested in the extent of the infestation we will view a tree that is infested as a success. Thus, the population success proportion p is the proportion of all of the apple trees in the entire orchard that are infested.

Two (related) questions of interest in this situation are:

- (1) What proportion of all of the trees in this orchard are infested? (What is p?)
- (2) Is there sufficient evidence to justify the application of the insecticide? (Is p > .20?)

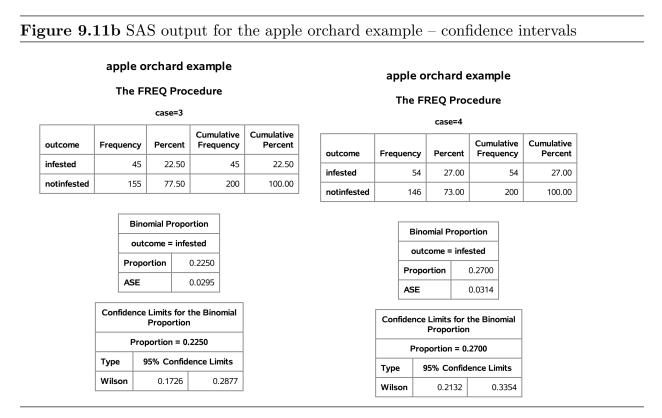
We will consider four hypothetical outcomes for this scenario to demonstrate how a 95% confidence interval estimate can be used to address these questions. The SAS output for these examples is provided in Figures 9.11a and 9.11b.

Figure 9.11a SAS output for the apple orchard example – confidence intervals

apple orchard example The FREQ Procedure case=1							app	ole o	rchard	examp	ole		
							т	he Fl	REQ Pro	cedure	9		
			case-1						case=2				
outcome	Freque	ncy	Percen	Cumulative Frequency		outcome	Frequency		Percent			Cumulative Percent	
infested		35	17.50	35	17.50								
notinfested		165	82.50	200	100.00	infested		26	13.00		26	13.00	
						notinfested		174	87.00		200	100.00	
	Binomial Proportion outcome = infested Proportion 0.1750						omial Propo come = infe	·					
	1	SE		0.0269				ASE		0.0238			
	Confidence Limits for the Binomial Proportion Proportion = 0.1750			-		Confidence Limits for the Binomial Proportion							
	Type 95% Confidence Limits			dence Limits	1		Proportion = 0.1300						
	Wilsor		0.1286	0.2336	-		Type 95% Confidence Limits						
			0.1200	0.2330			Wilso	n _	0.0903	0.	1837		

Case 1. Suppose that 35 of the 200 apple trees in the sample are infested so that $\hat{p} = .175$. In this case we know that 17.5% of the 200 trees in the sample are infested and we can conjecture that a similar proportion of all of the trees in the entire orchard are infested. However, we need a confidence interval estimate to get a handle on which values of the population success proportion p are plausible when we observe 17.5% infested trees in a sample of size 200. Using the Wilson method we get a 95% confidence interval ranging from .1286 to .2336 (see Figure 9.11a). Thus we can conclude that we are 95% confident that between 12.86% and 23.36% of all of the trees in this orchard are infested. Notice that this confidence interval does not exclude the possibility that more than 20% of the trees in the entire orchard are infested, since the upper limit of the confidence interval 23.36% is greater than 20%. In other words, even though less than 20% of the trees in the sample were infested, when we take sampling variability into account we find that it is possible that more than 20% (as high as 23.36%) of the trees in the entire orchard are infested. Of course the interval also indicates that it is possible that less than 20% (as low as 12.86%) of the trees in the entire orchard are infested.

Case 2. Suppose that 26 of the 200 apple trees in the sample are infested so that $\hat{p} = .13$. In this case we know that 13% of the 200 trees in the sample are infested. Using the Wilson method we get a 95% confidence interval ranging from .0903 to .1837 (see Figure 9.11a). Thus we can conclude that we are 95% confident that between 9.03% and 18.37% of all of the trees in this orchard are infested. In this case the entire confidence interval is below 20% excluding the possibility that more than 20% of the trees in the entire orchard are infested. Therefore, in this case we have sufficient evidence to conclude that less than 20% of the trees in the entire orchard are infested, *i.e.*, that p < .20.



Case 3. Suppose that 45 of the 200 apple trees in the sample are infested so that $\hat{p} = .225$. In this case we know that 22.5% of the 200 trees in the sample are infested. Using the Wilson method we get a 95% confidence interval ranging from .1726 to .2877 (see Figure 9.11b). Thus we can conclude that we are 95% confident that between 17.26% and 28.77% of all of the trees in this orchard are infested. Notice that this confidence interval does not exclude the possibility that less than 20% of the trees in the entire orchard are infested, since the lower limit of the confidence interval 17.26% is less than 20%. In other words, even though more than 20% of the trees in the sample were infested, when we take sampling variability into account we find that it is possible that less than 20% (as low as 17.26%)

of the trees in the entire orchard are infested. Of course the interval also indicates that it is possible that more than 20% (as high as 28.77%) of the trees in the entire orchard are infested.

Case 4. Finally, suppose that 54 of the 200 apple trees in the sample are infested so that $\hat{p} = .27$. In this case we know that 27% of the 200 trees in the sample are infested. Using the Wilson method we get a 95% confidence interval ranging from .2132 to .3354 (see Figure 9.11b). Thus we can conclude that we are 95% confident that between 21.32% and 33.54% of all of the trees in this orchard are infested. In this case the entire confidence interval is above 20% excluding the possibility that less than 20% of the trees in the entire orchard are infested. Therefore, in this case we have sufficient evidence to conclude that more than 20% of the trees in the entire orchard are infested, *i.e.*, that p > .20.

A formula for the Wilson (score) 95% confidence interval for p.

Some readers may find an algebraic expression for the Wilson confidence interval useful. Refer to the computer code and output of Figures 9.7–9.10 for an illustration of the computations of the Wilson 95% confidence interval for a simple example. For greater generality we will provide these expressions in terms of k, where k is the multiplier for the desired confidence level. For a 95% confidence level k = 1.96 and for a 90% confidence level k = 1.645. The Wilson confidence interval estimate of p is given by

$$\tilde{p}_k - \operatorname{ME}(\tilde{p}_k) \le p \le \tilde{p}_k + \operatorname{ME}(\tilde{p}_k),$$

(read \tilde{p}_k as p tilde sub k) where

$$\tilde{p}_k = \frac{n\hat{p} + \frac{k^2}{2}}{n+k^2}$$

determines the center of the interval, and the margin of error of \tilde{p}_k

$$\mathrm{ME}(\tilde{p}_k) = \sqrt{\tilde{p}_k^2 - \frac{n\hat{p}^2}{n+k^2}}$$

determines the length of the interval.

If we use k = 1.96 in these expressions, then we can claim that we are 95% confident that the population success proportion p is between $\tilde{p}_k - \text{ME}(\tilde{p}_k)$ and $\tilde{p}_k + \text{ME}(\tilde{p}_k)$. As noted above, there is some chance for confusion about what this statement actually means. The important thing to remember is that it is the statistic \tilde{p}_k and the margin of error $ME(\tilde{p}_k)$ that vary from sample to sample. The population proportion p is a fixed, unknown parameter which does not vary. Therefore, the 95% confidence level applies to the method used to generate the confidence interval estimate. That is, the method (obtain a simple random sample and compute the numbers $\tilde{p} - ME(\tilde{p})$ and $\tilde{p} + ME(\tilde{p})$) used to generate the limits of the confidence interval is such that 95% of the time it will yield a pair of confidence interval limits which bracket the population success proportion p. Therefore, when we obtain a sample, compute the confidence interval, and say that we are 95% confident that this interval contains p what we mean is that we feel "pretty good" about claiming that p is in this interval, since the method works for 95% of all possible samples and so it probably worked for our sample.

Aside – derivation of the Wilson interval formula.

This aside contains an algebraic derivation of the Wilson confidence interval for p. The starting point for this derivation is the interval $p - k\text{SE}(\hat{p}) \leq \hat{p} \leq p + k\text{SE}(\hat{p})$ for \hat{p} . Notice that we can re-express this relationship as $|\hat{p} - p| \leq k\text{SE}(\hat{p})$. Since $|\hat{p} - p|$, k, and $\text{SE}(\hat{p}) = \sqrt{p(1-p)/n}$ are positive, we can square each side of the inequality

$$|\hat{p} - p| \le k \mathrm{SE}(\hat{p})$$

to get the equivalent inequality

$$(\hat{p} - p)^2 \le \frac{k^2}{n}(p - p^2).$$

Straightforward algebra allows us to re–express this inequality as the following quadratic inequality in p

$$(n+k^2)p^2 - 2(n\hat{p} + \frac{k^2}{2})p + n\hat{p}^2 \le 0.$$

Treating this inequality as an equality and solving for p gives the two values

$$\tilde{p}_k \pm \mathrm{ME}(\tilde{p}_k),$$

where
$$\tilde{p}_k = \frac{n\hat{p} + \frac{k^2}{2}}{n+k^2}$$
 and $\operatorname{ME}(\tilde{p}_k) = \sqrt{\tilde{p}_k^2 - \frac{n\hat{p}^2}{n+k^2}}$

Thus, letting C denote the desired confidence level, the probability statement

$$P[|\hat{p} - p| \le k \operatorname{SE}(\hat{p})] = C.$$

is equivalent to the probability statement

$$P[\tilde{p}_k - \operatorname{ME}(\tilde{p}_k) \le p \le \tilde{p}_k + \operatorname{ME}(\tilde{p}_k)] = C.$$

The endpoints of this interval, which are functions of n, \hat{p} , and k, are computable. Therefore, the Wilson confidence interval is given by

$$\tilde{p}_k - \operatorname{ME}(\tilde{p}_k) \le p \le \tilde{p}_k + \operatorname{ME}(\tilde{p}_k).$$

9.4 Testing a hypothesis about p

In the apple orchard example we used a confidence interval estimate of p to decide whether the data supported the contention that more than 20% of the apple trees in the entire orchard were infested. We will now develop some formal methodology for assessing the evidence in favor of such a contention. We will start with some terminology.

A hypothesis (statistical hypothesis) is a conjecture about the nature of the population. When the population is dichotomous, a hypothesis is a conjecture about the value of the population success proportion p.

A hypothesis test (test of significance) is a formal procedure for deciding between two complementary hypotheses. These hypotheses are known as the null hypothesis (H_0 for short) and the research (or alternative) hypothesis (H_1 for short). The research hypothesis is the hypothesis of primary interest, since the testing procedure is designed to address the question: "Do the data support the research hypothesis?" The null hypothesis is defined as the negation of the research hypothesis. The test begins by tentatively assuming that the null hypothesis is true (the research hypothesis is false). The data are then examined to determine whether the null hypothesis can be rejected in favor of the research hypothesis. The probability of observing data as unusual (surprising) or more unusual as that actually observed under the tentative assumption that the null hypothesis is true is computed. This probability is known as the P-value of the test. (The P in P-value indicates that it is a probability it does not refer to the population success proportion p.) A small P-value indicates that the observed data would be unusual (surprising) if the null hypothesis was actually true. Thus if the P-value is small enough, then the null hypothesis is judged untenable and the test rejects the null hypothesis in favor of the research (alternative) hypothesis. On the other hand, a large (not small) P-value indicates that the observed data would not be unusual (not surprising) if the null hypothesis was actually true. Thus

toc

if the P-value is large (not small enough), then the null hypothesis is judged tenable and the test fails to reject the null hypothesis.

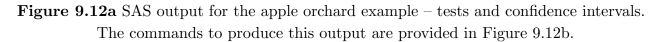
There is a strong similarity between the reasoning used for a hypothesis test and the reasoning used in the trial of a defendant in a court of law. In a trial the defendant is presumed innocent (tentatively assumed to be innocent) and this tentative assumption is not rejected unless sufficient evidence is provided to make this tentative assumption untenable. In this situation the research hypothesis states that the defendant is guilty and the null hypothesis states that the defendant is not guilty (is innocent). The P-value of a hypothesis test is analogous to a quantification of the weight of the evidence that the defendant is guilty with small values indicating that the evidence is unlikely under the assumption that the defendant is innocent.

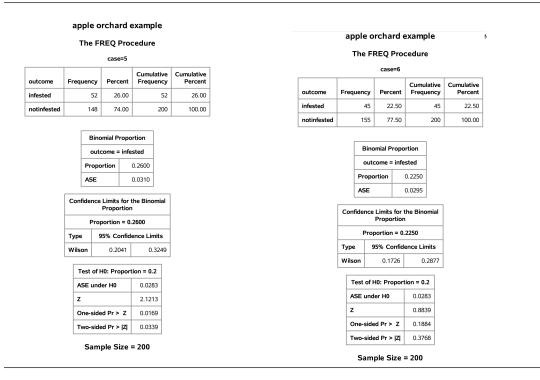
We will introduce hypothesis testing in the context of the apple orchard example. Details and formalization will follow the example.

Example 9.1 Insects in an apple orchard (revisited). Recall that the manager of a large apple orchard examined a simple random sample of 200 apple trees to gauge the extent of insect infestation in the orchard. The manager has determined that applying the insecticide is not economically justifiable unless more than 20% of the apple trees in the orchard are infested. Since the manager does not want to apply the insecticide unless there is evidence that it is needed, the question of interest here is: "Is there sufficient evidence to justify application of the insecticide?" In terms of the population success proportion p(the proportion of all of the apple trees in this orchard that are infested) **the research hypothesis** is $H_1 : p > .20$ (more than 20% of all the trees in the orchard are infested); and **the null hypothesis** is $H_0 : p \leq .20$ (no more than 20% of all the trees in the orchard are infested).

A test of the null hypothesis $H_0: p \leq .20$ versus the research hypothesis $H_1: p > .20$ begins by tentatively assuming that no more than 20% of all the trees in the orchard are infested. Under this tentative assumption it would be surprising to observe a proportion of infested trees in the sample, \hat{p} , that was much larger than .20. For example, when n = 200 and p = .2, the central 95% interval for \hat{p} ranges from .1446 to.2554 and the central 99% interval for \hat{p} ranges from .1272 to.2728. Therefore, if exactly 20% of all the trees in the orchard were infested, then it would be surprising to see much more than about

25% infested trees in the sample (above the central 95% interval) and it would be very surprising to see more than about 27% infested trees in the sample (above the central 99% interval). On the other hand, if exactly 30% of all the trees in the orchard were infested, then the central 95% interval for the sample percentage ranges from 23.65% to 36.35% and the central 99% interval for the sample percentage ranges from 21.65% to 38.35%, so that sample percentages around 25% to 28% would not be surprising. Thus the test should reject $H_0: p \leq .20$ in favor of $H_1: p > .20$ if the observed value of \hat{p} is sufficiently large relative to .20. That is, if \hat{p} is large enough to make us doubt our tentative assumption that p itself is not larger than .20.





Case 1. Suppose that 52 of the 200 apple trees in the sample are infested so that $\hat{p} = .26$. In this case we know that 26% of the 200 trees in the sample are infested and we need to decide whether this suggests that the proportion of all the trees in the orchard that are infested, p, exceeds .20. More specifically, we need to determine whether observing 52 or more infested trees in a simple random sample of 200 trees would be surprising if in fact no more than 20% of all the trees in the orchard were infested. Assuming that exactly 20% of all the trees in the orchard are infested, we find that the probability of observing 52 or more infested trees in a sample of 200 trees (seeing $\hat{p} \ge .26$), is .0169 (this is the *P*-value of the test). In the SAS output of Figure 9.12a (case 5) this *P*-value is labeled "one-sided Pr > Z". In other words, if no more than 20% of all the trees in the orchard were infested, then a simple random sample of 200 trees would give $\hat{p} \ge .26$ about 1.69% of the time. Therefore, observing 52 infested trees in a sample of 200 would be very surprising if no more than 20% of all the trees in the orchard were infested and we have sufficient evidence to reject the null hypothesis $H_0: p \le .20$ in favor of the research hypothesis $H_1: p > .20$. In the case we would conclude that there is sufficient evidence to contend that more than 20% of all the trees in the orchard are infested and, in this sense, application of the insecticide is justifiable. Referring to the SAS output, we see that we can also conclude with 95% confidence that between 20.41% and 32.49% of all the trees in the entire orchard are infested. As expected, we find that the entire 95% confidence interval is above 20%. Note, however, that values as low as 20.41%, which is not much above 20%, are deemed plausible here.

The SAS commands which produced the output in Figure 9.12a are provided in Figure 9.12b.

Case 2. Next suppose that 45 of the 200 apple trees in the sample are infested so that $\hat{p} = .225$. Assuming that exactly 20% of all the trees in the orchard are infested, we find that the probability of observing 45 or more infested trees in a sample of 200 trees (seeing $\hat{p} \ge .225$), is .1884 (this is the *P*-value of the test). In the SAS output of Figure 9.12a (case 6) this *P*-value is labeled "one-sided $\Pr > Z$ ". In other words, if no more than 20% of all the trees in the orchard were infested, then a simple random sample of 200 trees would give $\hat{p} \ge .225$ about 18.84% of the time. Therefore, observing 45 infested trees in a sample of 200 would not be very surprising if no more than 20% of all the trees in the orchard were infested and we do not have sufficient evidence to reject the null hypothesis $H_0 : p \le .20$ in favor of the research hypothesis $H_1 : p > .20$. In the case we would conclude that there is not sufficient evidence to a contend that more than 20% of all the trees in the orchard are infested and, in this sense, application of the insecticide not is justifiable. Referring to the SAS output, we see that we can also conclude with 95% confidence that between 17.26% and 28.77% of all the trees in the entire orchard are infested. As expected, we see that the 95% confidence interval contains values which are both above and below 20%.

Figure 9.12b SAS commands for the apple orchard example – tests and confidence intervals

SAS command file: Apple orchard examples cases 5 and 6

```
/* apple orchard example */
data orchard;
   input case outcome : $ 11. count;
/* the coding outcome : $ 11. indicates that we want
                      to allocate 11 characters for outcome */
cards;
5 infested 52
5 notinfested 148
6 infested 45
6 notinfested 155
proc freq data=orchard order=data;
   tables outcome / alpha=.05 binomial(cl=wilson p=.2 level='infested');
   weight count;
   by case;
title 'apple orchard example';
/* binomial options
   -- cl=wilson Wilson confidence interval
   -- p=.2 use p_0=.2 in the hypothesis
   -- level='infested' define "success" default is
         the first level which is infested in this example */
run;
```

The research hypothesis in the apple orchard example is a **directional hypothesis** of the form $H_1: p > p_0$, where $p_0 = .20$. We will now discuss the details of a hypothesis test for a directional research hypothesis of this form. For the test procedure to be valid the specified value p_0 and the direction of the research hypothesis must be motivated from subject matter knowledge before looking at the data that are to be used to perform the test.

Testing a directional hypothesis of the form $p > p_0$

Research question. Is there sufficient evidence to conclude that the population proportion p is greater than the hypothesized value p_0 ?

Research hypothesis. $H_1: p > p_0$, The population proportion p is greater than the hypothesized value p_0 .

Tentative assumption – null hypothesis. $H_0: p \leq p_0$, We tentatively assume that the population proportion p is not greater than the hypothesized value p_0 .

Evidence in favor of the research hypothesis. The relationship between the observed proportion of successes in the sample \hat{p} and the hypothesized value p_0 will be used to assess the strength of the evidence in favor of the research hypothesis. Generally, we would expect to observe larger values of \hat{p} more often when the research hypothesis $H_1: p > p_0$ is true than when the null hypothesis $H_0: p \leq p_0$ is true. In particular, we can view the observation of a value of \hat{p} that is sufficiently large relative to p_0 as constituting evidence against the null hypothesis $H_0: p \leq p_0$ and in favor of the research hypothesis $H_1: p > p_0$.

Assessment of the strength of the evidence – the P–value of the test. Deciding whether the observed value of \hat{p} is "sufficiently large relative to p_0 " is based on the *P*–value of the test. The *P*–value for testing the null hypothesis $H_0: p \leq p_0$ versus the research hypothesis $H_1: p > p_0$ is the probability of observing a value of \hat{p} as large or larger than the value of \hat{p} that we actually do observe. The *P*–value quantifies the consistency of the observed data with the null hypothesis and may be interpreted as a, somewhat indirect, measure of the strength of the evidence in the data in favor of the research hypothesis and against the null hypothesis. Because the *P*–value is computed under the assumption that the null hypothesis is true (and the research hypothesis is false), the smaller the *P*–value is, the less consistent the observed data are with the null hypothesis. Therefore, since one of the hypothesis is more consistent with the observed data than is the null hypothesis.

Computation of the P-value. The *P*-value of the test is computed under the assumption that the research hypothesis $H_1: p > p_0$ is false and the null hypothesis $H_0: p \le p_0$ is true. Because the null hypothesis only specifies that $p \le p_0$, we need to choose a particular value of p (that is no larger than p_0) in order to compute the *P*-value. It is most appropriate to use $p = p_0$ for this computation. (Recall that in the apple orchard example we used $p_0 = .20$ to compute the *P*-value.) Using $p = p_0$, which defines the boundary between $p \le p_0$, where the null hypothesis is true, and $p > p_0$, where the research hypothesis is true, provides some protection against incorrectly rejecting $H_0: p \le p_0$.

The derivation below is meant to clarify the procedure. We can use a suitable calculator or computer program to perform these computations. We will use the normal approximation to the sampling distribution of \hat{p} to compute the *P*-value. As noted above we will use the

hypothesized value p_0 in our computation of the *P*-value. Thus we will use the population standard deviation of \hat{p}

$$\operatorname{SE}(\hat{p}) = \sqrt{\frac{p_0(1-p_0)}{n}}$$

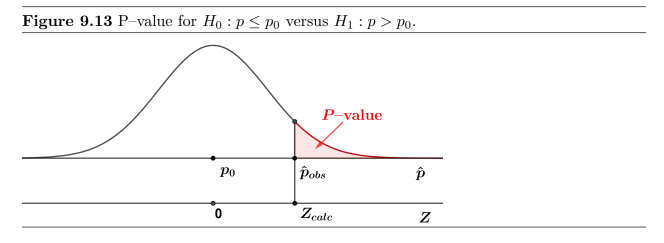
in our computation of the Z-score. The calculated Z statistic or Z score corresponding to the observed value of \hat{p} , denoted by Z_{calc} , is

$$Z_{calc} = \frac{\hat{p} - p_0}{\text{SE}(\hat{p})} = \frac{\hat{p} - p_0}{\sqrt{p_0(1 - p_0)/n}}$$

Recall that the P-value for testing the null hypothesis $H_0: p \leq p_0$ versus the research hypothesis $H_1: p > p_0$ is the probability of observing a value of \hat{p} as large or larger than the value of \hat{p} that we actually do observe, computed assuming that $p = p_0$. Using the normal approximation, this P-value is equal to the probability that a standard normal variable takes on a value at least as large as Z_{calc} . This P-value is

$$P$$
-value = $P(Z \ge Z_{calc}),$

where Z denotes a standard normal variable, *i.e.*, this P-value is the area under the standard normal density curve to the right of Z_{calc} , as shown in Figure 9.13. Notice that the P value (the area to the right of Z_{calc}) is small when Z_{calc} is far to the right of zero which is equivalent to \hat{p} being far to the right of p_0 .



Once the P-value has been computed we need to decide whether the P-value is small enough to justify rejecting the null hypothesis in favor of the research hypothesis. In the apple orchard example we argued that observing 52 infested trees in a sample of 200 would be very surprising if no more than 20% of all the trees in the orchard were infested, since the corresponding P-value of .0169 was very small. We also argued that observing 45 infested trees in a sample of 200 would not be very surprising if no more than 20% of all the trees in the orchard were infested, since the corresponding P-value of .1884 is fairly large. Deciding whether a P-value is small enough to reject a null hypothesis requires a subjective judgment by the investigator in the context of the problem at hand. Some guidelines for interpreting a P-value are provided below.

Returning to our discussion for the directional research hypothesis $H_1: p > p_0$. The final steps for performing a hypothesis test for

$$H_0: p \le p_0$$
 versus $H_1: p > p_0$

are summarized below.

1. Use a suitable calculator or computer program to find the P-value = $P(Z \ge Z_{calc})$, where Z denotes a standard normal variable, $Z_{calc} = (\hat{p} - p_0)/\text{SE}(\hat{p})$, and $\text{SE}(\hat{p}) = \sqrt{p_0(1-p_0)/n}$. This P-value is the area under the standard normal density curve to the right of Z_{calc} , as shown in Figure 9.13. This P-value is the "one-sided Pr > Z" of the SAS output.

2a. If the *P*-value is small enough (less than .05 for a test at the 5% level of significance), conclude that the data favor $H_1: p > p_0$ over $H_0: p \le p_0$. That is, if the *P*-value is small enough, then there is sufficient evidence to conclude that the population success proportion p is greater than p_0 .

2b. If the *P*-value is not small enough (is not less than .05 for a test at the 5% level of significance), conclude that the data do not favor $H_1: p > p_0$ over $H_0: p \le p_0$. That is, if the *P*-value is not small enough, then there is not sufficient evidence to conclude that the population success proportion p is greater than p_0 .

Remarks and guidelines about P-values.

The following general remarks regarding the use of P-values to assess the evidence against a null hypothesis and in favor of a research hypothesis apply to hypothesis tests in general, not just hypothesis tests for a proportion.

One approach to hypothesis testing is to use a fixed cutoff value to decide whether the P-value is "large" or "small". The most common application of this approach is to conclude that there is sufficient evidence to reject the null hypothesis in favor of the research

hypothesis only when the P-value is less than .05. When a fixed cutoff value like .05 (5%) is used to decide whether to reject the null hypothesis in favor of the research hypothesis this cutoff value is known as the **significance level** of the test. Hence, if we adopt the rule of rejecting the null hypothesis in favor of the research hypothesis only when the P-value is less than .05, then we are performing a hypothesis test at the 5% level of significance. In accordance with this terminology, the P-value is also known as the **observed significance level** of the test and if the P-value is less than the prescribed significance level, then the results are said to be **statistically significant**.

To perform a hypothesis test at the 5% level of significance we compute the appropriate P-value and compare it to the fixed significance level .05. If the P-value is less than .05, then we conclude that there is sufficient evidence, at the 5% level of significance, to reject the null hypothesis H_0 in favor of the research hypothesis H_1 , *i.e.*, if the P-value **is less than** .05, then the data **do** support H_1 . If the P-value is not less than .05, then we conclude that there is not sufficient evidence, at the 5% level of significance, to reject the null hypothesis H_0 in favor of the research hypothesis H_1 , *i.e.*, if the P-value **is not less than** .05, then we conclude that there is not sufficient evidence, at the 5% level of significance, to reject the null hypothesis H_0 in favor of the research hypothesis H_1 , *i.e.*, if the P-value **is not less than** .05, then the data **do not** support H_1 .

Instead of, or in addition to, using a fixed significance level like 5% we can use the P-value as a measure of the evidence (in the data) against the null hypothesis H_0 and in favor of the research hypothesis H_1 . Some guidelines for deciding how strong the evidence is in favor of the research hypothesis H_1 are given below.

Guidelines for interpreting a P-value:

1. If the *P*-value is greater than .10, there is no evidence in favor of H_1 .

2. If the *P*-value is between .05 and .10, there is suggestive but very weak evidence in favor of H_1 .

3. If the *P*-value is between .04 and .05, there is weak evidence in favor of H_1 .

4. If the P-value is between .02 and .04, there is moderately strong evidence in favor of H_1 .

5. If the *P*-value is between .01 and .02, there is strong evidence in favor of H_1 .

6. If the *P*-value is less than .01, there is very strong evidence in favor of H_1 .

Whether you choose to use a fixed significance level or the preceding guidelines based on the P-value you should always report the P-value since this allows someone else to interpret the evidence in favor of H_1 using their personal preferences regarding the size of a P-value.

In the U.S. legal system there is a similar set of guidelines for assessing the level of proof or weight of the evidence against the null hypothesis of innocence and in favor of the research hypothesis of guilt. The weakest level of proof is "the preponderance of the evidence" (this is similar to a reasonably small P-value), the next level of proof is "clear and convincing evidence" (this is similar to a small P-value), and the highest level of proof is "beyond a reasonable doubt" (this is similar to a very small P-value).

Example 9.2 Acceptance sampling for electronic devices. A large retailer receives a shipment of 10,000 electronic devices from a supplier. The supplier guarantees that no more than 6% of these devices are defective. In fact, if more than 6% of the devices in the shipment are defective, then the supplier will allow the retailer to return the entire shipment, provided this is done with 10 days of receiving the shipment. Therefore, the retailer needs to decide between accepting the shipment and returning the shipment to the supplier. This decision will be based on the information provided by examining a simple random sample of electronic devices selected from the shipment.

In this example one of these electronic devices is a unit and the collection of 10,000 units constituting the shipment is the population. Notice that, in this example, the target population and the sampled population are the same (each is the shipment of 10,000 devices). A suitable variable for the indicated objective is whether an electronic device is defective with the two possible values: yes (it is defective) and no (it is not defective). A relevant parameter is the proportion p of defective devices in the shipment of 10,000 devices. The corresponding statistic \hat{p} is the proportion of defective devices in the sample of devices that is examined.

The boundary between the null and research hypotheses is clearly $p_0 = .06$, since we need to decide whether the population proportion of defective devices p exceeds .06. Assuming that the supplier is trustworthy, it would seem to be a reasonable business practice to accept the shipment of electronic devices unless we find sufficient evidence, by examining the sample of devices, to conclude that more than 6% of the devices in the shipment

are defective. Hence, we will use a hypothesis test to determine whether there is sufficient evidence to conclude that the population defective proportion p exceeds .06. More formally, our research hypothesis is $H_1: p > .06$ and our null hypothesis is $H_0: p \leq .06$.

To continue with this example we need to know the sample size n and the results of the examination of the sample of electronic devices. Suppose that the simple random sample contains n = 200 electronic devices. For a sample of size n = 200 the standard error of \hat{p} for testing a hypothesis with $p_0 = .06$ is

$$SE(\hat{p}) = \sqrt{\frac{(.06)(.94)}{200}} = .0168.$$

acceptance sampling example М acceptance sampling example Mo The FREQ Procedure The FREQ Procedure case=1 case=2 Cumulative Cumulative Percent outcom Frequency Percent Frequency Cumulative Frequency Cumulative outcome Frequency Percent Percent defective 16 8.00 16 8.00 10.00 20 defective 20 10.00 notdefective 184 92.00 200 100.00 90.00 200 180 100.00 notdefective **Binomial Proportion Binomial Proportion** outcome = defective outcome = defective Proportion 0.0800 0.1000 Proportion 0.0192 ASE ASE 0.0212 Confidence Limits for the Binomial Proportion Confidence Limits for the Binomial Proportion Proportion = 0.0800 Proportion = 0.1000 Туре 95% Confidence Limits 95% Confidence Limits Туре Wilson 0.0498 0.1260 Wilson 0.0657 0.1494 Test of H0: Proportion = 0.06 Test of H0: Proportion = 0.06 ASE under H0 0.0168 ASE under H0 0.0168 1.1910 7 z 2.3820 One-sided Pr > Z 0.1168 One-sided Pr > Z 0.0086 Two-sided Pr > |Z| 0.2337 Two-sided Pr > |Z| 0.0172 Sample Size = 200 Sample Size = 200

Figure 9.14 SAS output for the acceptance sampling example

Case 1. Suppose that 16 of the 200 devices in the sample are defective so that $\hat{p} = .08$. In this case we know that 8% of the 200 devices in the sample are defective and we need to decide whether this suggests that more than 6% of all the devices in the shipment are defective. The calculated Z statistic is

$$Z_{calc} = \frac{\hat{p} - p_0}{\mathrm{SE}(\hat{p})} = \frac{.08 - .06}{.0168} = 1.1910$$

and the P-value is

$$P$$
-value = $P(Z \ge Z_{calc}) = P(Z \ge 1.1910) = .1168.$

In the SAS output of Figure 9.14 this P-value is labeled "one-sided $\Pr > Z$. Since this P-value is large there is not sufficient evidence to reject the null hypothesis $p \leq .06$ in favor of the research hypothesis p > .06. Therefore, if we observe 16 defective devices in a random sample of n = 200 devices, then we should accept the shipment of devices, since there is not sufficient evidence to conclude that more than 6% of the shipment of 10,000 devices is defective.

Case 2. Now suppose that 20 of the 200 devices in the sample are defective so that $\hat{p} = .10$. In this case

$$Z_{calc} = \frac{\hat{p} - p_0}{\mathrm{SE}(\hat{p})} = \frac{.10 - .06}{.0168} = 2.3820$$

and the P-value is

$$P$$
-value = $P(Z \ge Z_{calc}) = P(Z \ge 2.3820) = .0086.$

In the SAS output of Figure 9.14 this P-value is labeled "one-sided $\Pr > Z$. This P-value is very small indicating that we have strong evidence against the null hypothesis $p \leq .06$ and in favor of the research hypothesis p > .06. Therefore, if we observe 20 defective devices in a random sample of n = 200 devices, then we are justified in returning the shipment of devices, since there is strong evidence that more than 6% of the shipment of 10,000 devices is defective.

In both of the cases described above, in addition to the conclusion of the hypothesis test the retailer might also wonder exactly what proportion of devices in the shipment of 10,000 devices are defective. We can use a 95% confidence interval estimate of p to answer this question.

In the first case there are 16 defective devices in the sample of n = 200 giving an observed proportion of defective devices of $\hat{p} = .08$. From the SAS output in Figure 9.14, we are 95% confident that the actual proportion of defective devices in the shipment of 10,000 is between .0498 and .1260. As expected, since we did not reject the tentative assumption that $p \leq .06$, we see that this confidence interval includes proportions that are both less than .06 and greater than .06.

In the second case there are 20 defective devices in the sample of n = 200 giving $\hat{p} = .10$. From the SAS output in Figure 9.14, we are 95% confident that the actual proportion of defective devices in the shipment of 10,000 is between .0656 and .1494. As expected, since we did reject the tentative assumption that $p \leq .06$, we see that all of the values in this confidence interval are greater than .06. Notice that in this case the *P*-value .0086 is quite small indicating that there is very strong evidence that the proportion of defective devices in the shipment is larger than .06. However, from the 95% confidence interval estimate of *p* we find that this proportion of defective devices might actually be as small as .0656, which is not much larger than .06. Thus, the small *P*-value indicates strong evidence that *p* is greater than .06 but it does not necessarily indicate that *p* is a lot larger than .06. Of course the 95% confidence interval estimate also indicates that *p* may be as large as .1494 which is a good bit larger than .06.

The scenario in the acceptance sampling example where there is strong evidence that p > .06 (*P*-value .0086) but the lower limit of the 95% confidence interval .0656 is not much larger than .06 highlights the need for a confidence interval to estimate the value of p in addition to a hypothesis test to clarify the practical importance of the result of the test. Bear in mind that a hypothesis test addresses a very formal distinction between two complementary hypotheses and that in some situations the results may be statistically significant (in the sense that the *P*-value is small) but of little practical significance (in the sense that p is not very different from p_0).

Testing a directional hypothesis of the form $p < p_0$

The procedure for testing the null hypothesis $H_0: p \leq p_0$ versus the research hypothesis $H_1: p > p_0$ given above is readily modified for testing the null hypothesis $H_0: p \geq p_0$ versus the research hypothesis $H_1: p < p_0$. The essential modification is to change the direction of the inequality in the definition of the *P*-value. Consider a situation where the research hypothesis specifies that the population success proportion p is less than the particular, hypothesized value p_0 , *i.e.*, consider a situation where the research hypothesis is $H_1: p < p_0$ and the null hypothesis is $H_0: p \geq p_0$. For these hypotheses values of the observed success proportion \hat{p} that are sufficiently small relative to p_0 provide evidence in favor of the research hypothesis $H_1: p < p_0$ and against the null hypothesis $H_0: p \geq p_0$. Therefore, the *P*-value for testing $H_0: p \geq p_0$ versus $H_1: p < p_0$ is the probability of observing a value of \hat{p} as small or smaller than the value actually observed. As before, the

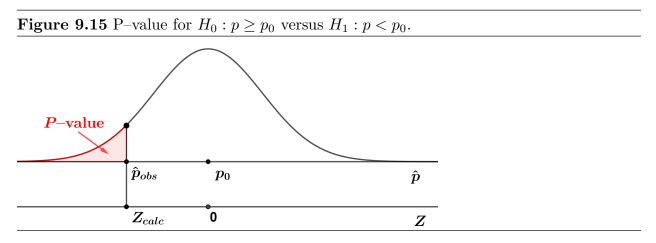
P-value is computed under the assumption that $p = p_0$. The calculated Z statistic Z_{calc} is defined as before; however, in this situation the P-value is the area under the standard normal density curve to the left of Z_{calc} , since values of \hat{p} that are small relative to p_0 constitute evidence in favor of the research hypothesis.

The steps for performing a hypothesis test for

$$H_0: p \ge p_0$$
 versus $H_1: p < p_0$

are summarized below.

1. Use a suitable calculator or computer program to find the P-value = $P(Z \leq Z_{calc})$, where Z denotes a standard normal variable, $Z_{calc} = (\hat{p} - p_0)/\text{SE}(\hat{p})$, and $\text{SE}(\hat{p}) = \sqrt{p_0(1-p_0)/n}$. This P-value is the area under the standard normal density curve to the left of Z_{calc} as shown in Figure 9.15. This P-value is the "one-sided Pr < Z" of the SAS output.



2a. If the P-value is small enough (less than .05 for a test at the 5% level of significance), conclude that the data favor $H_1: p < p_0$ over $H_0: p \ge p_0$. That is, if the P-value is small enough, then there is sufficient evidence to conclude that the population success proportion p is less than p_0 .

2b. If the *P*-value is not small enough (is not less than .05 for a test at the 5% level of significance), conclude that the data do not favor $H_1: p < p_0$ over $H_0: p \ge p_0$. That is, if the *P*-value is not small enough, then there is not sufficient evidence to conclude that the population success proportion p is less than p_0 .

Bernoulli trials

Some applications of these inferential methods for a proportion, such as the following machine parts example, correspond to a sequence of n trials. In this context a dichotomous trial is a process of observation or experimentation which results in one of two distinct outcomes (success or failure).

A sequence of n trials is said to constitute a sequence of **n** Bernoulli trials with success **probability p** if the following conditions are satisfied.

1. There is a common probability of success p for every trial. That is, on every trial the probability that the outcome of the trial will be a success is p.

2. The outcomes of the trials are independent of each other. That is, if we knew the outcome of a particular trial or trials this would provide no additional information about the probability of observing a success (or failure) on any other trial. For example, if we knew that a success (or failure) occurred in the first trial, this would not change the probability of success in any other trial.

The simple example below will help to clarify the definition of a sequence of n Bernoulli trials and the connection between sampling from a dichotomous population and Bernoulli trials.

Example 9.3 Tossing a fair die. Let a trial consist of tossing a fair (balanced) die and observing the number of dots on the upturned face. Define a success to be the occurrence of a 1, 2, 3, or 4. Since the die is fair, the probability of a success on a single trial is p = 4/6 = 2/3. Furthermore, if the die is always tossed in the same fashion, then the outcomes of the trials are independent. Therefore, with success defined as above, tossing the fair die n times yields a sequence of n Bernoulli trials with success probability p = 2/3.

Note that this process of tossing a die is abstractly the same as the process of selecting a ball at random from a box containing six balls with the balls numbered from 1 to 6. Thus tossing the die n times is equivalent to selecting a simple random sample of size n with replacement from this box containing six balls.

Example 9.4 Machine parts. The current production process used to manufacture a particular machine part is known (from past experience) to produce parts which are unacceptable, in the sense that they require further machining, 35% of the time. A new production process has been developed with the hope that it will reduce the chance of producing unacceptable parts. Suppose that 200 parts are produced using the new production process and that 54 of these parts are found to be unacceptable.

In this example we have a sequence of 200 dichotomous trials, where a trial consists of producing a part with the new production process and determining whether it is unacceptable. In this example p denotes the probability that a part produced using the new production process will be unacceptable. We will model these 200 trials as a sequence of n = 200 Bernoulli trials with population success probability p. This assumption is reasonable provided: (1) the probability that a part is unacceptable is essentially constant from part to part; and, (2) whether a specific part is unacceptable or not has no effect on the probability that any other part is unacceptable.

In this example the boundary between the null and research hypotheses is clearly $p_0 = .35$. Since these data were collected to determine if the new production process is better than the old process, we want to know whether there is sufficient evidence to conclude that less than 35% of the parts produced using the new production process would be unacceptable. Thus our research hypothesis is $H_1: p < .35$ and our null hypothesis is $H_0: p \ge .35$. Since 54 of the 200 parts in our sample are unacceptable we know that $\hat{p} = .27$ and we need to determine whether this is small enough to suggest that the corresponding population probability p is also less than .35. For a sample of size n = 200 the standard error of \hat{p} for testing a hypothesis with $p_0 = .35$ is

$$\operatorname{SE}(\hat{p}) = \sqrt{\frac{(.35)(.65)}{200}} = .0337.$$

The calculated Z statistic is

$$Z_{calc} = \frac{\hat{p} - p_0}{\mathrm{SE}(\hat{p})} = \frac{.27 - .35}{.0337} = -2.3739$$

and the P-value is

$$P$$
-value = $P(Z \le Z_{calc}) = P(Z \le -2.3739) = .0088.$

You can find these values in the SAS output of Figure 9.16. Since this P-value is very small, there is sufficient evidence to reject the null hypothesis $p \ge .35$ in favor of the research hypothesis p < .35. Hence, based on this sample of 200 parts there is very strong evidence that the new production process is superior in the sense that the probability of producing an unacceptable part is less than .35.

Clearly this conclusion should be accompanied by an estimate of how much smaller this probability is likely to be. Observing 54 unacceptable parts in the sample of n = 200 gives $\hat{p} = .27$ and a 95% confidence interval ranging from .2132 to .3354. Therefore, we are 95% confident that the probability of a part produced using the new production process being unacceptable is between .2132 and .3354. As expected, since we did reject the tentative assumption that $p \ge .35$, we see that all of the values in this confidence interval are less than .35. The *P*-value .0088 is quite small indicating that there is very strong evidence that the probability of producing an unacceptable part is less than .35. However, from the 95% confidence interval estimate of p we find that this probability might actually be as large as .3354 which is not much smaller than .35. Of course the 95% confidence interval estimate also indicates that p may be as small as .2132 which is a good bit smaller than .35.

	mach	ine	e pa	rts e	xampl	e					
The FREQ Procedure											
case=1											
outcome	Frequer	Frequency Perce			Cumul Frequ						
unacceptable		54	2	27.00	1		27.0				
acceptable		146	7	3.00	200		100.00				
]										
	out	outcome = unacceptabl			ptable]					
	Prop	Proportion]					
	ASE	ASE			0.0314	1					
	Confide	vial									
	connaci										
	F										
	Туре	Type 95% Confidence Limits									
	Wilson 0.2132 0.3354					354					
	Test										
	ASE u	ASE under H0				-					
	z				-2.3720						
	One-s	One-sided Pr < Z			0.0088	1					
	Two-sided Pr > Z				0.0177	-					
					200						
	Sa	imp	ne S	ize =	200						

Figure 9.16 SAS output for the machine parts example

Testing a nondirectional research hypothesis

The hypothesis tests we have discussed thus far are only appropriate when we have enough a priori information, *i.e.*, information that does not depend on the data to be used for the hypothesis test, to postulate that the population success proportion p is on one side of a particular value p_0 . That is, we have only considered situations where the research hypothesis is directional in the sense of specifying either that $p > p_0$ or that $p < p_0$. In some situations we will not have enough a priori information to allow us to choose the appropriate directional research hypothesis. Instead, we might only conjecture that the population success proportion p is different from some particular value p_0 . In a situation like this our research hypothesis specifies that the population success proportion p is different from some particular value p_0 . In a situation like this our research hypothesis specifies that the population success proportion p is different from p_0 , *i.e.*, $H_1 : p \neq p_0$ and the corresponding null hypothesis specifies that p is exactly equal to p_0 , *i.e.*, $H_0 : p = p_0$. As we will see in the inheritance model considered below, when testing to see whether p is equal to a specified value p_0 the null hypothesis $H_0 : p = p_0$ often corresponds to the validity of a particular theory or model and the research hypothesis or alternative hypothesis specifies that the theory is invalid.

Testing a nondirectional research (alternative) hypothesis of the form $p \neq p_0$.

Research question. Is there sufficient evidence to conclude that the population proportion p is different from the hypothesized value p_0 ?

Research hypothesis. $H_1: p \neq p_0$, The population proportion p is not equal to the hypothesized value p_0 .

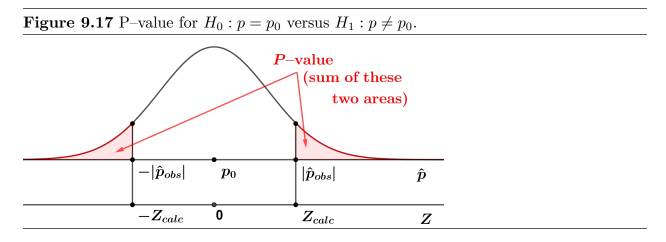
Tentative assumption – null hypothesis. $H_0: p = p_0$, We tentatively assume that the population proportion p is exactly equal to the hypothesized value p_0 .

Evidence in favor of the research hypothesis. As with the directional hypothesis cases, the relationship between the observed proportion of successes in the sample \hat{p} and the hypothesized value p_0 will be used to assess the strength of the evidence in favor of the research hypothesis. Generally, we would expect to observe values of \hat{p} farther away from p_0 more often when the research hypothesis $H_1: p \neq p_0$ is true than when the null hypothesis $H_0: p = p_0$ is true. In particular, we can view the observation of a value of \hat{p} that is sufficiently far away from p_0 , in either direction, as constituting evidence against the null hypothesis $H_0: p = p_0$ and in favor of the research hypothesis $H_1: p \neq p_0$. Assessment of the strength of the evidence – the P–value of the test. Deciding whether the observed value of \hat{p} is "sufficiently far away from p_0 in either direction" is based on the P–value of the test. The P–value for testing the null hypothesis $H_0: p = p_0$ versus the research hypothesis $H_1: p \neq p_0$ is the probability of observing a value of \hat{p} for which the distance $|\hat{p} - p_0|$ (the absolute value of the difference between \hat{p} and p_0) is as large or larger than the value of this distance that we actually do observe.

Computation of the P-value. The *P*-value of the test is computed under the assumption that the research hypothesis $H_1: p \neq p_0$ is false and the null hypothesis $H_0: p = p_0$ is true. In this situation the calculated *Z* statistic Z_{calc} is the absolute value of the *Z* statistic that would be used for testing a directional hypothesis. That is, the calculated *Z* statistic is

$$Z_{calc} = \left| \frac{\hat{p} - p_0}{\mathrm{SE}(\hat{p})} \right| = \left| \frac{\hat{p} - p_0}{\sqrt{p_0(1 - p_0)/n}} \right|.$$

In terms of this Z statistic the P-value is the probability that the absolute value of a standard normal variable Z would take on a value as large or larger than Z_{calc} assuming that $p = p_0$. This probability is the sum of the area under the standard normal density curve to the left of $-Z_{calc}$ and the area under the standard normal density curve to the right of Z_{calc} , as shown in Figure 9.17. We need to add these two areas (probabilities) since we are finding the probability that the observed success proportion \hat{p} would be as far or farther away from p_0 in either direction as is the value that we actually observe, when $p = p_0$. Notice that this P value (the area to the left of $-Z_{calc}$ plus the area to the right of Z_{calc}) is small when Z_{calc} is far away from zero in one direction or the other which is equivalent to \hat{p} being far away from p_0 in one direction or the other.



The steps for performing a hypothesis test for

$$H_0: p = p_0$$
 versus $H_1: p \neq p_0$

are summarized below.

1. Use a suitable calculator or computer program to find the P-value = $P(|Z| \ge Z_{calc}) = P(Z \le -Z_{calc}) + P(Z \ge Z_{calc})$, where Z denotes a standard normal variable, $Z_{calc} = |(\hat{p} - p_0)/\text{SE}(\hat{p})|$, and $\text{SE}(\hat{p}) = \sqrt{p_0(1 - p_0)/n}$. This P-value is the sum of the area under the standard normal density curve to the left of $-Z_{calc}$ and the area under the standard normal density curve to the right of Z_{calc} as shown in Figure 9.17. This P-value is the "two-sided $\Pr > |Z|$ " of the SAS output.

2a. If the *P*-value is small enough (less than .05 for a test at the 5% level of significance), conclude that the data favor $H_1: p \neq p_0$ over $H_0: p = p_0$. That is, if the *P*-value is small enough, then there is sufficient evidence to conclude that the population success proportion p is different from p_0 .

2b. If the *P*-value is not small enough (is not less than .05 for a test at the 5% level of significance), conclude that the data do not favor $H_1: p \neq p_0$ over $H_0: p = p_0$. That is, if the *P*-value is not small enough, then there is not sufficient evidence to conclude that the population success proportion p is different from p_0 .

Example 9.5 Inheritance in peas (flower color). In his investigations, during the years 1856 to 1868, of the chromosomal theory of inheritance Gregor Mendel performed a series of experiments on ordinary garden peas. One characteristic of garden peas that Mendel studied was the color of the flowers (red or white). When Mendel crossed a plant with red flowers with a plant with white flowers, the resulting offspring all had red flowers. But when he crossed two of these first generation plants, he observed plants with white as well as red flowers. We will use the results of one of Mendel's experiments to test a simple model for inheritance of flower color. Mendel observed 929 pea plants arising from a cross of two of these first generation plants. Of these 929 plants he found 705 plants with red flowers and 224 plants with white flowers.

The gene which determines the color of the flower occurs in two forms (alleles). Let R denote the allele for red flowers (which is dominant) and r denote the allele for white flowers

(which is recessive). When two plants are crossed the offspring receives one allele from each parent, thus there are four possible genotypes (ordered combinations) RR, Rr, rR, and rr. The three genotypes RR, Rr, and rR, which include the dominant R allele, will yield red flowers while the fourth genotype rr will yield white flowers. If a red flowered RR genotype parent is crossed with a white flowered rr genotype parent, then all of the offspring will have genotype Rr and will produce red flowers. If two of these first generation Rr genotype plants are crossed, each of the four possible genotypes RR, Rr, rR, and rr is equally likely and plants with white as well as red flowers will occur. Under this simple model for inheritance, with each of the four genotypes having the same probability of occurring (and with each plant possessing only one genotype), the probability that a plant will have red flowers is p = 3/4 and the probability that a plant will have white flowers is 1 - p = 1/4. In other words, this model for inheritance of flower color says that we would expect to see red flowers 3/4 of the time and white flowers 1/4 of the time.

We can test the validity of this model by testing the null hypothesis $H_0: p = 3/4$ versus the alternative hypothesis $H_1: p \neq 3/4$. Notice that the model is valid under the null hypothesis and the model is not valid under the alternative hypothesis. Mendel observed 705 plants with red flowers out of the n = 929 plants giving an observed proportion of plants with red flowers of $\hat{p} = 705/929 = .7589$. The standard error of \hat{p} , computed under the assumption that $p = p_0 = 3/4$, is

$$\operatorname{SE}(\hat{p}) = \sqrt{\frac{(.75)(.25)}{929}} = .0142$$

and the calculated Z statistic is $Z_{calc} = .6251$ giving a P-value of

$$P$$
-value = $P(|Z| \ge Z_{calc}) = P(|Z| \ge .6251) = .5319.$

You can find these values in the SAS output of Figure 9.18. This P-value is quite large and we are not able to reject the null hypothesis; therefore, we conclude that the observed data are consistent with Mendel's model. Technically, we should say that the data are not inconsistent with the model in the sense that we cannot reject the hypothesis that p = 3/4. In this example, the 95% confidence interval estimate of p ranges from .7303 to .7853.

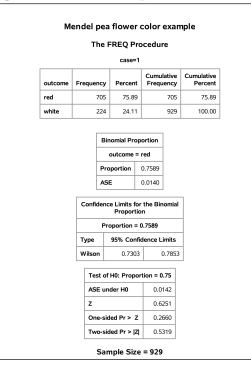


Figure 9.18 SAS output for the Mendel pea flower color example

9.5 Directional confidence bounds

toc

In our discussion of hypothesis testing we considered directional research hypotheses of the form $p > p_0$ and $p < p_0$ as well as nondirectional research hypotheses of the form $p \neq p_0$. However, in our discussion of 95% confidence intervals for p we only considered "nondirectional" confidence intervals of the form $p_L \leq p \leq p_U$. A 95% confidence interval of this form, consisting of a lower bound p_L for p and an upper bound p_U for p, gives a range of plausible values for p. In a situation where we have enough a priori information to justify a directional research hypothesis we might argue that it would be more appropriate to determine a 95% confidence bound (a lower bound or an upper bound) for p instead of a range of values.

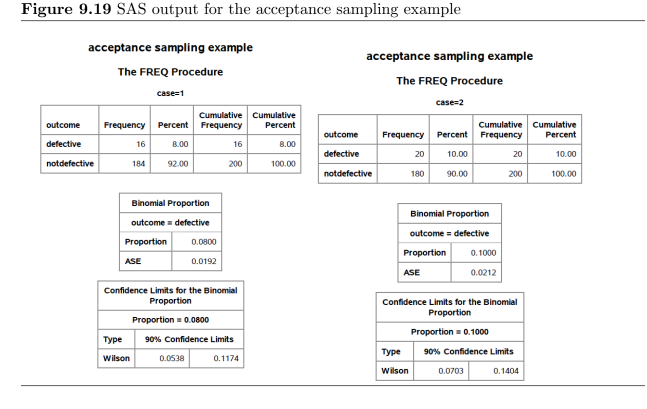
A lower confidence bound on p allows us to estimate the smallest value of p which is plausible in light of the observed value of \hat{p} . Similarly, an upper confidence bound on pallows us to estimate the largest value of p which is plausible in light of the observed value of \hat{p} . For example, in the acceptance sampling example we might argue that we are less concerned with a limit on how large p might be than with a limit on how small it might be. Therefore, we might be satisfied with an estimate of the smallest value of p which would be consistent with the data, *i.e.*, we might only need a 95% confidence lower bound for p.

The reasoning which led to the "nondirectional" Wilson interval (a range of values for p) can be adapted to yield a directional interval (a lower or upper confidence bound for p). Recall that our development of the Wilson 95% confidence interval estimate of p was based on the observation that, for each possible value of p, we can view the central 95% interval from $p - 1.96\text{SE}(\hat{p})$ to $p + 1.96\text{SE}(\hat{p})$ as an interval which is likely to contain \hat{p} . This starting point led us to a confidence interval estimate of p which provided a range of values between a lower limit and an upper limit. If we use an upper 95% interval for \hat{p} or a lower 95% interval for \hat{p} instead, we will end up with a directional confidence interval, *i.e.*, we will end up with a 95% confidence bound (lower or upper) for p. In practice, 95% confidence interval. That is, if $p_L \leq p \leq p_U$ is a 90% confidence interval estimate of p, then P_L is a 95% lower confidence bound for p and p_U is a 95% upper confidence bound for p.

We will illustrate some applications of confidence bounds in the context of some of the examples we discussed earlier. Detailed derivations of 95% lower and upper confidence bounds are provided after the examples.

Example 9.2 Acceptance sampling for electronic devices (revisited). Recall that in this example the retailer had received a shipment of 10,000 electronic devices from a supplier with a guarantee that no more than 6% of these devices were defective. The retailer is interested in p, the value of the proportion of defective devices in the shipment of 10,000 devices. In particular, the retailer is concerned that this proportion might be too large (greater than .06). Thus we proposed a test of the null hypothesis $H_0: p \leq .06$ versus the research hypothesis $H_1: p > .06$. We considered two cases to illustrate the corresponding hypothesis testing procedure. We will now show how a directional confidence bound can supplement the formal hypothesis test.

As noted above, in this example, we are less concerned with a limit on how large p might be than with a limit on how small it might be. Therefore, we might be satisfied with an estimate of the smallest value of p which would be consistent with the data, *i.e.*, we might only need a 95% confidence lower bound for p. Note that a lower confidence bound will always include values greater than .06 (6%). The important consideration is whether it also includes values less than .06.



Case 1 If there are 16 defective devices in a sample of n = 200, then the observed proportion of defective devices is $\hat{p} = .08$. The lower endpoint .0538 of the 90% confidence interval in the SAS output for case 1 in Figure 9.19 is the 95% confidence lower bound for p. Hence, we can conclude that we are 95% confident that the actual percentage of defective devices in the shipment of 10,000 is at least 5.38%. More importantly, with 95% confidence we can say that the percentage could be as low as 5.38%. Since this lower bound is less than 6% we do not have sufficient evidence to claim that more than 6% of the 10,00 devices are defective.

Recall that, in this case, the *P*-value for testing $H_0: p \leq .06$ versus $H_1: p > .06$ was .1168 and we concluded that there is not sufficient evidence to claim that more than 6% of the 10,000 devices are defective. Thus the lower confidence bound and the formal hypothesis test lead to the same conclusion.

Case 2 If there are 20 defective devices in a sample of n = 200, then the observed proportion of defective devices is $\hat{p} = .10$. The lower endpoint .0703 of the 90% confidence

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interval in the SAS output for case 2 in Figure 9.19 is the 95% confidence lower bound for p. In this case we can conclude that we are 95% confident that the actual percentage of defective devices in the shipment of 10,000 is at least 7.03%. That is, with 95% confidence we can say that the percentage is not less than 7.03%. Since this lower bound is greater than 6% we have sufficient evidence to claim that more than 6% of the 10,000 devices are defective. Note also that the lower bound 7.03% indicates that the percentage of defective devices among the 10,000 is at least 1.03 percentage points higher than the 6% cutoff value.

In this case, the *P*-value for testing $H_0: p \leq .06$ versus $H_1: p > .06$ was .0086 and we concluded that there is strong evidence that more than 6% of the 10,000 devices are defective. Again, the lower confidence bound and the test lead to the same conclusion.

Example 9.4 Machine parts (revisited). Recall that, in this example, the current production process used to manufacture a particular machine part is known (from past experience) to produce parts which are unacceptable, in the sense that they require further machining, 35% of the time. A new production process has been developed with the hope that it will reduce the chance of producing unacceptable parts. In this example p denotes the probability that a part produced using the new production process will be unacceptable and our goal is to decide whether this probability is less than .35. A sample of 200 parts was produced using the new production process and 54 of these parts were found to be unacceptable. The SAS output for this example is provided in Figure 9.20. In this example, the *P*-value for testing $H_0: p \ge .35$ versus $H_1: p < .35$ is .0088 and we concluded that there is very strong evidence that the new production process is superior in the sense that the probability of producing an unacceptable part is less than .35. Using the upper endpoint of the 90% confidence interval in the SAS output of Figure 9.20, we can conclude that we are 95% confident that the probability that a part produced using the new production process will be unacceptable is no larger than .3245. Since this value is less than .35, the 95% confidence upper bound leads to the conclusion that the new production process is superior to the old process and indicates the extent of the improvement.

	mach	nine	e part	s e	xampl	е					
	The	e FR	EQ P	roc	edure						
case=1											
outcome	Freque	ncy	ncy Percent		Cumula Freque		Cumulative Percent				
unacceptable	54		27	.00		54	27.00				
acceptable		146		.00	200		100.00				
	Binomial Proportion				tion						
	ou	outcome = unacceptable									
	Pro	oportion 0.2700									
	ASE		0.0314								
	Confidence Limits for the Binomial Proportion										
	I										
	Туре	e 90% Confidence Limits									
	Wilson	0.2217 0.32				245					
	Test										
	ASE	SE under H0 0.0337				1					
	z	-2.372		-2.3720	1						
	One-	sided	sided Pr < Z 0.0088			1					
	Two-	sided	Pr >	zi	0.0177	1					

Figure 9.20 SAS output for the machine parts example

For completeness, we will now provide detailed derivations of 95% lower and upper confidence bounds. The probability that a standard normal variable Z takes on a value less than 1.645 is equal to .95, $P(Z \le 1.645) = .95$. That is, when we observe the value of a standard normal variable Z, 95% of the time we will find that $Z \le 1.645$. Graphically this means that the area under the standard normal density curve over the interval from $-\infty$ to 1.645 is .95. Thus, for sufficiently large values of n we have the approximation,

$$P\left[\frac{\hat{p}-p}{\mathrm{SE}(\hat{p})} \le 1.645\right] = .95.$$

Note that this indicates that 95% of the time when a simple random sample is selected and \hat{p} is computed the observed value of \hat{p} will be between zero and $p + 1.645 \text{SE}(\hat{p})$, *i.e.*, \hat{p} will be no more than 1.645 population standard error units above p. We will refer to the interval from zero to $p + 1.645 \text{SE}(\hat{p})$ as the lower 95% interval of the distribution of \hat{p} , since it will contain the observed value of \hat{p} 95% of the time.

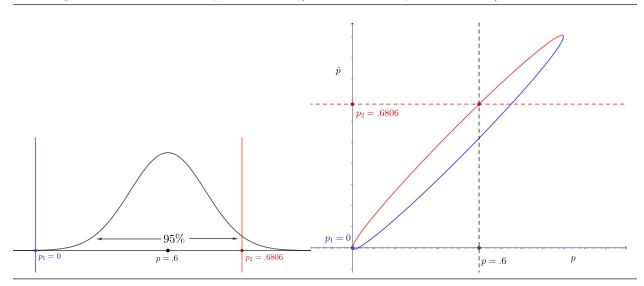
The plots in Figure 9.21 show how the lower 95% interval of the distribution of \hat{p} depends on the value of p. On the left we have a representation of the lower 95% interval, the interval from $p_1 = 0$ to p_2 . The plot on the right shows how p_2 depends on p. We want to determine the values of p which yield lower 95% intervals which contain \hat{p}_{obs} . To do this

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we need to use the graph on the right of Figure 9.21 in the other direction, as shown in Figure 9.22. In Figure 9.22 a horizontal line is drawn at \hat{p}_{obs} and its intersection, p_L , with the upper curve is indicated. Notice that p_L is the smallest value of p for which the lower 95% interval of the distribution of \hat{p} contains \hat{p}_{obs} . (Figure 9.22 is drawn for n = 100 and $\hat{p}_{obs} = .55$. In this case, $p_L = .4679$.) Thus, if we draw a vertical line, as in Figure 9.21, at any value of p between p_L and $p_U = 1$, then the corresponding lower 95% interval of the distribution of \hat{p}_{obs} . Note that p_L is the 95% confidence lower bound for p.

Figure 9.21 The plot on the left shows the lower 95% interval of the distribution of \hat{p} for n = 100 and p = .6.

The upper (red) curve in the plot on the right shows the upper endpoint, $p + 1.645\sqrt{p(1-p)/n}$, of the lower 95% interval of the distribution of \hat{p} as a function of p for n = 100. The upper endpoint for the case p = .6 is indicated by the line marking the intersections at $p_2 = .6806$. (The lower endpoint is zero.)



In order to determine the value of p_L we simply set $p + 1.645 \text{SE}(\hat{p})$ equal to \hat{p}_{obs} and solve for p (draw the horizontal line as in Figure 9.22 and project down at the intersection with the upper (red) curve).

An analogous argument starting with the upper 95% interval of the distribution of \hat{p} , *i.e.*, the interval from $p - 1.645 \text{SE}(\hat{p})$ to one, leads to a 95% confidence upper bound for p. In this case, as shown in Figure 9.23, we start with the upper 95% interval of the distribution of \hat{p} , the interval from $p_1 = p - 1.645 \text{SE}(\hat{p})$ to one. The graph on the right in Figure 9.23 shows how the upper 95% interval of the distribution of \hat{p} depends on value of p.

Figure 9.22 The smallest value of p, p_L , for which the lower 95% interval of the distribution of \hat{p} contains \hat{p}_{obs} . (The interval goes from p_L to $p_U = 1$.) This example is drawn for n = 100 and $\hat{p}_{obs} = .55$. Here $p_L = .4679$.

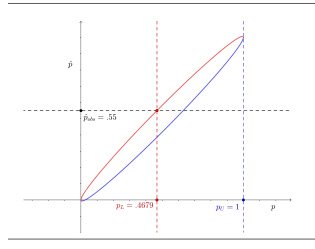
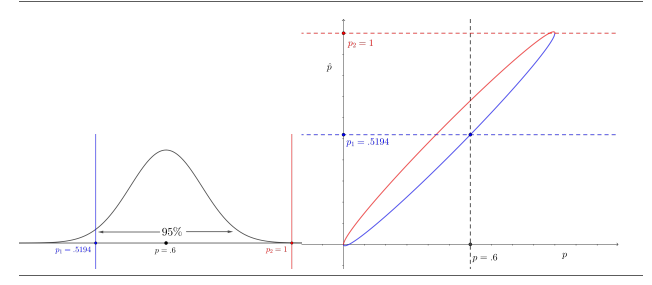


Figure 9.23 The plot on the left shows the upper 95% interval of the distribution of \hat{p} for n = 100 and p = .6.

The lower (blue) curve in the plot on the right shows the lower endpoint, $p - 1.645\sqrt{p(1-p)/n}$, of the upper 95% interval of the distribution of \hat{p} as a function of p for n = 100. The lower endpoint for the case p = .6 is indicated by the line marking the intersections at $p_1 = .5194$. (The upper endpoint is one.)



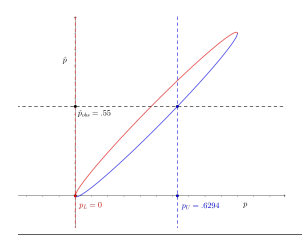
In Figure 9.24 a horizontal line is drawn at \hat{p}_{obs} and its intersection, p_U , with the lower curve is indicated. Notice that p_U is the largest value of p for which the upper central 95% interval of the distribution of \hat{p} contains \hat{p}_{obs} . (Figure 9.24 is drawn for n = 100 and

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 $\hat{p}_{obs} = .55$. In this case, $p_U = .6294$.) Thus, if we draw a vertical line, as in Figure 9.23, at any value of p between $p_L = 0$ and p_U , then the corresponding upper 95% interval of the distribution of \hat{p} contains \hat{p}_{obs} . Note that p_U is the 95% confidence upper bound for p.

In order to determine the value of p_U we simply set $p - 1.645 \text{SE}(\hat{p})$ equal to \hat{p}_{obs} and solve for p (draw the horizontal line as in Figure 9.24 and project down at the intersection with the lower (blue) curve).

Figure 9.24 The largest value of p, p_U , for which the upper 95% interval of the distribution of \hat{p} contains \hat{p}_{obs} . (The interval goes from $p_L = 0$ to p_U .) This example is drawn for n = 100 and $\hat{p}_{obs} = .55$. Here $p_U = .6294$.



9.6 Problems

The problem sets and solutions can be found on the book web page at https://www.ucs.louisiana.edu/~jcb0773/.

toc

10 Inference for Two Proportions

10.1 Introduction

In Chapter 9 we developed the basic theory of statistical inference in the context of inference for a population proportion. We will now discuss how these inferential methods confidence intervals and hypothesis tests — can be extended to make inferences about the relationship between two population proportions. More specifically, we will discuss methods for making inferences about the difference $p_1 - p_2$ between two population proportions p_1 and p_2 . The inferential methods for a single proportion p discussed in Chapter 9 are based on a large sample size normal approximation to the sampling distribution of \hat{p} . The inferential methods we will discuss in this chapter are based on an analogous large sample size normal approximation to the sampling distribution of $\hat{p}_1 - \hat{p}_2$.

In some applications there are two actual physical dichotomous populations so that p_1 denotes the population success proportion for population one and p_2 denotes the population success proportion for population two. In other applications, such as randomized comparative experiments p_1 and p_2 denote hypothetical population success probabilities corresponding to two treatments.

We will assume that the data correspond to two independent sequences of Bernoulli trials: a sequence of n_1 Bernoulli trials with population success probability p_1 and an independent sequence of n_2 Bernoulli trials with population success probability p_2 . The assumption that these are independent sequences of Bernoulli trials means that the outcomes of all $n_1 + n_2$ trials are independent. When sampling from physical populations these assumptions are equivalent to assuming that the data consist of two independent simple random samples (of sizes n_1 and n_2) selected with replacement from dichotomous populations with population success proportions p_1 and p_2 . In this context the assumption of independence basically means that the method used to select the random sample from the first population is not influenced by the method used to select the random sample from the second population, and vice versa.

The box of balls model for the present situation requires two boxes of balls, box one and box two. As we did in Chapter 9, we assume that each box contains balls some of which are red (successes) and the rest of which are blue (failures). The population success

toc

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proportion, p_1 , is the proportion of red balls in the box one. Similarly, the population success proportion, p_2 , is the proportion of red balls in the box two.

We will denote the sample sizes by n_1 and n_2 . The sample success proportion, \hat{p}_1 , is the proportion of red balls in the sample of n_1 balls selected from box one. Similarly, the sample success proportion, \hat{p}_2 , is the proportion of red balls in the sample of n_2 balls selected from box two. As noted above we will assume that these random samples are selected independently of each other so that the sample proportions \hat{p}_1 and \hat{p}_2 , when viewed as random variables, are independent.

In this model, a ball is a unit, the two boxes of balls are the populations, selecting a red ball is a success, the proportions of red balls the boxes, p_1 and p_2 , are the parameters (the population success proportions), and the proportions of red balls in the samples, \hat{p}_1 and \hat{p}_2 , are the statistics (the sample success proportions).

10.2 The sampling distribution and the normal approximation toc

The observed success proportions \hat{p}_1 and \hat{p}_2 are the obvious estimates of the two population success proportions p_1 and p_2 ; and, the difference $\hat{p}_1 - \hat{p}_2$ between these observed success proportions is the obvious estimate of $p_1 - p_2$, the difference between the two population success proportions. The behavior of $\hat{p}_1 - \hat{p}_2$ as an estimator of $p_1 - p_2$ can be determined from its sampling distribution. As you might expect, since \hat{p}_1 and \hat{p}_2 are unbiased estimators of p_1 and p_2 , $\hat{p}_1 - \hat{p}_2$ is an unbiased estimator of $p_1 - p_2$. Thus the sampling distribution of $\hat{p}_1 - \hat{p}_2$ has mean equal to $p_1 - p_2$. The standard deviation of the sampling distribution of $\hat{p}_1 - \hat{p}_2$ is the **population standard error** of $\hat{p}_1 - \hat{p}_2$

SE
$$(\hat{p}_1 - \hat{p}_2) = \sqrt{\frac{p_1(1-p_1)}{n_1} + \frac{p_2(1-p_2)}{n_2}}.$$

Notice that the population variance $\operatorname{var}(\hat{p}_1 - \hat{p}_2)$, *i.e.*, the square of $\operatorname{SE}(\hat{p}_1 - \hat{p}_2)$, is equal to the sum of the population variance of \hat{p}_1 and the population variance of \hat{p}_2 . This property is a consequence of our assumption that the random samples are independent. This expression for the standard error of the difference between two sample success proportions is not appropriate if the random samples are not independent.

As noted earlier, we will consider inferential methods based on a large sample size normal approximation to the sampling distribution of $\hat{p}_1 - \hat{p}_2$. This normal approximation is

analogous to the normal approximation to the sampling distribution of \hat{p} of Section 9.2. In the present context the normal approximation simply says that, when both n_1 and n_2 are large, the standardized value of $\hat{p}_1 - \hat{p}_2$, obtained by subtracting the population difference $p_1 - p_2$ and dividing by the population standard error of $\hat{p}_1 - \hat{p}_2$, behaves in approximate accordance with the standard normal distribution.

More formally, the normal approximation to the sampling distribution of $\hat{p}_1 - \hat{p}_2$ says that, for large values of n_1 and n_2 , the standardized value of $\hat{p}_1 - \hat{p}_2$ obtained by subtracting the difference between the population proportions $p_1 - p_2$ from $\hat{p}_1 - \hat{p}_2$ and dividing this difference by the population standard error of $\hat{p}_1 - \hat{p}_2$, behaves in approximate accordance with the standard normal distribution. That is, for large values of n_1 and n_2 the quantity

$$Z = \frac{\hat{p}_1 - \hat{p}_2 - (p_1 - p_2)}{\operatorname{SE}(\hat{p})}$$

behaves like a standard normal variable. This means, as will be shown below, that we can use an area under the standard normal density curve (a probability in terms of Z) to approximate an area in the probability histogram of the sampling distribution of $\hat{p}_1 - \hat{p}_2$ (a probability in terms of $\hat{p}_1 - \hat{p}_2$). The relationship between $\hat{p}_1 - \hat{p}_2$ and $p_1 - p_2$ indicated by this expression for Z and the normal distribution itself allow us to use $\hat{p}_1 - \hat{p}_2$ to make formal, quantifiable inferences about $p_1 - p_2$.

10.3 Estimation of
$$p_1 - p_2$$
 toc

The starting point for using this normal approximation to construct a 95% confidence interval estimate of the difference $p_1 - p_2$ between the two population success proportions is the approximate probability statement

$$P\left(\left|(\hat{p}_1 - \hat{p}_2) - (p_1 - p_2)\right| \le 1.96 \cdot \operatorname{SE}(\hat{p}_1 - \hat{p}_2)\right) \approx .95.$$

This probability statement indicates that the probability that the actual difference $p_1 - p_2$ is within $1.96 \cdot \text{SE}(\hat{p}_1 - \hat{p}_2)$ units of the observed difference $\hat{p}_1 - \hat{p}_2$ is approximately .95. As was the case with the analogous interval for one proportion, this interval is not computable, since it involves the population standard error $\text{SE}(\hat{p}_1 - \hat{p}_2)$ which depends on the unknown parameters p_1 and p_2 and is therefore also unknown.

The method we used to derive the Wilson confidence interval for a single proportion will not work in the present context. Therefore, in the present context we will consider a

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confidence interval estimate of the difference $p_1 - p_2$ based on the estimated difference $\hat{p}_1 - \hat{p}_2$ and the estimated standard error of $\hat{p}_1 - \hat{p}_2$

$$\widehat{SE}(\hat{p}_1 - \hat{p}_2) = \sqrt{\frac{\hat{p}_1(1 - \hat{p}_1)}{n_1} + \frac{\hat{p}_2(1 - \hat{p}_2)}{n_2}}$$

We will refer to this estimated standard error as the standard error for estimation.

For this application the confidence interval is centered at the estimate $\hat{p}_1 - \hat{p}_2$. The interval of plausible values for $p_1 - p_2$ is obtained by subtracting from and adding to this estimate a quantity know as the margin of error. The margin of error of $\hat{p}_1 - \hat{p}_2$ is obtained by multiplying this estimated standard error by a suitable constant k. (Recall that: for a 95% confidence level k = 1.96, for a 90% confidence level k = 1.645, and for a 99% confidence level k = 2.576.) The 95% margin of error of $\hat{p}_1 - \hat{p}_2$ is

$$ME(\hat{p}_1 - \hat{p}_2) = 1.96 \cdot \widehat{SE}(\hat{p}_1 - \hat{p}_2)$$

and the interval from $(\hat{p}_1 - \hat{p}_2) - \text{ME}(\hat{p}_1 - \hat{p}_2)$ to $(\hat{p}_1 - \hat{p}_2) + \text{ME}(\hat{p}_1 - \hat{p}_2)$ is a 95% confidence interval estimate of the difference $p_1 - p_2$. Thus we can claim that we are 95% confident that the difference $p_1 - p_2$ between the population success proportions is between

$$(\hat{p}_1 - \hat{p}_2) - \operatorname{ME}(\hat{p}_1 - \hat{p}_2)$$
 and $(\hat{p}_1 - \hat{p}_2) + \operatorname{ME}(\hat{p}_1 - \hat{p}_2)$.

Recall that it is the estimate $\hat{p}_1 - \hat{p}_2$ and the margin of error $ME(\hat{p}_1 - \hat{p}_2)$ which vary from sample to sample. Therefore, the 95% confidence level applies to the method used to generate the confidence interval estimate.

Example 10.1. Rural versus urban voter preferences. Suppose that a polling organization has separate listings of all the registered voters in a large rural district and all the registered voters in a large urban district and wishes to compare the proportions of voters in these districts who favor a proposition which is to appear on an upcoming election ballot. Let p_1 denote the proportion of all registered voters in the rural district who favor the proposition at the time of the poll and let p_2 denote the proportion of all registered voters in the urban district who favor the proposition at the time of the poll. (As noted above, in terms of the box of balls analogy of Chapter 9, we now have two boxes of balls with p_1 denoting the proportion of red balls in box one and p_2 denoting the proportion of red balls in box two.) The most obvious way to obtain independent random samples in this scenario is to: (1) randomly generate a set of n_1 labels for the rural district, contact the corresponding voters, and compute the estimate \hat{p}_1 for the voters in the rural district; and, (2) randomly generate a set of n_2 labels for the urban district, contact the corresponding voters, and compute the estimate \hat{p}_2 for the voters in the urban district. (Select a simple random sample of balls from box one and compute \hat{p}_1 and, independently, select a simple random sample of balls from box two and compute \hat{p}_2 .) Assuming that simple random samples are selected (with replacement or from large populations) this method clearly yields independent samples and the confidence interval method described above is valid.

Now suppose that we do not have separate listings of the rural voters and the urban voter but instead have a single listing of all registered voters in a large district which includes both rural and urban voters. In this situation we could randomly generate a set of n labels for the entire district, contact the corresponding voters, and in addition to determining whether the voter favors the proposition also determine whether the voter lives in a rural or urban area. We could then partition the simple random sample of n voters into the subsample of n_1 voters who live in a rural area and the subsample of n_2 voters who live in an urban area. (This is like labeling the balls in box one with a one, labeling the balls in box two with a two, then combining the balls in a single box, selecting a simple random sample of n balls from this box, and dividing it to get a sample of n_1 balls from box one and a sample of n_2 balls from box two.) This approach yields independent random samples but, technically (based on the formal definition), these random samples are not simple random samples, since the sample sizes n_1 and n_2 were not selected in advance. Actually this is not a problem, since it is readily verified that the samples can be viewed as independent sequences of Bernoulli trials (exactly if selection is with replacement and approximately if selection is without replacement from a large population and both subpopulations are also large). Therefore, the confidence interval method described above is also valid when this alternate method of forming independent random samples by partitioning a simple random sample is used.

Example 10.2. An opinion poll. The purpose of this example is to demonstrate the application of a 95% confidence interval for $p_1 - p_2$. To make the numbers more realistic we will use numbers from a Gallup poll conducted October 2–23, 2023. Like all such national

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polls this poll was not based on a simple random sample; it employed a complex random sampling method involving stratification and clustering.

For this poll a random sample of n = 1,009 adults, ages 18+, living in all U.S. states and the District of Columbia was selected. Therefore, we will think of our data as forming a simple random sample of n = 1,009 adults selected from the population of all adults in the U.S. and D.C. who could have been contacted by the pollsters at the time of this survey. We will concentrate our attention on the responses of these adults to the following question (Question 7 on the survey). Are you in favor of the death penalty for a person convicted of murder? For the purposes of this example we will restrict our attention to the subpopulation of adults who would have responded favor or oppose if everyone had been asked question 7. We will use the responses of the adults to four demographic questions (what is your gender, political party affiliation, race, and income) to subdivide the sample into samples from the corresponding subpopulations.

The computer output (SAS) which yields the sample proportions and confidences intervals for each part of this example is provided in the figure which follows the discussion of that part.

First, let's consider the relationship between gender and opinion about this topic. Let p_1 denote the proportion of all adult men in this population who would have responded "favor" if all such men had been asked question 7 and let p_2 denote the proportion of all adult women in this population who would have responded "favor" if all such women had been asked question 7. Of the $n_1 = 481$ men in the sample 285 responded "favor" giving $\hat{p}_1 = .5925$ while 252 of the $n_2 = 490$ women in the sample responded "favor" giving $\hat{p}_2 = .5143$. These data clearly suggest that the population proportion p_1 is greater than the population proportion p_2 , since 59.25% of the men responded "favor" while only 51.43% of the women responded this way. In this situation we are 95% confident that $p_1 - p_2$ is between .0159 and .1406. Since this entire interval is positive we can conclude that we are 95% confident that the population proportion of men who would have responded "favor" if all men had been asked exceeds the analogous population proportion of women by at least .0159 and perhaps as much as .1406. In other words, we are 95% confident that the percentage of all men who would have responded "favor" exceeds the corresponding percentage of women by between 1.59 and 14.06 percentage points.

Figure 10.1. The SAS output providing the estimates and confidence interval for the relationship between gender and opinion about the death penalty.

The FREQ Procedure				Statistics for Table of gender by response								
Frequency	Table	e of gende	er by respo	onse	Column 1 Risk Estimates							
Row Pct		respo	onse(favor o penalty)	death		Risk	ASE	95 Confiden	% ce Limits	Exact Confiden	95% ce Limits	
	gender	favor	oppose	Total	Row 1	0.5925	0.0224	0.5486	0.6364	0.5471	0.6368	
	men	285	196	481	Row 2	0.5143	0.0226	0.4700	0.5585	0.4690	0.5594	
		59.25	40.75		Total	0.5530	0.0160	0.5218	0.5843	0.5211	0.5846	
	women	252 51.43	238 48.57	490	Difference	0.0782	0.0318	0.0159	0.1406			
	Total	537	434	971			Differe	nce is (Row	1 - Row 2)			

Support for death penalty by gender: difference

Now, let's consider the relationship between political party affiliation and opinion about this topic. For this part we will restrict our attention to the subpopulations of adults who express affiliation with the Republican or Democratic Party. Let p_1 denote the proportion of all Republican adults in this population who would have responded "favor" if all such adults had been asked question 7 and let p_2 denote the proportion of all Democratic adults in this population who would have responded "favor" if all such adults had been asked question 7. Of the $n_1 = 268$ Republicans in the sample 218 responded "favor" giving $\hat{p}_1 = .8134$ while 86 of the $n_2 = 261$ Democrats in the sample responded "favor" giving $\hat{p}_2 = .3295$. These data clearly suggest that the population proportion p_1 is greater than the population proportion p_2 , since 81.34% of the Republicans responded "favor" while only 32.95% of the Democrats responded this way. In this situation we are 95% confident that $p_1 - p_2$ is between .4103 and .5576. Since this entire interval is positive we can conclude that we are 95% confident that the population proportion of Republicans who would have responded "favor" if all Republicans had been asked exceeds the analogous population proportion of Democrats by at least .4103 and perhaps as much as .5576. In other words, we are 95% confident that the percentage of all Republicans who would have responded "favor" exceeds the corresponding percentage of Democrats by between 41.03 and 55.76 percentage points.

Figure 10.2. The SAS output providing the estimates and confidence interval for the relationship between political party affiliation and opinion about the death penalty.

Frequency Row Pct	Table	of party b	by respons	e			Colu	mn 1 Risk E	stimates		
		response(favor death penalty)				Risk	ASE	95% Confidence Limits		Exact 95% Confidence Limits	
	party	favor	oppose	Total	Row 1	0.8134	0.0238	0.7668	0.8601	0.7615	0.858
	republican	218 81.34	50 18.66	268	Row 2	0.3295	0.0291	0.2725	0.3865	0.2728	0.390
	democratic	86	175	261	Total	0.5747	0.0215	0.5325	0.6168	0.5313	0.617
		32.95	67.05		Difference	0.4839	0.0376	0.4103	0.5576		
	Total	304	225	529			Differe	nce is (Row	1 - Row 2)	I	

Support for death penalty by political party: difference The FREQ Procedure

Next, let's consider the relationship between race and opinion about this topic. For this part we will restrict our attention to the subpopulations of (nonhispanic) whites and nonwhites. Let p_1 denote the proportion of all white adults in this population who would have responded "favor" if all such adults had been asked question 7 and let p_2 denote the proportion of all nonwhite adults in this population who would have responded "favor" if all such adults had been asked question 7. Of the $n_1 = 638$ whites in the sample 388 responded "favor" giving $\hat{p}_1 = .6082$ while 141 of the $n_2 = 329$ nonwhites in the sample responded "favor" giving $\hat{p}_2 = .4286$. These data clearly suggest that the population proportion p_1 is greater than the population proportion p_2 , since 60.82% of the whites in the sample responded "favor" while only 42.86% of the nonwhites in the sample responded this way. In this situation we are 95% confident that $p_1 - p_2$ is between .1140 and .2451. Since this entire interval is positive we can conclude that we are 95% confident that the population proportion of whites who would have responded "favor" if all whites had been asked exceeds the analogous population proportion of nonwhites by at least .1140 and perhaps as much as .2451. In other words, we are 95% confident that the percentage of all whites who would have responded "favor" exceeds the corresponding percentage of nonwhites by between 11.40 and 24.51 percentage points.

Statistics for Table of party by response

Statistics for Table of race by response

Figure 10.3. The SAS output providing the estimates and confidence interval for the relationship between race and opinion about the death penalty.

	The FREQ Procedure					Statist	tics for	Table of r	ace by re	sponse		
Frequency	Table	e of race	by respons	se		Column 1 Risk Estimates						
Row Pct		respo	onse(favor penalty)	death		Risk ASE 25% Confidence Limits				Exact 95% Confidence Limits		
	race	favor	oppose	Total	Row 1	0.6082	0.0193	0.5703	0.6460	0.5691	0.6462	
	white	388	250	638	Row 2	0.4286	0.0273	0.3751	0.4820	0.3744	0.4840	
		60.82	39.18		Total	0.5471	0.0160	0.5157	0.5784	0.5151	0.5788	
	nonwhite	141 42.86	188 57.14	329	Difference	0.1796	0.0334	0.1140	0.2451			
	Total	529	438	967			Differe	nce is (Row	1 - Row 2)			

Support for death penalty by race: difference

Finally, let's consider the relationship between income and opinion about this topic. For this part we will restrict our attention to the subpopulations of low or moderate income (no more than \$100,000) adults and high income (greater than \$100,000) adults. Let p_1 denote the proportion of all low or moderate income adults in this population who would have responded "favor" if all such adults had been asked question 7 and let p_2 denote the proportion of all high income adults in this population who would have responded "favor" if all such adults had been asked question 7. Of the $n_1 = 616$ low or moderate income adults in the sample 330 responded "favor" giving $\hat{p}_1 = .5357$ while 153 of the $n_2 = 274$ high income adults in the sample responded "favor" giving $\hat{p}_2 = .5584$. These data suggest that the population proportion p_1 is at least slightly smaller than the population proportion p_2 , since only 53.57% of the low or moderate income adults in the sample responded "favor" while 55.84% of the high income adults in the sample responded this way. In this situation we are 95% confident that $p_1 - p_2$ is between -.0934 and .0481. Since this interval contains values that are both negative (indicating that $p_1 < p_2$) and positive (indicating that $p_1 > p_2$) we must conclude that, based on these data, we cannot tell which of these population proportions is larger. We can conclude that we are 95% confident that the population proportion of low or moderate income adults who would have responded "favor" if all low or moderate income adults had been asked this question my be as much as .0934 lower than the analogous population proportion of high income adults, as much as .0481 higher, or anywhere within this range of values. In other words, we are 95% confident that the percentage of all low or moderate adults who would have responded "favor" could be as much as 9.34 percentage points lower than the corresponding proportion of high income adults, as much as 4.81 percentage points higher than the corresponding percentage of high income adults, or anywhere within this range of values.

Figure 10.4. The SAS output providing the estimates and confidence interval for the relationship between income and opinion about the death penalty.

Support for death penalty by income level: difference The FREO Procedure

Frequency Row Pct	Table of income by response								
		respons							
	income	favor	oppose	Total					
	lowormoderate	330 53.57	286 46.43	616					
	high	153 55.84	121 44.16	274					
	Total	483	407	890					

Statistics	for T	able	of	income	by	response
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	Column 1 Risk Estimates										
	Risk	ASE	95 Confider	% Ice Limits	Exact Confiden	: 95% ice Limits					
Row 1	0.5357	0.0201	0.4963	0.5751	0.4954	0.5757					
Row 2	0.5584	0.0300	0.4996	0.6172	0.4974	0.6181					
Total	0.5427	0.0167	0.5100	0.5754	0.5093	0.5758					
Difference	-0.0227	0.0361	-0.0934	0.0481							
	Difference is (Row 1 - Row 2)										

Another common application of this confidence interval for the difference between two population proportions is for randomized comparative experiments. Consider a randomized comparative experiment where $N = n_1 + n_2$ available units are randomly assigned to receive one of two treatments (with n_1 units assigned to treatment 1 and the remaining n_2 units assigned to treatment 2). We can imagine two hypothetical populations of responses and two population success proportions corresponding to the two treatments. The first hypothetical population is the collection of responses (S or F), corresponding to all Navailable units, which we would observe if all N available units were subjected to treatment 1 and p_1 is the proportion of successes among these units. The second hypothetical population and population success proportion p_2 are defined similarly to correspond to the responses we would observe if all N available units were subjected to treat-

The model corresponding to the assumptions we made to justify the confidence interval for $p_1 - p_2$ treats the data as if they constitute independent simple random samples selected with replacement from these two hypothetical populations. In terms of balls in a box, this means that we are assuming that we have independent simple random samples selected with replacement from two separate boxes of balls, with each box containing N balls. Clearly this model is not appropriate for this application; a more appropriate model treats the data as two dependent random samples selected without replacement from a single box of N balls. Fortunately, even though the underlying assumptions are not valid for this

application, provided that n_1 and n_2 are reasonably large, the confidence interval based on the assumptions of independent simple random samples selected with replacement works well for randomized comparative experiments.

Example 10.3. Leading questions. The wording of questions in surveys can have a major impact on the responses elicited. The effect of wording of questions was investigated in Schuman and Presser, Attitude measurement and the gun control paradox, *Public Opinion Quarterly*, **41** winter 1977–1978, 427–438. Two groups of adults were used to estimate the difference in response to the following two versions of a question regarding gun control.

1. Would you favor or oppose a law which would require a person to obtain a police permit before he could buy a gun?

2. Would you favor a law which would require a person to obtain a police permit before he could buy a gun, or do you think that such a law would interfere too much with the right of citizens to own guns?

We might expect the second version of the question, with the added remark about the right of citizens to own guns, to lead to less responses in favor of requiring a permit.

This study was conducted in 1976. The researchers began with a group of 1263 adults which had been obtained by a random sampling method for a survey conducted by the Survey Research Center of the University of Michigan. These 1263 adults were randomly divided into two groups with 642 adults in the first group and 621 adults in the second group. The adults in the first group were asked to respond to the first version of the gun control question and the adults in the second group were asked to respond to the second version of the gun control question. Twenty-seven adults in the first group and 36 adults in the second group would not respond to the question. Therefore, we will restrict our attention to the 1200 adults who were willing to respond to a question about gun control, and we will use the $n_1 = 615$ adults in the first group and the $n_2 = 585$ adults in the second group who responded to the question as our samples.

In this randomized comparative experiment the group of available units is the group of 1200 adults who were willing to respond to a question about gun control in 1976. Let p_1 denote the proportion of these 1200 adults who would respond "favor" (in 1976) if all 1200 were asked the first question and let p_2 denote the proportion of these 1200 adults who would respond "favor" (in 1976) if all 1200 were asked the first question and let p_2 denote the proportion of these 1200 adults who would respond "favor" (in 1976) if all 1200 were asked the second question. Our goal

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is to estimate the difference $p_1 - p_2$ between these proportions. You do not have to use the formulae given above to perform the computations given below; you can find all of these values in the SAS output given in Figure 10.5. When the study was conducted 463 of the 615 adults in the first group responded "favor" and 403 of the 585 adults in the second group responded "favor". The observed proportions of adults who respond "favor" are $\hat{p}_1 = .7528$ and $\hat{p}_2 = .6889$ giving a difference of $\hat{p}_1 - \hat{p}_2 = .0639$. The standard error is $\widehat{SE}(\hat{p}_1 - \hat{p}_2) = .02586$ and the margin of error is $ME(\hat{p}_1 - \hat{p}_2) = .0507$; therefore, we are 95% confident that the difference $p_1 - p_2$ is between .0639 - .0507 = .0132 and .0639 + .0507 = .1146. That is, we are 95% confident that modifying the first question about gun control by adding the comment about the right of citizens to own guns lowers the probability that an individual adult (from this group of 1200 adults) would respond "favor" (in 1976) by at least .0132 and at most .1146.

Figure 10.5. SAS output for the leading question example.

leading question regarding gun permits example (success = favor permit)

Frequency	Table of group by response								
Row Pct		respon	onse(favor permit)						
	group	favor	not	Total					
	leadingQ	403 68.89	182 31.11	585					
	neutralQ	463 75.28	152 24.72	<mark>615</mark>					
	Total	866	334	1200					

The FREQ Procedure

		Colu	mn 1 Risk E	stimates			Risk Differe	ence Test
			95	%	Exact	95%	H0: P1 - P2 = 0	Wald Method
	Risk	ASE		ice Limits		ce Limits	Risk Difference	-0.0640
Row 1	0.6889	0.0191	0.6514	0.7264	0.6496	0.7262	ASE (H0)	0.0259
Row 2	0.7528	0.0174	0.7188	0.7869	0.7168	0.7865	z	-2.4709
Total	0.7217	0.0129	0.6963	0.7470	0.6954	0.7469	One-sided Pr < Z	0.0067
Difference	-0.0640	0.0259	-0.1146	-0.0133			Two-sided Pr > Z	0.0135
		Differe	nce is (Row	1 - Row 2)			Column 1 (resp	onse = favor)

In summary, we estimate that, in 1976, about 75.28% of these 1200 adults would respond "favor" if asked the first question and we estimate that, if these same people had instead been asked the second question with the comment about the right of citizens to own guns,

then we would see a reduction of this percentage in the range of 1.32 to 11.46 percentage points. Thus we find sufficient evidence to conclude that the added comment has the anticipated effect of lowering the percentage who would respond "favor"; note, however, that this reduction might be as small as 1.32 percentage points, as large as 11.46 percentage points, or anywhere within this range. As we noted above these 1200 adults can be viewed as a random sample from the population of adults sampled by the University of Michigan researchers who would respond to a question about gun control, thus it is reasonable to claim that this inference applies to this entire population of adults (in 1976) not just these 1200.

10.4 Testing a hypothesis about $p_1 - p_2$ toc

In this section we will consider hypothesis tests for hypotheses relating two population success proportions p_1 and p_2 . The tests we consider are based on the same normal approximation to the sampling distribution of $\hat{p}_1 - \hat{p}_2$ that we used for confidence estimation. Thus we will assume that the data on which the hypothesis test is based correspond to two independent simple random samples of sizes n_1 and n_2 , selected with replacement, from dichotomous populations with population success proportions p_1 and p_2 , or equivalently, that the data correspond to the outcomes of two independent sequences of n_1 and n_2 Bernoulli trials with success probabilities p_1 and p_2 . However, as with confidence estimation, for practical purposes, we do not need to worry about whether the samples are selected with or without replacement, provided both of the populations are very large; and, these tests are also applicable to randomized comparative experiments.

Many hypotheses about the relationship between the population proportions p_1 and p_2 can be expressed as hypotheses about the relationship between $p_1 - p_2$ and zero, e.g., $p_1 > p_2$ is equivalent to $p_1 - p_2 > 0$. Therefore, we will consider tests which are based on a suitably standardized value of the difference $\hat{p}_1 - \hat{p}_2$ between the observed success proportions.

The *P*-value for a hypothesis about the relationship between a single proportion p and a hypothesized value p_0 is computed under the assumption that $p = p_0$, therefore, we used $p = p_0$ in the standard error of \hat{p} for the *Z*-statistic of the test. The *P*-value for a hypothesis about the relationship between p_1 and p_2 is computed under the assumption that $p_1 = p_2$, therefore, we need to determine a suitable standard error of $\hat{p}_1 - \hat{p}_2$ (the standard error for testing) under this assumption. Notice that $p_1 = p_2$ ($p_1 - p_2 = 0$)

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specifies a common value for p_1 and p_2 but does not specify what this common value is, e.g., we might have $p_1 = p_2 = .5$ or $p_1 = p_2 = .1$. When $p_1 = p_2$, \hat{p}_1 and \hat{p}_2 are estimates of the same population success proportion. This suggests that we can pool or combine the information in the two random samples to obtain a pooled estimate, \hat{p} , of this common population success proportion. This pooled estimate \hat{p} can then be used to get an estimate of $SE(\hat{p}_1 - \hat{p}_2)$ that is suitable for use in the hypothesis test. If we let p denote the common population success proportion under the assumption that $p_1 = p_2$, then the population standard error of $\hat{p}_1 - \hat{p}_2$ simplifies to

$$\operatorname{SE}(\hat{p}_1 - \hat{p}_2) = \sqrt{p(1-p)\left(\frac{1}{n_1} + \frac{1}{n_2}\right)}.$$

Replacing p in this population standard error by the pooled estimate \hat{p} gives the standard error for testing

$$\widehat{SE}(\hat{p}_1 - \hat{p}_2) = \sqrt{\hat{p}(1 - \hat{p})\left(\frac{1}{n_1} + \frac{1}{n_2}\right)},$$

where

$$\hat{p} = \frac{\text{the total number of successes in both samples}}{\text{the total number of observations in both samples}} = \frac{n_1 \hat{p}_1 + n_2 \hat{p}_2}{n_1 + n_2}.$$

When testing $H_0: p_1 \leq p_2$ versus $H_1: p_1 > p_2$ values of $\hat{p}_1 - \hat{p}_2$ which are sufficiently larger than zero provide evidence against the null hypothesis $H_0: p_1 \leq p_2$ and in favor of the research hypothesis $H_1: p_1 > p_2$. Thus large (positive) values of

$$Z_{calc} = \frac{\hat{p}_1 - \hat{p}_2}{\widehat{\mathrm{SE}}(\hat{p}_1 - \hat{p}_2)},$$

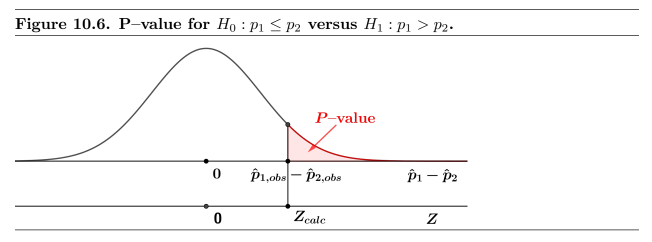
where $\widehat{SE}(\hat{p}_1 - \hat{p}_2)$ denotes the standard error for testing, favor the research hypothesis and the *P*-value is the probability that a standard normal variable takes on a value at least as large as Z_{calc} , *i.e.*, the *P*-value is the area under the standard normal density curve to the right of Z_{calc} .

The steps for performing a hypothesis test for

$$H_0: p_1 \le p_2 \quad \text{versus} \quad H_1: p_1 > p_2$$

are summarized below.

1. Use a suitable calculator or computer program to find the P-value = $P(Z \ge Z_{calc})$, where Z denotes a standard normal variable and Z_{calc} is as defined above. This P-value is the area under the standard normal density curve to the right of Z_{calc} as shown in Figure 10.6.



2a. If the *P*-value is small enough (less than .05 for a test at the 5% level of significance), conclude that the data favor $H_1: p_1 > p_2$ over $H_0: p_1 \leq p_2$. That is, if the *P*-value is small enough, then there is sufficient evidence to conclude that the population proportion p_1 is greater than the population success proportion p_2 .

2b. If the *P*-value is not small enough (is not less than .05 for a test at the 5% level of significance), conclude that the data do not favor $H_1: p_1 > p_2$ over $H_0: p_1 \leq p_2$. That is, if the *P*-value is not small enough, then there is not sufficient evidence to conclude that the population proportion p_1 is greater than the population success proportion p_2 .

When testing $H_0: p_1 \ge p_2$ versus $H_1: p_1 < p_2$ values of $\hat{p}_1 - \hat{p}_2$ which are sufficiently smaller than zero provide evidence against the null hypothesis $H_0: p_1 \ge p_2$ and in favor of the research hypothesis $H_1: p_1 < p_2$. Thus sufficiently negative values of Z_{calc} (as defined above) favor the research hypothesis and the *P*-value is the probability that a standard normal variable takes on a value no larger than Z_{calc} , *i.e.*, the *P*-value is the area under the standard normal density curve to the left of Z_{calc} .

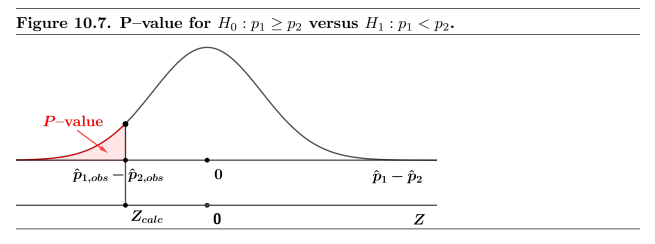
The steps for performing a hypothesis test for

$$H_0: p_1 \ge p_2 \quad \text{versus} \quad H_1: p_1 < p_2$$

are summarized below.

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1. Use a suitable calculator or computer program to find the P-value = $P(Z \leq Z_{calc})$, where Z denotes a standard normal variable and Z_{calc} is as defined above. This P-value is the area under the standard normal density curve to the left of Z_{calc} as shown in Figure 10.7.



2a. If the *P*-value is small enough (less than .05 for a test at the 5% level of significance), conclude that the data favor $H_1: p_1 < p_2$ over $H_0: p_1 \ge p_2$. That is, if the *P*-value is small enough, then there is sufficient evidence to conclude that the population proportion p_1 is less than the population success proportion p_2 .

2b. If the *P*-value is not small enough (is not less than .05 for a test at the 5% level of significance), conclude that the data do not favor $H_1: p_1 < p_2$ over $H_0: p_1 \ge p_2$. That is, if the *P*-value is not small enough, then there is not sufficient evidence to conclude that the population proportion p_1 is less than the population success proportion p_2 .

When testing $H_0: p_1 = p_2$ versus $H_1: p_1 \neq p_2$ values of $\hat{p}_1 - \hat{p}_2$ which are sufficiently far away from zero in either direction provide evidence against the null hypothesis $H_0: p_1 = p_2$ and in favor of the research hypothesis $H_1: p_1 \neq p_2$. Thus sufficiently large values of the absolute value of Z_{calc} (as defined above) favor the research hypothesis and the *P*-value is the probability that a standard normal variable takes on a value below $-|Z_{calc}|$ or above $|Z_{calc}|, i.e.$, the *P*-value is the combined area under the standard normal density curve to the left of $-|Z_{calc}|$ and to the right of $|Z_{calc}|$.

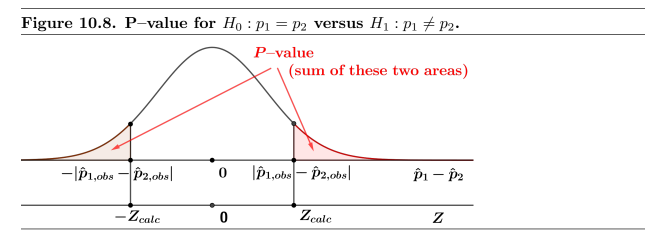
The steps for performing a hypothesis test for

$$H_0: p_1 = p_2$$
 versus $H_1: p_1 \neq p_2$

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are summarized below.

1. Use a suitable calculator or computer program to find the P-value = $P(|Z| \ge |Z_{calc}|) = P(Z \le -|Z_{calc}|) + P(Z \ge |Z_{calc}|)$, where Z denotes a standard normal variable and Z_{calc} is as defined above. This P-value is the combined area under the standard normal density curve to the left of $-|Z_{calc}|$ and to the right of $|Z_{calc}|$ as shown in Figure 10.8.



2a. If the *P*-value is small enough (less than .05 for a test at the 5% level of significance), conclude that the data favor $H_1 : p_1 \neq p_2$ over $H_0 : p_1 = p_2$. That is, if the *P*-value is small enough, then there is sufficient evidence to conclude that the population success proportions p_1 and p_2 are different.

2b. If the *P*-value is not small enough (is not less than .05 for a test at the 5% level of significance), conclude that the data do not favor $H_1: p_1 \neq p_2$ over $H_0: p_1 = p_2$. That is, if the *P*-value is not small enough, then there is not sufficient evidence to conclude that the population success proportions p_1 and p_2 are different.

Example 10.4. An HIV vaccine trial. This example is based on a study described in Flynn et al., Placebo-controlled phase 3 trial of a recombinant glycoprotein 120 vaccine to prevent HIV-1 infection, J. of Infect. Dis., **191** Mar. 1, 2005, 654–665. A double-blind randomized trial was conducted to investigate the effect of an rgp120 vaccine among men who have sex with men and among women at high risk for heterosexual transmission of type 1 HIV. A group of 5403 volunteers (5095 men and 308 women) was randomly divided into two groups (a vaccine group ($n_1 = 3598$) and a control group ($n_2 = 1805$)). Each volunteer received 7 injections of either placebo or vaccine over a 30 month period. These individuals were tracked for a period of 3 years to see whether they developed HIV-1.

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We can envision two hypothetical populations based on this group of 5403 individuals and these two experimental treatments. Since these 5403 volunteers do not form a random sample from some well defined population of people at high risk for developing HIV–1 we should restrict our inferences to these 5403 volunteers. Let p_1 denote the proportion of this group of 5403 volunteers who would develop HIV–1 within 3 years if all 5403 volunteers were given the vaccine. Let p_2 denote the proportion of this group of 5403 volunteers who would develop HIV–1 within 3 years if all 5403 volunteers were given the placebo. We can also think of these proportions as the probabilities that one of these 5403 volunteers would develop HIV–1 within 3 years if he or she was treated with the vaccine (p_1) or if he or she was treated with the placebo (p_2). In terms of these parameters our research hypothesis is $H_1 : p_1 < p_2$ (the vaccine reduces the risk of developing HIV–1) and our null hypothesis is $H_0 : p_1 \ge p_2$ (the vaccine does not reduce the risk of developing HIV–1).

Figure 10.9. SAS output for analysis of the effects of the HIV vaccine.

HIV vaccine trial (success = positive)

The FREQ	Procedure
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Frequency Row Pct	Tak	Table of group by response						
ROWPCI		response(HIV status)						
	group	positive	negative	Total				
	vaccine	241 6.70	3357 93.30	3598				
	control	126 6.98	1679 93.02	1805				
	Total	367	5036	5403				

		Colu	mn 1 Risk E	stimates			Risk Differe	ence Test
		Colu	95		Exact	05%	H0: P1 - P2 = 0	Wald Method
	Risk	ASE		ice Limits		ice Limits	Risk Difference	-0.0028
Row 1	0.0670	0.0042	0.0588	0.0752	0.0590	0.0756	ASE (H0)	0.0073
Row 2	0.0698	0.0060	0.0581	0.0816	0.0585	0.0826	z	-0.3892
Total	0.0679	0.0034	0.0612	0.0746	0.0614	0.0750	One-sided Pr < Z	0.3486
Difference	-0.0028	0.0073	-0.0171	0.0115			Two-sided Pr > Z	0.6971
		Differe	nce is (Row	1 - Row 2)			Column 1 (respo	nse = positive)

By the end of the 3 years, 126 of the 1805 individuals treated with the placebo developed HIV–1 while 241 of the 3598 individuals treated with the vaccine developed HIV–1. The observed proportions are $\hat{p}_1 = .0670$ and $\hat{p}_2 = .0698$, and the difference is $\hat{p}_1 - \hat{p}_2 = -.0028$.

The fact that this difference is negative $(\hat{p}_1 \text{ is less than } \hat{p}_2)$ shows that there is some evidence in favor of the research hypothesis $p_1 < p_2$. We need to determine whether observing a difference of -.0028, with samples of size $n_1 = 3598$ and $n_2 = 1805$, is sufficiently surprising under the assumption that $p_1 \ge p_2$ to allow us to reject this null hypothesis as untenable. When we use the standard error for testing to standardize this difference we get $Z_{calc} = -.3892$. The corresponding P-value= $P(Z \le -.38929) = .3486$ is quite large. In words, this means that (for these sample sizes) if the null hypothesis was true (p_1 was actually no less than p_2), then we would observe a difference this far below zero about 34.86% of the time. In other words, for the volunteers used in this study, these data do not provide enough evidence to allow us to claim that this vaccine is better than a placebo.

Example 10.5. Scotland coronary prevention study. This example is based on the West of Scotland Coronary Prevention Study as described in Shepherd *et al.*, Prevention of coronary heart disease with pravastatin in men with hypercholesterolemia, *New England Journal of Medicine*, **333** Nov. 16, 1995, 1994–1307, and Ford *et al.*, Long-term follow-up of the West of Scotland coronary prevention study, *New England Journal of Medicine*, **357** Oct. 11, 2007, 1477–1486. The primary goal of this study was to determine whether the administration of pravastatin to middle–aged men with high cholesterol levels and no history of myocardial infarction over a period of five years reduces the risk of coronary events. In this context a coronary event is defined as a nonfatal myocardial infarction or death from coronary heart disease. A group of 6595 men, aged 45 to 64 years, with high plasma cholesterol levels (mean 272 mg/dl) was randomly divided into two groups (a control group and a treatment group). The 3302 men in the treatment group received 40 mg of pravastatin daily while the 3293 men in the control group received a placebo. All of the men were given smoking cessation and dietary advice throughout the study.

We can envision two hypothetical populations based on this group of 6595 men and these two experimental treatments. Since these 6595 men do not form a random sample from some well defined population of middle–aged men with high cholesterol levels we should restrict our inferences to these 6595 men. However, the investigators examined these men to see if extrapolations beyond this group may be reasonable and they concluded that: "The subjects in this study were representative of the general population in terms of socioeconomic status and risk factors (Table 1). Their plasma cholesterol levels were in the highest quartile of the range found in the British population. A number had evidence of

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minor vascular disease, and in order to make the findings of the trial applicable to typical middle-aged men with hypercholesterolemia, they were not excluded."

Let p_1 denote the proportion of this group of 6595 men who would experience a coronary event (as defined above) if all 6595 men were subjected to the five year pravastatin treatment. Let p_2 denote the proportion of this group of 6595 men who would experience a coronary event if all 6595 men were subjected to the five year placebo treatment. We can also think of these proportions as the probabilities that one of these 6595 men would have a coronary event within five years if he was treated with pravastatin (p_1) or if he was treated with placebo (p_2) . In terms of these parameters our research hypothesis is $H_1 : p_1 < p_2$ (pravastatin reduces the risk of a coronary event) and our null hypothesis is $H_0 : p_1 \ge p_2$ (pravastatin does not reduce the risk of a coronary event).

By the end of this five year trial, 174 of the 3302 men treated with pravastatin had experienced a coronary event and 248 of the 3293 men treated with a placebo had experienced a coronary event. The observed proportions are $\hat{p}_1 = .0527$ and $\hat{p}_2 = .0753$, and the difference is $\hat{p}_1 - \hat{p}_2 = -.0226$. The fact that this difference is negative $(\hat{p}_1 \text{ is less than } \hat{p}_2)$ shows that there is some evidence in favor of the research hypothesis $p_1 < p_2$. We need to determine whether observing a difference of -.0226, with samples of size $n_1 = 3302$ and $n_2 = 3293$, is sufficiently surprising under the assumption that $p_1 \ge p_2$ to allow us to reject this null hypothesis as untenable. When we use the standard error for testing to standardize this difference we get $Z_{calc} = -3.7523$. The corresponding P-value $P(Z \leq -3.7523)$ is less than .0001 (approximately 8.8×10^{-5}). In words, this means that (for these sample sizes) if the null hypothesis was true $(p_1 \text{ was actually no less than } p_2)$, then we would almost never (less than .01% of the time) observe a difference this far below zero. Therefore, these data provide very strong evidence in favor of the research hypothesis that pravastatin reduces the probability of a coronary event in the sense that the probability that one of these 6595 men would have a coronary event within five years would be lower if he was treated with pravastatin than if he was treated with placebo.

In addition to this conclusion that pravastatin reduces the probability of a coronary event we can construct a confidence interval to quantify the practical importance of this reduction. In this example we are 95% confident that $p_1 - p_2$ is between -.0344 and -.0108 $(p_2 - p_1$ is between .0108 and .0344). In summary, for these 6595 men, we have very strong evidence (*P*-value < .0001) that pravastatin reduces the risk of a coronary event (versus placebo). We estimate that about 7.53% of these men would have a coronary event if they all were treated with a placebo, and we are 95% confident that if they all were treated with pravastatin we would see a 1.08 to 3.44 percentage point reduction in this percentage. Since we are dealing with small percentages it is instructive to note that a reduction from 7.53% (\hat{p}_2) to 5.27% (\hat{p}_1) is a 30% reduction ((7.53 - 5.27)/7.53 = .3001) in the risk of a man having a coronary event. Using 7.53% as a baseline and using the confidence interval limits we are 95% confident that we would see a 14.34 percent to 45.68 percent decrease (1.08/7.53 = .1434 and 3.44/7.53 = .4568) in the risk of a man having a coronary event.

Figure 10.10. SAS output for 5 year analysis of the effects of Pravastatin.

West of Scotland coronary prevention study: 5 years (success=yes) The FREQ Procedure

Frequency Row Pct	Table of drug by response							
		response(has a coronary event)						
	drug	yes	no	Total				
	pravastatin	174 5.27	3128 94.73	3302				
	placebo	248 7.53	3045 92.47	3293				
	Total	422	6173	6595				

Column 1 Risk Estimates								Risk Difference Test		
	95% Exact 95%				H0: P1 - P2 = 0	Wald Method				
	Risk	ASE		ice Limits	Confidence Limits			Risk Difference	-0.0226	
Row 1	0.0527	0.0039	0.0451	0.0603	0.0453	0.0609		ASE (H0)	0.0060	
Row 2	0.0753	0.0046	0.0663	0.0843	0.0665	0.0849	ŀ	z	-3.7523	
Total	0.0640	0.0030	0.0581	0.0699	0.0582	0.0702	ľ	One-sided Pr < Z	<.0001	
Difference	-0.0226	0.0060	-0.0344	-0.0108				Two-sided Pr > Z	0.0002	
	Difference is (Row 1 - Row 2)						Column 1 (res	oonse = yes)		

A follow-up to this study tracked the men used in this trial for ten additional years to assess the long term effects of treatment with pravastatin. At the end of the five year trial, treatment with pravastatin or placebo ceased, and the patients returned to the care of their primary care physicians. Five years after the conclusion of the trial 38.7% of the original pravastatin group and 35.2% of the original placebo group were being treated with statin

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drugs. The purpose of the follow–up study was to assess long–term effects regardless of treatment received after the initial trial period.

For this part of the study, let p_3 denote the proportion of this group of 6595 men who would experience a coronary event within 15 years of the beginning of the initial trial if all 6595 men were subjected to the five year pravastatin treatment. Let p_4 denote the analogous proportion if all the men were subjected to the placebo treatment. In terms of these parameters our research hypothesis is $H_1: p_3 < p_4$ (pravastatin reduces the longterm risk of a coronary event) and our null hypothesis is $H_0: p_3 \ge p_4$ (pravastatin does not reduce the long-term risk of a coronary event).

By the end of the 15 year period, 390 of the 3302 men treated with pravastatin had experienced a coronary event and 509 of the 3293 men treated with a placebo had experienced a coronary event. The observed proportions are $\hat{p}_3 = .1181$ and $\hat{p}_4 = .1546$, and the difference is $\hat{p}_3 - \hat{p}_4 = -.0365$. The fact that this difference is negative (\hat{p}_3 is less than \hat{p}_4) shows that there is some evidence in favor of the research hypothesis $p_3 < p_4$. Since the sample sizes for this test are the same as for the test above and since the difference in this case is more extreme than before, we know that the *P*-value will be even smaller. In this case, when we use the standard error for testing to standardize this difference we get $Z_{calc} = -4.3136$. The corresponding P-value= $P(Z \leq -4.3146)$ is less than .0001 (approximately 8.0×10^{-6}). Therefore, these data provide very strong evidence in favor of the research hypothesis that the five year pravastatin treatment reduces the probability of a coronary event in the long-term in the sense that the probability that one of these 6595 men would have a coronary event within 15 years would be lower if he was treated with pravastatin than if he was treated with placebo. In this case we are 95% confident that p_4 exceeds p_3 by at least .0199 and perhaps as much as .0530. Here we would estimate that about 15.46% of these men would have a coronary event within 15 years if they were all given the five year placebo treatment and we are 95% confident that the five year pravastatin treatment would reduce this percentage by between 1.99 and 5.30 percentage points. Using 15.46% as a baseline and using the confidence interval limits we are 95%confident that we would see a 12.87 to 34.28 percent decrease (1.99/15.46 = .1287) and 5.30/15.46 = .3428) in the risk of a man having a coronary event.

Figure 10.11. SAS output for 15 year analysis of the effects of Pravastatin.

West of Scotland coronary prevention study: 15 years (success=yes)

The FREQ Procedure

Frequency Row Pct	Table of drug by response					
RowPct		response(has a coronary event)				
	drug	yes	no	Total		
	pravastatin	390 11.81	2912 88.19	3302		
	placebo	509 15.46	2784 84.54	3293		
	Total	899	5696	6595		

Column 1 Risk Estimates							Risk Difference Test		
	95% Exact 95%		H0: P1 - P2 = 0	Wald Method					
	Risk	ASE		ce Limits	Confidence Limits		Risk Difference	-0.0365	
Row 1	0.1181	0.0056	0.1071	0.1291	0.1073	0.1296	ASE (H0)	0.0085	
Row 2	<mark>0.1546</mark>	0.0063	0.1422	0.1669	0.1424	0.1674	z	-4.3146	
Total	0.1363	0.0042	0.1280	0.1446	0.1281	0.1448	One-sided Pr < Z	<.0001	
Difference	-0.0365	0.0084	-0.0530	-0.0199			Two-sided Pr > Z	<.0001	
	Difference is (Row 1 - Row 2)					Column 1 (res	ponse = yes)		

10.5 Problems

toc

The problem sets and solutions can be found on the book web page at https://www.ucs.louisiana.edu/~jcb0773/.

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11 Inference for a Mean

11.1 Introduction

There are many situations when we might wish to make inferences about the location of the "center" of the population distribution of a quantitative variable. The most commonly used measure of the location of the center of the distribution of a quantitative variable is the population mean.

In this chapter we will discuss inference for the mean μ of the population distribution of a continuous variable X. The basic problem we will treat is that of using a random sample of values of the continuous variable X to estimate the corresponding population mean or to test a hypothesis about this population mean.

Before we get into the details of inference, it is instructive to consider some situations where inference about a population mean could be used and the way in which we might interpret a population mean.

In some applications the population mean represents an actual physical constant such as the speed of light. Let μ denote the true value of the physical constant (such as the speed of light under well defined conditions) which we wish to estimate. Suppose that an experiment has been devised to produce a measurement X of the physical constant μ . A probability model for the distribution of X provides a model for the behavior of an observed value of X by specifying probabilities which X must satisfy. Thus, the probability model provides an explanation of the variability in X as an estimate of μ . It would be unreasonable to expect an observed value of X to be exactly equal to μ ; however, if the experiment is carefully planned and executed it would be reasonable to expect the average value of X based on a long series of replications of the experiment to be equal to μ . If this is the case, the population mean of the probability model for X will be equal to the physical constant μ and the standard deviation σ of the probability model will serve as a useful quantification of the variability of the measurement process.

When interest centers on an actual, physical population of units the population mean is the average value of the variable of interest corresponding to all of the units in the population. Imagine a large population of units, *e.g.*, a population of humans or animals. Let the continuous variable X denote a characteristic of a unit, *e.g.*, X might be some measurement of the size of a unit. For concreteness recall Example 8.1, in that example we

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considered the distribution of the heights of adult men and the heights of adult women in the United States. Let X denote the height of an adult woman and consider the population of all adult women in the United States. A probability model for the distribution of X provides a model for the behavior of an observed value of the height X of an adult woman selected at random from this population. In this situation a probability model explains the variability among the heights of the adult women in this population. Let μ denote the population mean height of all adult women in the United States, *i.e.*, let μ be the average height we would get if we averaged the heights of all the adult women in this population. In this context the population mean height is obviously not the "true height" of each adult woman; however, we can think of the height X of a particular adult woman as being equal to the population mean height μ plus or minus an adjustment for this particular woman which is due to hereditary, environmental, and other factors. The standard deviation σ of the probability model serves to quantify the variability among this population of heights.

In many applications interest centers on the population mean difference between two paired values. For example, consider a population of individuals with high cholesterol levels and a drug designed to reduce cholesterol levels. Let X_1 denote the cholesterol level of an individual before taking the drug, let X_2 denote this same individual's cholesterol level after being treated with the drug, and let $D = X_1 - X_2$ denote the difference between the two cholesterol levels (the decrease in cholesterol level). A probability model for the distribution of D provides a model for the behavior of an observed value of the difference D for an individual selected at random from this population. In this situation a probability model explains the variability among the differences in cholesterol level due to treatment with the drug for the individuals in this population. The corresponding population mean difference μ is the average difference (decrease) in cholesterol level that we would observe if all of the individuals in this population were treated with this drug. The standard deviation σ of the probability model serves to quantify the variability among the differences for the individuals in this population.

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In Chapter 9 we considered inferences about the distribution of a dichotomous variable. Since the distribution of a dichotomous variable is completely determined by the corresponding population success proportion, we found that the sampling distribution of the sample proportion \hat{p} was determined by the sampling method. In general, the distribution of a quantitative variable is not completely determined by a single parameter. Therefore, before we can make inferences about the distribution of a quantitative variable we need to make some assumptions about the distribution of the variable.

As we noted in Chapter 7, we can envision a probability model for the distribution of a quantitative variable X in terms of a box model. If X has a finite number of possible values, then a probability model specifies the probabilities with which these possible values will occur. If the balls in a box are labeled with the possible values of X and the proportion of balls with each label (value of X) in the box is equal to the probability for that value specified by the probability model, then, according to the probability model, observing a value of X is equivalent to selecting a single ball at random from this box and observing the label on the ball. For a continuous variable X observing the value of X is like selecting a ball at random from a box containing an infinite collection of suitably labeled balls.

Given a probability model for the distribution of X, a collection of n values of X is said to form a random sample of size n if it satisfies the two properties given below.

1. Each value of the variable X that we observe can be viewed as a single value chosen at random from a (usually infinite) population of values which are distributed according to the specified probability model for the distribution of X.

2. The observed values of X are independent. That is, knowing the value of one or more of the observed values of X has no effect on the probabilities associated with other values of X.

In other words, in terms of the box model for the probability distribution of X a random sample of n values of X can be viewed as a collection of n labels corresponding to a simple random sample selected with replacement from a box of suitably labeled balls.

Given a random sample of values of X it seems obvious that the sample mean \overline{X} is an appropriate estimate of the corresponding population mean μ . The sampling distribution of \overline{X} , which describes the sample to sample variability in \overline{X} , serves as the starting point for our study of the behavior of the sample mean \overline{X} as an estimator of the population mean μ . The exact form of the sampling distribution of \overline{X} depends on the form of the distribution of X. However, the two important properties of the sampling distribution of the sample mean given below are valid regardless of the exact form of the distribution of X.

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Let \overline{X} denote the sample mean of a random sample of size n from a population (distribution) with population mean μ and population standard deviation σ . The sampling distribution of the sample mean \overline{X} has the following characteristics.

1. The mean of the sampling distribution of \overline{X} is the corresponding population mean μ . This indicates that the sample mean \overline{X} is unbiased as an estimator of the population mean μ . Recall that saying that a statistic is unbiased means that, even though the statistic will overestimate the parameter for some samples and will underestimate the parameter in other samples, it will do so in such a way that, in the long run, the values of the statistic will average to give the correct value of the parameter.

2. The population standard error of the sample mean \overline{X} (the standard deviation of the sampling distribution of \overline{X}) is $\operatorname{SE}(\overline{X}) = \sigma/\sqrt{n}$. That is, the standard deviation of the sampling distribution of \overline{X} is equal to the standard deviation of the distribution of X divided by the square root of the sample size. Notice that this implies that the sample mean is less variable than a single observation as an estimator of μ ; and that if μ and σ are held constant, then the variability in \overline{X} as an estimator of μ decreases as the sample size increases reflecting the fact that a larger sample provides more information than a smaller sample.

Since the form of the sampling distribution of \overline{X} depends on the form of the distribution of X, we will need to make some assumptions about the distribution of X before we can proceed with our discussion of inference for the population mean μ . These assumptions correspond to the choice of a probability model (density curve) to represent the distribution of X. There is an infinite collection of probability models to choose from but we will restrict our attention to the normal probability model, which we discussed in Section 8.3. The normal distribution is appropriate for many situations when the distribution of Xis symmetric and mound shaped. However, this does not imply that all, or even most, distributions of continuous variables are normal distributions.

Some of the reasons that we will use the normal distribution as a probability model are:

(1) the theory needed for inference has been worked out for the normal model;

(2) there are many situations where a normal distribution provides a reasonable model for the distribution of a quantitative variable;

(3) even though the inferential methods we discuss are based on the assumption that the

distribution of the variable is exactly a normal distribution, it is known that these inferential methods actually perform reasonably well provided the true distribution of the variable is "reasonably similar to a normal distribution"; and,

(4) it is often possible to transform or redefine a variable so that its distribution is reasonably modeled by a normal distribution.

Strictly speaking, the inferential methods based on the Student's t distribution described in Sections 11.3 and 11.4 are only appropriate when the data constitute a random sample from a normal population. However, these methods are known to be generally reasonable even when the underlying population is not exactly a normal population, provided the underlying population distribution is reasonably symmetric and the true density curve has a more or less normal (bell–shaped) appearance. We cannot be sure that an underlying population (distribution) is normal; however, we can use descriptive methods to look for evidence of possible nonnormality, provided the sample size is reasonably large. The most easily detected and serious evidence of nonnormality you should look for is evidence of extreme skewness or evidence of extremely outlying observations. If there is evidence of extreme skewness or extremely outlying observations, then the inferential methods based on the Student's t distribution should not be used. You will need to consult a statistician or a more advanced statistics book for more information about alternate methods.

Normal probability plots. If you have access to a suitable computer program, you can construct a normal probability plot (QQ-plot) as an aid for assessing whether your data are consistent with what we would expect to see in a random sample from a normal distribution. A normal probability plot provides a graphical comparison of the observed sample values with idealized sample values which we might expect to observe if the sample came from a normal distribution. The idealized sample values used in a normal probability plot are known as normal scores. Ideally, we would expect a random sample of size n from a normal distribution to partition the region under the normal density curve into n + 1 regions of equal area, with each of these areas being 1/(n + 1). The n normal scores, which constitute our idealized random sample, can thus be formed by determining the n values which would partition the area under the normal density curve into n + 1 regions each of area 1/(n + 1) as suggested above. Once these normal scores are obtained we can plot the ordered normal scores versus the ordered observed data values and examine this normal probability plot looking for evidence of systematic disagreement between the actual sample values and the expected normal scores. If the sample really was a sample from a

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normal distribution, then we would expect the normal probability plot to approximate a straight line. Therefore, a normal probability plot which differs greatly in appearance from a straight line provides evidence that the sample may not come from a normal distribution.

11.3 Estimating a normal population mean toc

The sample mean \overline{X} provides a single number estimate of the population mean μ . We can think of \overline{X} as our "best guess" of the value of μ . As noted above we know that \overline{X} is unbiased as an estimator of μ ; therefore, on the average in the long run (under repeated sampling) we know that \overline{X} provides a good estimate of the unknown mean μ . Since this unbiasedness does not guarantee that the observed value of \overline{X} , based on a single sample, will be close to the true, unknown value of μ , it would be useful to have a confidence interval estimate of μ .

Recall that when the sample mean \overline{X} corresponds to a random sample from a population distribution with population mean μ and population standard deviation σ the sampling distribution of \overline{X} has mean μ and standard deviation $\operatorname{SE}(\overline{X}) = \sigma/\sqrt{n}$ (the population standard error of \overline{X}). When the underlying population distribution is normal we can say more about the form of this sampling distribution. If the random sample from which the sample mean \overline{X} is computed is a random sample from a normal population (distribution) with population mean μ and population standard deviation σ , then the sampling distribution of \overline{X} is a normal distribution with population mean μ and population standard deviation $\operatorname{SE}(\overline{X}) = \sigma/\sqrt{n}$. Thus, under these assumptions, the quantity

$$Z = \frac{\overline{X} - \mu}{\operatorname{SE}(\overline{X})} = \frac{\overline{X} - \mu}{\sigma/\sqrt{n}}$$

behaves in accordance with the standard normal distribution.

We know that a standard normal variable Z will take on a value between -1.96 and 1.96 with probability .95 ($P(-1.96 \le Z \le 1.96) = .95$). This implies that

$$P\left(-1.96 \le \frac{\overline{X} - \mu}{\operatorname{SE}(\overline{X})} \le 1.96\right) = .95$$

which is equivalent to

$$P(\overline{X} - 1.96\text{SE}(\overline{X}) \le \mu \le \overline{X} + 1.96\text{SE}(\overline{X})) = .95$$

This probability statement says that 95% of the time when we take a random sample from a normal distribution we will observe a value of \overline{X} such that the population mean μ will be between $\overline{X} - 1.96\text{SE}(\overline{X})$ and $\overline{X} + 1.96\text{SE}(\overline{X})$. Unfortunately, we cannot use this interval as a confidence interval for μ , since the population standard error $\text{SE}(\overline{X}) = \sigma/\sqrt{n}$ depends on the unknown population standard deviation σ and thus is not computable. We can avoid this difficulty by replacing the unknown population standard error

$$SE(\overline{X}) = \frac{\sigma}{\sqrt{n}}$$
 (population standard eror)

by the sample standard error

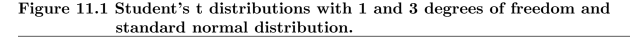
$$\widehat{\operatorname{SE}}(\overline{X}) = \frac{S}{\sqrt{n}}$$
 (sample standard eror),

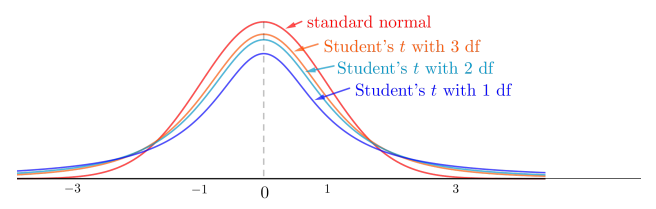
where $S = S_X$ is the sample standard deviation, and basing our confidence interval estimate on the quantity

$$T = \frac{\overline{X} - \mu}{\widehat{\operatorname{SE}}(\overline{X})} = \frac{\overline{X} - \mu}{S/\sqrt{n}}.$$

If the sample mean \overline{X} and the sample standard deviation S are computed from a random sample of size n from a normal population (distribution) with population mean μ and population standard deviation σ , then the quantity T defined above follows the **Student's** t distribution with n-1 degrees of freedom. The Student's t distribution with n-1degrees of freedom is symmetric about zero and has a density curve very similar to that of the standard normal distribution. The main difference between these two distributions is that the Student's t distribution has "heavier" tails than the standard normal distribution. That is, the tails of the Student's t density curve approach the x-axis more slowly than do the tails of the standard normal density curve. As the sample size (and the degrees of freedom) increases the Student's t distribution becomes more similar to the standard normal distribution. In fact, the standard normal distribution is the limiting version of the Student's t distribution in the sense that the Student's t density curve approaches the standard normal density curve when the degrees of freedom increases without bound. The relationship between Student's t distributions and the standard normal distribution is indicated by the plots in Figure 11.1.

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The 97.5 percentiles, k, of the Student's t distributions for selected d.f. are given in Table 11.1. These k values are used to form confidence intervals for μ . Note that you do not need to use the formulas we are about to give to construct confidence intervals by hand. You can use a calculator or computer program to find confidence intervals for μ .

				97.5%			
_				011070			
				0		k T	_
Table	11.1 97.	5 perce	entiles	of the S	tudent's	s t distrik	oution.
d.f.	k	d.f.	k	d.f.	k	d.f.	k
1	12.706	14	2.145	27	2.052	40	2.021
2	4.303	15	2.131	28	2.048	45	2.014
3	3.182	16	2.120	29	2.045	50	2.008
4	2.776	17	2.110	30	2.042	55	2.004
5	2.571	18	2.101	31	2.040	60	2.000
6	2.447	19	2.093	32	2.037	65	1.997
7	2.365	20	2.086	33	2.034	70	1.994
8	2.306	21	2.080	34	2.032	75	1.992
9	2.262	22	2.074	35	2.030	80	1.989
10	2.228	23	2.069	36	2.028	85	1.988
11	2.201	24	2.064	37	2.026	90	1.986
12	2.179	25	2.060	38	2.024	95	1.985
13	2.160	26	2.056	39	2.023	100	1.982

Given a constant k such that $P(-k \le T \le k) = .95$, where T denotes a Students t variable with n-1 degrees of freedom, we have

$$P\left(-k \le \frac{\overline{X} - \mu}{\widehat{\operatorname{SE}}(\overline{X})} \le k\right) = .95$$

which is equivalent to

$$P\left(\overline{X} - k \cdot \widehat{\operatorname{SE}}(\overline{X}) \le \mu \le \overline{X} + k \cdot \widehat{\operatorname{SE}}(\overline{X})\right) = .95.$$

The quantity

$$\operatorname{ME}(\overline{X}) = k \cdot \widehat{\operatorname{SE}}(\overline{X}) = \frac{k \cdot S}{\sqrt{n}}$$

is the 95% margin of error of \overline{X} . The preceding probability statement says that 95% of the time when we take a random sample from a normal distribution we will observe values of \overline{X} and S such that the population mean μ will be between $\overline{X} - \operatorname{ME}(\overline{X})$ and $\overline{X} + \operatorname{ME}(\overline{X})$. Thus, the interval from $\overline{X} - \operatorname{ME}(\overline{X})$ to $\overline{X} + \operatorname{ME}(\overline{X})$ is a 95% confidence interval estimate for μ . To compute this confidence interval we need to determine the value of the appropriate margin of error multiplier k. This multiplier depends on the size of the sample so that there is a different multiplier for each sample size. The symmetry of the Student's t distribution and the definition of k imply that k is the 97.5 percentile of the Student's t distribution with n - 1 degrees of freedom. The 95% margin of error multipliers (k's) based on the Student's t distribution, for several choices of the degrees of freedom (d.f.), are given in Table 11.1.

This confidence interval estimate may be reported using a statement such as: We are 95% confident that the population mean μ is between $\overline{X} - \operatorname{ME}(\overline{X})$ and $\overline{X} + \operatorname{ME}(\overline{X})$. Notice that it is the sample mean \overline{X} and the margin of error $\operatorname{ME}(\overline{X})$ that vary from sample to sample. The population mean μ is a fixed, unknown parameter that does not vary. Therefore, the 95% confidence level applies to the method used to generate the confidence interval estimate. That is, the method (obtain a simple random sample and compute the numbers $\overline{X} - \operatorname{ME}(\overline{X})$ and $\overline{X} + \operatorname{ME}(\overline{X})$ forming the limits of the confidence interval) is such that 95% of the time (when we take a random sample and compute these values) it will yield a pair of confidence interval limits which bracket the population mean μ .

Example 11.1 NHANES (Example 8.1 revisited) In Example 8.1 we showed that the 2013–2014 NHANES height data for women (n = 2888) and the 2013–2014 NHANES height

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data for men (n = 2642) could be reasonably modeled as random samples from normal distributions. We will now find confidence intervals for the corresponding population mean heights.

Heights of women First consider the random sample of n = 2888 heights of adult women. Let W denote the height of an adult woman. We will model the n = 2888 heights as a random sample from a normal distribution with population mean μ_W and population standard deviation σ_W . In this application population mean height μ_W is the unknown mean height of all adult women in the United States who could have been included in the 2013–2014 NHANES sample. The unknown population standard deviation σ_W is a measure of the variability among the heights of the women in this population.

The height data is summarized graphically in Figure 11.2a. Two density curves are superimposed on the histogram in this figure. There is a normal density curve, with μ and σ chosen to match the data, and there is a "kernel" density curve which is basically a smooth version of the histogram. As we noted in Chapter 8, it is somewhat tricky to properly compare distributions based on histograms or density curves. Therefore, a normal probability plot or QQ-plot (Quantile – Quantile plot) is provided. The QQ-plot is used to assess whether the data can reasonably be modeled as a random sample from a normal distribution. (Recall that we want to see a reasonably strong linear trend in the QQ-plot.) In this application the QQ-plot shows a strong linear trend and we can conclude that it is reasonable to model the data as a random sample from a normal distribution.

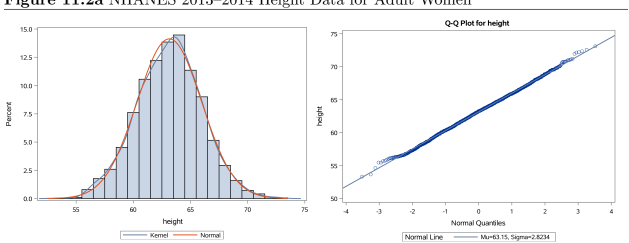


Figure 11.2a NHANES 2013–2014 Height Data for Adult Women

Figure 11.2b 95% Confidence Interval for Mean Height of Adult Women

The TTEST Procedure

Variable: height

N	Mean	Std Dev	Std Err	Minimum	Maximum
2888	63.1497	2.8234	0.0525	53.3071	73.1102

Mean	95% CL Mean		
63.1497	63.0467	63.2528	

The SAS output given in Figure 11.2b provides summary information and the 95% confidence interval for the population mean height of adult women in the USA at the time of the NHANES 2013–2014 survey.

The summary statistics provided indicated that the sample size is n = 2888, the sample mean height is $\overline{W} = 63.1497$ and the sample standard deviation is $S_W = 2.8234$. The sample standard error of \overline{W} is also given. Here $\widehat{SE}(\overline{W}) = .0525$, recall that this is $S_W/\sqrt{n} = 2.8234/\sqrt{2888}$.

The confidence limits (CL Mean in the output) reported are 63.0467 and 63.2528. (If you want to perform the computation of this confidence interval by hand, the 97.5 percentile of the Student's t distribution with n-1 = 2887 df is 1.9608.) Therefore, we can conclude that we are 95% confident that the mean height of all adult women in the USA at the time of the NHANES 2013–2014 survey is between 63.0467 and 63.2528 inches.

Be careful to distinguish between the sample mean height \overline{W} which is known and the population mean height μ_W which is unknown. In this example we know that the mean height of the n = 2888 women who were measured is 63.1497 inches. If we were able to measure the heights of all of the women in the population our sample came from, we are 95% confident that the mean height we would obtain would be between 63.0467 and 63.2528 inches.

Aside (technicality – not to worry about here): Here we are modeling the n = 2888 observed heights as a simple random sample from a normal distribution. These data were actually obtained using a more complex method of random sampling and adjustments could be made to the methods we employed to compute the confidence interval to account for this.

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Heights of men Next consider the random sample of n = 2642 heights of adult men. Let M denote the height of an adult man. We will model the n = 2642 heights as a random sample from a normal distribution with population mean μ_M and population standard deviation σ_M . In this application population mean height μ_M is the unknown mean height of all adult men in the United States who could have been included in the 2013–2014 NHANES sample. The unknown population standard deviation σ_M is a measure of the variability among the heights of the men in this population.

The height data is summarized graphically in Figure 11.3a. Two density curves are superimposed on the histogram in this figure. There is a normal density curve, with μ and σ chosen to match the data, and there is a "kernel" density curve which is basically a smooth version of the histogram. As we noted in Chapter 8, it is somewhat tricky to properly compare distributions based on histograms or density curves. Therefore, a normal probability plot or QQ-plot (Quantile – Quantile plot) is provided. The QQ-plot is used to assess whether the data can reasonably be modeled as a random sample from a normal distribution. (Recall that we want to see a reasonably strong linear trend in the QQ-plot.) In this application the QQ-plot shows a strong linear trend and we can conclude that it is reasonable to model the data as a random sample from a normal distribution.

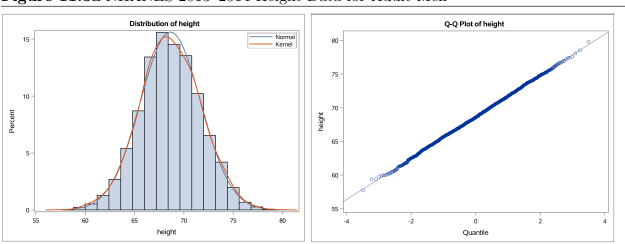


Figure 11.3a NHANES 2013–2014 Height Data for Adult Men

Figure 11.3b 95% Confidence Interval for Mean Height of Adult Men

The TTEST Procedure

Variable: height

N	Mean	Std Dev	Std Err	Minimum	Maximum
2642	68.6349	3.0649	0.0596	57.7165	79.7638

Mean	95% C	L Mean
68.6349	68.5180	68.7518

The SAS output given in Figure 11.3b provides summary information and the 95% confidence interval for the population mean height of adult men in the USA at the time of the NHANES 2013–2014 survey.

The summary statistics provided indicated that the sample size is n = 2642, the sample mean height is $\overline{M} = 68.6349$, and the sample standard deviation is $S_M = 3.0649$. The sample standard error of \overline{M} is also given. Here $\widehat{SE}(\overline{M}) = .0596$, recall that this is $S_M/\sqrt{n} = 3.0649/\sqrt{2642}$.

The confidence limits (CL Mean in the output) reported are 68.5180 and 68.7518. (If you want to perform the computation of this confidence interval by hand, the 97.5 percentile of the Student's t distribution with n-1 = 2642 df is 1.9609.) Therefore, we can conclude that we are 95% confident that the mean height of all adult men in the USA at the time of the NHANES 2013–2014 survey is between 68.5180 and 68.7518 inches.

Be careful to distinguish between the sample mean height \overline{M} which is known and the population mean height μ_M which is unknown. In this example we know that the mean height of the n = 2642 men who were measured is 68.6349 inches. If we were able to measure the heights of all of the men in the population our sample came from, we are 95% confident that the mean height we would obtain would be between 68.5180 and 68.7518 inches.

Aside (technicality – not to worry about here): Here we are modeling the n = 2642 observed heights as a simple random sample from a normal distribution. These data were actually obtained using a more complex method of random sampling and adjustments could be made to the methods we employed to compute the confidence interval to account for this.

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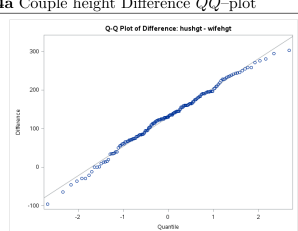
Example 11.2. Heights of husbands and wives. The data used in this example are part of data set 231 in Hand, Daly, Lunn, McConway, and Ostrowski (1994), A Handbook of Small Data Sets, Chapman and Hall, London. The original source is Marsh (1988), Exploring Data, Cambridge, UK. A random sample of n = 169 married couples was selected from the 1980 OPCS study of the heights and weights of the adult population of Great Britain. The data consist of paired heights (in mm) for these husbands and wives. A few of these paired heights are given in Table 11.2 and all of the differences (husband's height minus wife's height) are given in Table 11.3.

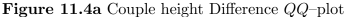
couple	husband's height	wife's height	difference	
1	1786	1590	196	
2	1754	1660	94	
3	1755	1590	165	
4	1725	1550	175	
5	1796	1550	246	
		•••		
169	1641	1570	71	

Table 11.2. Husband and wife height data (partial).

Table 11.3. Husband and wife height differences.

$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	-										
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$		-96	-65	-46	-37	-30	-30	-21	-12	0	0
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$		1	9	13	14	15	19	34	35	35	36
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$		39	40	50	55	56	59	60	60	65	65
9094949596100103103105110110113113114115118119120120123123123123125125125125125125128128128129130130130133134135135135135139140141141144144145145145145150151155155155159159160160160160161161164165165166169170174175175178180181183183188189190190191193194195195195196196196204205209210215219225228228229233235236239241243244246		66	68	70	70	71	71	73	74	75	75
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$		79	79	81	83	84	84	84	85	85	88
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$		90	94	94	95	96	100	103	103	105	110
128129130130133134135135135135139140141141144144145145145145150151155155155159159160160160160161161164165165165166169170174175175178180181183183188189190190191193194195195195196196196204205209210215219225228228229233235236239241243244246		110	113	113	114	115	118	119	120	120	123
135139140141141144144145145145145150151155155155159159160160160160161161164165165165166169170174175175178180181183183188189190190191193194195195195196196196204205209210215219225228228229233235236239241243244246		123	123	123	125	125	125	125	125	128	128
145150151155155159159160160160160161161164165165165166169170174175175178180181183183188189190190191193194195195195196196196204205209210215219225228228229233235236239241243244246		128	129	130	130	130	133	134	135	135	135
160160161161164165165165166169170174175175178180181183183188189190190191193194195195195196196196204205209210215219225228228229233235236239241243244246		135	139	140	141	141	144	144	145	145	145
170174175175178180181183183188189190190191193194195195195196196196204205209210215219225228228229233235236239241243244246		145	150	151	155	155	155	159	159	160	160
189190190191193194195195195196196196204205209210215219225228228229233235236239241243244246		160	160	161	161	164	165	165	165	166	169
196196204205209210215219225228228229233235236239241243244246		170	174	175	175	178	180	181	183	183	188
228 229 233 235 236 239 241 243 244 246		189	190	190	191	193	194	195	195	195	196
		196	196	204	205	209	210	215	219	225	228
250 255 258 259 271 276 281 295 303		228	229	233	235	236	239	241	243	244	246
	_	250	255	258	259	271	276	281	295	303	





Since the normal probability plot (QQ-plot) of Figure 11.4a is reasonably linear we can model these 169 differences as a random sample from a normal distribution of differences.

The population mean difference μ_D is the average difference in height corresponding to the population of all married couples in Great Britain in 1980. (Technically, we should restrict this mean to those married couples included in the 1980 COPS study from which the sample was taken.)

The sample mean for these n = 169 differences is 129.82 mm, the sample standard deviation is 76.02 mm, and the sample standard error of the sample mean is $76.0211/\sqrt{169} = 5.85$.

We are 95% confident that for the population of all married couples in Great Britain in 1980 (as represented by the COPS study group from which this sample came), on average, the husband's height exceeds his wife's height by something between 118.3 mm and 141.4 mm.

Figure 11.4b Couple height Difference SAS Output

The TTEST Procedure

Difference: husbandheight - wifeheight

Ν	Mean	Std Dev	Std Err	Minimum	Maximum
169	129.8	76.0211	5.8478	-96.0000	303.0

Mean		lean			
129.8	118.3	141.4			

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You should get the confidence interval from the output – this computation is included to illustrate the computations. The multiplier for the 95% margin of error, based on the Student's t distribution with n - 1 = 168 degrees of freedom, is k = 1.9742. Thus the 95% margin of error of the sample mean is (1.9742)(5.8478) = 11.5447 and the limits for the 95% confidence interval estimate of μ_D are 129.8225 - 11.5447 = 118.2778 and 129.8225 + 11.5447 = 141.3672.

11.4 Testing a hypothesis about a normal population mean toc

The hypothesis testing procedures discussed in this section are based on the fact that, when $\mu = \mu_0$, the **Student's** t **test statistic**

$$T_{calc} = \frac{\overline{X} - \mu_0}{\widehat{\operatorname{SE}}(\overline{X})} = \frac{\overline{X} - \mu_0}{S/\sqrt{n}},$$

follows the Student's t distribution with n-1 degrees of freedom. Recall that, technically, this result requires that the sample from which the sample mean \overline{X} and the sample standard deviation S are computed forms a random sample from a normal distribution with population mean μ . However, these methods are known to be generally reasonable even when the underlying population is not exactly a normal population, provided the underlying population distribution is reasonably symmetric and the true density curve has a more or less normal (bell-shaped) appearance.

Recall that a hypothesis (statistical hypothesis) is a conjecture about the nature of the population. When we considered hypotheses about dichotomous populations we noted that the population was completely determined by the population success proportion p. In the present context of sampling from a normal population two parameters, the mean and the standard deviation, must be specified to completely determine the normal population. A hypothesis about the value of the population mean μ of a normal distribution specifies where the center of this normal distribution, μ , is located on the number line but places no restriction on the population standard deviation.

Even though the logic behind a hypothesis test for a population mean μ is the same as the logic behind a hypothesis test for a population proportion p, we will introduce hypothesis testing for a mean in the context of a simple hypothetical example.

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Example 11.3. Strength of bricks. Consider a brick manufacturer that has produced a large batch of bricks and wants to determine whether these bricks are suitable for a particular construction project. The specifications for this project require bricks with a mean compressive strength that exceeds 3200 psi. In order to assess the suitability of this batch of bricks the manufacturer will obtain a simple random sample of bricks from this batch and measure their compressive strength. In this example a single brick is a unit, the entire large batch of bricks is the population, and a suitable variable X is the compressive strength of an individual brick (in psi). We will assume that the distribution of X is reasonably modeled by a normal distribution with population mean μ and population standard deviation σ . In this example the population mean μ represents the mean compressive strength for all of the bricks in this large batch and the population standard deviation quantifies the variability from brick to brick in (measured) compressive strength.

The manufacturer does not want to use these bricks for this construction project unless there is sufficient evidence to claim that the population mean compressive strength for this batch exceeds 3200 psi. Thus, the question of interest here is: "Is there sufficient evidence to justify using this batch of bricks for this project?" In terms of the population mean μ the research hypothesis is $H_1: \mu > 3200$ (the mean compressive strength for this batch of bricks exceeds 3200 psi); and the null hypothesis is $H_0: \mu \leq 3200$ (the mean compressive strength for this batch of bricks does not exceed 3200 psi). In other words, the manufacturer will tentatively assume that these bricks are not suitable for this project and will check to see if there is sufficient evidence against this tentative assumption to justify the conclusion that these bricks are suitable for the project.

A test of the null hypothesis $H_0: \mu \leq 3200$ versus the research hypothesis $H_1: \mu > 3200$ begins by tentatively assuming that the mean compressive strength for this batch of bricks is no larger than 3200 psi. Under this tentative assumption it would be surprising to observe a sample mean compressive strength \overline{X} that was much larger than 3200. Thus the test should reject $H_0: \mu \leq 3200$ in favor of $H_1: \mu > 3200$ if the observed value of \overline{X} is sufficiently large relative to 3200. In order to determine whether \overline{X} is large relative to 3200 we need an estimate of the sample to sample variability in \overline{X} . The sample standard error of the sample mean $\widehat{SE}(\overline{X}) = S/\sqrt{n}$, where S denotes the sample standard deviation,

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provides a suitable measure of the sample to sample variability in \overline{X} . Our conclusion will hinge on deciding whether the observed value of the Student's t test statistic

$$T_{calc} = \frac{\overline{X} - 3200}{\widehat{\operatorname{SE}}(\overline{X})} = \frac{\overline{X} - 3200}{S/\sqrt{n}}$$

is far enough above zero to make $\mu > 3200$ more tenable than $\mu \leq 3200$. We will base this decision on the probability of observing a value of T as large or larger than the actual calculated value T_{calc} of T, under the assumption that $\mu \leq 3200$. This probability (computed assuming that $\mu = 3200$) is the P-value of the test. We will use the fact that, when $\mu = 3200$, the Student's t statistic T follows the Student's t distribution with n - 1degrees of freedom to calculate the P-value.

First suppose that a simple random sample of n = 100 bricks is selected from the large batch. Further suppose that the sample mean compressive strength of these 100 bricks is $\overline{X} = 3481$ psi and the sample standard deviation is S = 1118.38. In this case we know that the mean compressive strength of the bricks in the sample $\overline{X} = 3481$ exceeds 3200 and we need to decide whether this suggests that the mean compressive strength of all of the bricks in the batch μ exceeds 3200. In this case the sample standard error of \overline{X} is $\widehat{SE}(\overline{X}) = 1118.38/\sqrt{100} = 111.838$. Thus \overline{X} exceeds 3200 by 281 psi which is 281/111.838 = 2.5126 standard error units, *i.e.*, $T_{calc} = 2.5126$. In this case the *P*-value .0068 is the probability of observing a calculated T value as large or larger than 2.5126when $\mu \leq 3200$. That is, if $\mu \leq 3200$ and we perform this experiment with n = 100, then we would expect to see a value of \overline{X} that is 2.5126 standard error units above the hypothesized value 3200 (2.5126 times $\widehat{SE}(\overline{X})$ psi above 3200) about 0.68% of the time. Therefore, observing values of \overline{X} and S in a sample of size n = 100 which yield a value of T_{calc} as large or larger than 2.5126 would be very surprising if the true mean compressive strength for the batch μ was not larger than 3200 and we have sufficient evidence to reject the null hypothesis $H_0: \mu \leq 3200$ in favor of the research hypothesis $H_1: \mu > 3200$. In this case we would conclude that there is sufficient evidence to contend that the mean compressive strength for this batch of bricks exceeds 3200 psi, *i.e.*, these bricks are suitable for the construction project.

Now suppose that a simple random sample of n = 100 bricks is selected from the large batch and the sample mean compressive strength of these 100 bricks is $\overline{X} = 3481$ psi as before. However, suppose that in this case the sample standard deviation is S = 2329.3. As

before we know that the mean compressive strength of the bricks in the sample $\overline{X} = 3481$ exceeds 3200 and we need to decide whether this suggests that the mean compressive strength of all of the bricks in the batch μ exceeds 3200. In this case the sample standard error of \overline{X} is $\widehat{SE}(\overline{X}) = 2329.3/\sqrt{100} = 232.93$. Thus \overline{X} exceeds 3200 by 281 psi which is 281/232.93 = 1.2064 standard error units, *i.e.*, $T_{calc} = 1.2064$. In this case the *P*-value .1153 is the probability of observing a calculated T value as large or larger than 1.2064when $\mu \leq 3200$. That is, if $\mu \leq 3200$ and we perform this experiment with n = 100, then we would expect to see a value of \overline{X} that is 1.2064 standard error units above the hypothesized value 3200 (1.2064 times $\widehat{SE}(\overline{X})$ psi above 3200) about 11.53% of the time. Therefore, observing values of \overline{X} and S in a sample of size n = 100 which yield a value of T_{calc} as large or larger than 1.2064 would not be very surprising if the true mean compressive strength for the batch μ was not larger than 3200 and we do not have sufficient evidence to reject the null hypothesis $H_0: \mu \leq 3200$ in favor of the research hypothesis $H_1: \mu > 3200$. In the case we would conclude that there is not sufficient evidence to contend that the mean compressive strength for this batch of bricks exceeds 3200 psi, *i.e.*, these bricks are not suitable for the construction project.

The research hypothesis in the brick example is a directional hypothesis of the form H_1 : $\mu > \mu_0$, where $\mu_0 = 3200$. We will now discuss the details of a hypothesis test for a directional research hypothesis of this form. For the test procedure to be valid the specified value μ_0 and the direction of the research hypothesis must be motivated from subject matter knowledge before looking at the data that are to be used to perform the test.

Let μ_0 denote the hypothesized value which we wish to compare with μ . The research hypothesis states that μ is greater than μ_0 ; in symbols we will indicate this research hypothesis by writing

$$H_1: \mu > \mu_0.$$

The null hypothesis is the negation of $H_1: \mu > \mu_0$ which states that μ is not greater than μ_0 ; in symbols we will indicate this null hypothesis by writing

$$H_0: \mu \le \mu_0.$$

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The research hypothesis $H_1: \mu > \mu_0$ specifies that the normal distribution is one of the normal distributions for which the population mean μ is greater than μ_0 . The null hypothesis $H_0: \mu \leq \mu_0$ specifies that the normal distribution is one of the normal distributions for which the population mean μ is at most μ_0 . The population standard deviation σ is not restricted by either hypothesis. Assuming that the population standard deviation is the same regardless of which hypothesis is true, this competing pair of hypotheses provides a decomposition of all possible normal distributions with this σ into the collection of normal distributions where $\mu > \mu_0$ and the research hypothesis is true and the collection of normal distributions where $\mu \leq \mu_0$ and the null hypothesis is true. Our goal is to use the data to decide which of these two collections of normal distributions contains the normal distribution we are actually sampling from.

Since a hypothesis test begins by tentatively assuming that the null hypothesis is true, we need to decide what constitutes evidence against the null hypothesis $H_0: \mu \leq \mu_0$ and in favor of the research hypothesis $H_1: \mu > \mu_0$. We will assume that the unknown population standard deviation σ is fixed regardless of the value of μ . The difference $\overline{X} - \mu_0$ between the sample mean \overline{X} and the hypothesized value μ_0 , expressed in standard error units, will be used to assess the strength of the evidence in favor of the research hypothesis. Generally, we would expect to observe larger values of \overline{X} more often when the research hypothesis $H_1: \mu > \mu_0$ is true than when the null hypothesis $H_0: \mu \leq \mu_0$ is true. In particular, we can view the observation of a value of \overline{X} that is sufficiently large relative to μ_0 as constituting evidence against the null hypothesis $H_0: \mu \leq \mu_0$ and in favor of the research hypothesis $H_1: \mu > \mu_0$. To decide whether the observed value of \overline{X} is "sufficiently large relative to μ_0 " we need to take the variability in the data into account. We can do this by basing our decision on the corresponding Student's t test statistic,

$$T_{calc} = \frac{\overline{X} - \mu_0}{\widehat{SE}(\overline{X})} = \frac{\overline{X} - \mu_0}{S/\sqrt{n}},$$

instead of \overline{X} alone. Notice that values of T_{calc} which are large relative to zero correspond to values of \overline{X} which are large relative to μ_0 . Deciding whether the observed value of T_{calc} is sufficiently large relative to zero to allow rejection of H_0 is based on the corresponding P-value, which is defined below.

The *P*-value for testing the null hypothesis $H_0: \mu \leq \mu_0$ versus the research hypothesis $H_1: \mu > \mu_0$ is the probability of observing a value of a Student's *t* variable *T* as large or

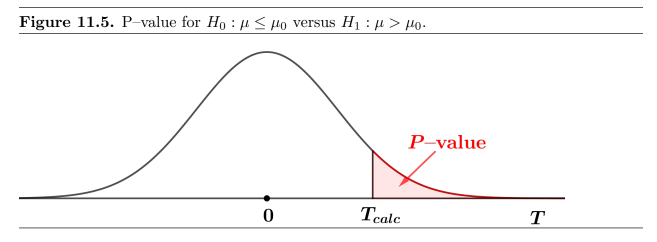
larger than the calculated value T_{calc} that we actually do observe, *i.e.*, P-value = $P(T \ge T_{calc})$, where T denotes a Student's t variable with n-1 degrees of freedom. This P-value is computed under the assumption that the research hypothesis $H_1 : \mu > \mu_0$ is false and the null hypothesis $H_0 : \mu \le \mu_0$ is true. Because the null hypothesis only specifies that $\mu \le \mu_0$, we need to choose a particular value of μ (that is no larger than μ_0) in order to compute the P-value. It is most appropriate to use $\mu = \mu_0$ for this computation. Using $\mu = \mu_0$, which defines the boundary between $\mu \le \mu_0$, where the null hypothesis is true, and $\mu > \mu_0$, where the research hypothesis is true, provides some protection against incorrectly rejecting $H_0 : \mu \le \mu_0$.

The steps for performing a hypothesis test for

$$H_0: \mu \leq \mu_0$$
 versus $H_1: \mu > \mu_0$

are summarized below.

1. Use a suitable calculator or computer program to find the P-value = $P(T \ge T_{calc})$, where T denotes a Student's t variable with n - 1 degrees of freedom and $T_{calc} = (\overline{X} - \mu_0)/\widehat{SE}(\overline{X})$ as described above. This P-value is the area to the right of T_{calc} under the density curve for the Student's t distribution with n - 1 degrees of freedom as indicated in Figure 11.5.



2a. If the *P*-value is small enough (less than .05 for a test at the 5% level of significance), conclude that the data favor $H_1: \mu > \mu_0$ over $H_0: \mu \leq \mu_0$. That is, if the *P*-value is small enough, then there is sufficient evidence to conclude that the population mean μ is greater than μ_0 .

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2b. If the *P*-value is not small enough (is not less than .05 for a test at the 5% level of significance), conclude that the data do not favor $H_1: \mu > \mu_0$ over $H_0: \mu \le \mu_0$. That is, if the *P*-value is not small enough, then there is not sufficient evidence to conclude that the population mean μ is greater than μ_0 .

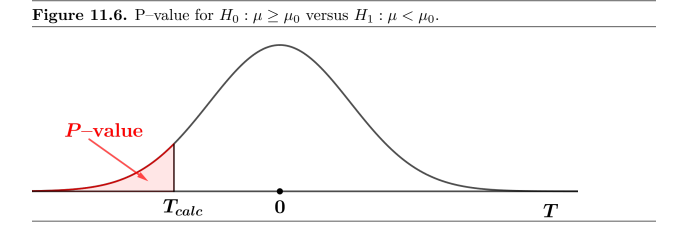
The procedure for testing the null hypothesis $H_0: \mu \leq \mu_0$ versus the research hypothesis $H_1: \mu > \mu_0$ given above is readily modified for testing the null hypothesis $H_0: \mu \geq \mu_0$ versus the research hypothesis $H_1: \mu < \mu_0$. The essential modification is to change the direction of the inequality in the definition of the P-value. Consider a situation where the research hypothesis specifies that the population mean μ is less than the particular, hypothesized value μ_0 . For these hypotheses values of the sample mean \overline{X} that are sufficiently small relative to μ_0 provide evidence in favor of the research hypothesis $H_1: \mu < \mu_0$ and against the null hypothesis $H_0: \mu \geq \mu_0$. Therefore, the appropriate P-value is the probability of observing a value of a Student's t variable T as small or smaller than the value actually observed. As before, the P-value is computed under the assumption that $\mu = \mu_0$. The calculated t statistic T_{calc} is defined as before; however, in this situation the P-value is the area under the density curve of the Student's t distribution with n-1 degrees of freedom to the left of T_{calc} , since values of \overline{X} that are small relative to μ_0 constitute evidence in favor of the research hypothesis.

The steps for performing a hypothesis test for

$$H_0: \mu \ge \mu_0$$
 versus $H_1: \mu < \mu_0$

are summarized below.

1. Use a suitable calculator or computer program to find the P-value = $P(T \leq T_{calc})$, where T denotes a Student's t variable with n - 1 degrees of freedom and $T_{calc} = (\overline{X} - \mu_0)/\widehat{SE}(\overline{X})$ as described above. This P-value is the area to the left of T_{calc} under the density curve for the Student's t distribution with n - 1 degrees of freedom as indicated in Figure 11.6.



2a. If the *P*-value is small enough (less than .05 for a test at the 5% level of significance), conclude that the data favor $H_1 : \mu < \mu_0$ over $H_0 : \mu \ge \mu_0$. That is, if the *P*-value is small enough, then there is sufficient evidence to conclude that the population mean μ is less than μ_0 .

2b. If the *P*-value is not small enough (is not less than .05 for a test at the 5% level of significance), conclude that the data do not favor $H_1: \mu < \mu_0$ over $H_0: \mu \ge \mu_0$. That is, if the *P*-value is not small enough, then there is not sufficient evidence to conclude that the population mean μ is less than μ_0 .

Example 11.4. Alzheimer's disease and homophone spelling confusion A study was conducted to investigate the relationship between Alzheimer's disease and homophone spelling confusion. A homophone pair is a pair of words with the same pronunciation having different meanings and spellings. Twenty patients with Alzheimer's disease were asked to spell 24 homophone pairs (given in random order) and the number of homophone confusions, *e.g.* spelling doe given the context bake bread dough, was recorded for each patient. One year later, the same patients were again asked to spell the same 24 homophone pairs and the number of homophone confusions was again recorded. The data given below are the numbers of homophone confusions at the two times of measurement for the 20 Alzheimer's patients. This example comes from Neils, J., D.P. Roeltgen, and F. Constantinidou, "Decline in homophone spelling associated with loss of semantic influence on spelling in Alzheimer's disease", *Brain and Language*, **49**, (1995).

Patient	Time 1	Time 2	Patient	Time 1	Time 2
1	5	5	11	7	10
2	1	3	12	0	3
3	0	0	13	3	9
4	1	1	14	5	8
5	0	1	15	7	12
6	2	1	16	10	16
7	5	6	17	5	5
8	1	2	18	6	3
9	0	9	19	9	6
10	5	8	20	11	8

Table 11.4. Alzheimer's homophone confusion data.

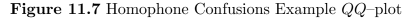
For this discussion we will assume that the 20 patients used in this study were randomly selected from a well–defined population of Alzheimer's patients (let's call it "the population of Alzheimer's patients this sample was selected from".) This assumption makes this problem analogous to the couple height difference example.

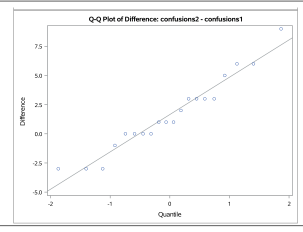
As noted above, each patient completed the homophone confusion test twice yielding a pair of scores (A_i, B_i) . Here A_i is the number of homophone confusions (homophone spelling errors) made by patient *i* on the initial test and B_i is the number of homophone confusions (homophone spelling errors) made by the same patient one year later on the second test. We will use the difference $D_i = B_i - A_i$ to assess the effect of the progression of the disease on homophone spelling ability. Thus our response variable, D, is the difference (increase from time one to time two) in the number of homophone spelling errors made by the patient.

The population mean difference is μ_D the mean difference (from time1 to time2) in the number of homophone confusion errors on this twenty-four homophone pair identification test that we would observe if all of the patients in the population of Alzheimer's patients this sample was selected from were examined using the same protocol as was used in this study.

The obvious conjecture here would be that the progression of the disease will decrease the patient's ability to select the appropriate word. Therefore, we would anticipate that patients will tend to make more errors (more homophone confusions) the second time they are tested. Therefore, we can formalize our research hypothesis as $H_1: \mu_D > 0$. That is, if all the patients in the population of Alzheimer's patients this sample was selected from were examined using the same protocol as was used in this study then, after a year, on average, we would see an increase in the number of homophone confusion errors on the twenty-four homophone pair identification test used in this study.

The normal probability plot (QQ-plot) for these 20 differences provided in Figure 11.7 is reasonably linear so we can model these 20 differences as a random sample from a normal distribution of differences (increases in homophone confusions).







Homophone confusions example

The TTEST Procedure

Difference: confusions2 - confusions1

Ν	Mean	Std De	v	Std E	r	Minim	um	Maximum
	mean	0.0 00			·			
20	1.6500	3.199	9	0.715	5	-3.00	000	9.0000
			_				7	
		Mear	n	95%	CL	Mean		
		1.650	0	0.1524	-	3.1476]	
		DF	t	Value	Р	∙r> t		
		10	-	2.24		0005		
		19		2.31	0	0.0325		

The sample mean for these n = 20 differences is $\overline{D} = 1.65$, the sample standard deviation is $S_D = 3.1999$, and the sample standard error of the sample mean is $\widehat{SE}(\overline{D}) = 3.1999/\sqrt{20} = .7155$.

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The observed value of the Student's t test statistic is $T_{calc} = 2.31$ and the *P*-value for $H_1: \mu_D > 0$ is $P(T \ge 2.31) = \frac{.0325}{2} = .01625$ (The *P*-value given in the SAS output, which is labeled $\Pr > |t|$, is the *P*-value for the nondirectional research hypothesis $H_1: \mu_D \neq 0$. This is called the two-sided or two-tailed *P*-value.)

Since the P-value .01625 is very small we have strong evidence that if we performed this experiment using all of the Alzheimer's patients in the population this sample was selected from, then, on average, the patients would make more mistakes (homophone confusions) the second time they took the test.

In addition to this conclusion it would be helpful to have an estimate of how high this mean increase in homophone confusions would be. Using the confidence interval values in the SAS output, we are 95% confident that the mean difference (from time1 to time2) in the number of homophone confusion errors on this twenty-four homophone pair identification test that we would observe if all of the patients in the population of Alzheimer's patients this sample was selected from were examined using the same protocol as was used in this study is between .1524 and 3.1476. That is, for this population of Alzheimer's patients, on average, a patient would make between .1524 and 3.1476 more mistakes the second time that they took the test. An average increase of .1524 errors is clearly not very substantial; however, an increase of 3.1476 errors (remember that there are 24 homophone pairs on the test) is a substantial increase. Therefore, based on the data we have for these 20 patients it is hard to say whether the progression of the disease in a one year time frame is very serious or not, since even though the confidence interval allows for a μ_D value that is very close to zero.

The hypothesis tests we have discuss thus far are only appropriate when we have enough a priori information, *i.e.*, information that does not depend on the data to be used for the hypothesis test, to postulate that the population mean μ is on one side of a particular value μ_0 . That is, we have only considered situations where the research hypothesis is directional. There are situations when we will not have enough a priori information to allow us to choose the appropriate directional research hypothesis. Instead, we might only conjecture that the population mean μ is different from some particular value μ_0 . In a situation like this our research hypothesis specifies that the population mean μ is different from μ_0 , *i.e.*, $H_1: \mu \neq \mu_0$. To decide between the null hypothesis $H_0: \mu = \mu_0$ and the research hypothesis $H_1: \mu \neq \mu_0$, we need to decide whether the sample mean \overline{X} supports the null hypothesis by being "close to μ_0 ", or supports the research hypothesis by being "far away from μ_0 ". In this situation the *P*-value is the probability that the sample mean \overline{X} would be as far or farther away from μ_0 in either direction as is the value that we actually observe. In other words, the *P*-value is the probability that the standardized distance from \overline{X} to μ_0 (the standardized absolute value of the difference between \overline{X} and μ_0) is as large or larger than the actual observed value of this standardized distance. As before, the *P*-value is computed under the assumption that the null hypothesis is true and $\mu = \mu_0$. In this situation the calculated *t* statistic T_{calc} is the absolute value of the *t* statistic that would be used for testing a directional hypothesis. That is, the calculated *t* statistic is

$$T_{calc} = \left| \frac{\overline{X} - \mu_0}{\widehat{\operatorname{SE}}(\overline{X})} \right|$$

In terms of this t statistic the P-value is the probability that the absolute value of a Students's t variable with n - 1 degrees of freedom T would take on a value as large or larger that T_{calc} , assuming that $\mu = \mu_0$. This probability is the sum of the area under the appropriate Student's t density curve to the left of $-T_{calc}$ and the area under this Student's t density curve to the right of T_{calc} . We need to add these two areas (probabilities) since we are finding the probability that the sample mean \overline{X} would be as far or farther away from μ_0 in either direction as is the value that we actually observe, when $\mu = \mu_0$.

The steps for performing a hypothesis test for

$$H_0: \mu = \mu_0$$
 versus $H_1: \mu \neq \mu_0$

are summarized below.

1. Use a suitable calculator or computer program to find the P-value = $P(|T| \ge T_{calc}) = P(T \le -T_{calc}) + P(T \ge T_{calc})$, where T denotes a Student's t variable with n - 1 degrees of freedom and $T_{calc} = |\overline{X} - \mu_0| / \widehat{SE}(\overline{X})$ as described above. This P-value is the sum of the area to the left of $-T_{calc}$ and the area to the right of T_{calc} , where each area is that under the density curve for the Student's t distribution with n - 1 degrees of freedom over the appropriate region on the x-axis as indicated in Figure 11.9.

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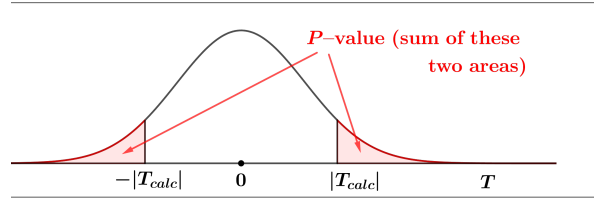


Figure 11.9. P-value for $H_0: \mu = \mu_0$ versus $H_1: \mu \neq \mu_0$.

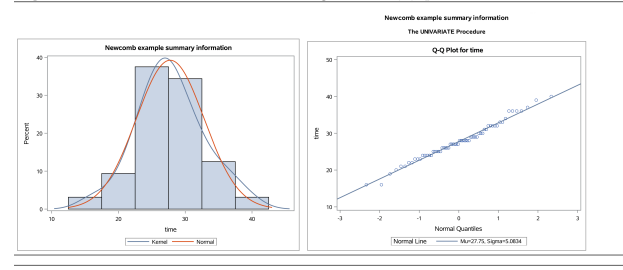
2a. If the *P*-value is small enough (less than .05 for a test at the 5% level of significance), conclude that the data favor $H_1 : \mu \neq \mu_0$ over $H_0 : \mu = \mu_0$. That is, if the *P*-value is small enough, then there is sufficient evidence to conclude that the population mean μ is not equal to μ_0 .

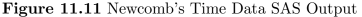
2b. If the *P*-value is not small enough (is not less than .05 for a test at the 5% level of significance), conclude that the data do not favor $H_1: \mu \neq \mu_0$ over $H_0: \mu = \mu_0$. That is, if the *P*-value is not small enough, then there is not sufficient evidence to conclude that the population mean μ differs from μ_0 .

Example 11.5. Newcomb's measurements of the speed of light. In 1882 Simon Newcomb conducted an investigation to measure the speed of light. The essence of Newcomb's experiment was to determine the time it took for light to travel from Fort Myer on the west bank of the Potomac river to a fixed mirror at the foot of the Washington monument 3721 meters away and back. (More details about this and similar examples can be found in Stigler (1977), Do robust estimators work with real data? (with discussion), Annals of Statistics, 5, 1055–1098.) Data from 64 replications of this experiment are provided in Table 11.5. For ease of handling the times in Table 11.5 are simplified. The values given in Table 11.5 are times expressed as billionths of a second in excess of 24.8 millionths of a second, *i.e.*, if a time value from Table 11.5 is multiplied by 10^{-3} and added to 24.8, the result is the time which Newcomb observed measured in millionths of a second. For example, the first observation in Table 11.5 is 28 which corresponds to an observed time of 24.828 millionths of a second.

Tab	ole 11	L.5. I	Newc	omb	's tin	ne da	ata.								
28	22	36	26	28	28	26	24	32	30	27	24	33	21	36	32
31	25	24	25	28	36	27	32	34	30	25	26	26	25	23	21
30	33	29	27	29	28	22	26	27	16	31	29	36	32	28	40
19	37	23	32	29	24	25	27	24	16	29	20	28	27	39	23

Figure 11.10 Newcomb's Time Data Histogram and QQ-plot





newcomb example: test H_0: mu = 33.02

The TTEST Procedure

Variable: time

Ν	Mean	Std De	v	Std E	rr	Minin	num	Maximum
64	27.7500	5.083	84	0.63	54	16.0	0000	40.0000
	[Mean		95%	CL	. Mean		
		27.7500		26.480	2	29.019	98	
	-							
		DF	t١	Value	Р	r > t		
		63		-8.29	<	.0001		

Based on the histogram and the QQ (normal probability) plot in Figure 11.10 it seems reasonable to model these data as a random sample from a normal population. More specifically, it seems reasonable to assume that these 64 measurements are independent realizations of a normally distributed variable with population mean μ . Actually, the

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assumption of independence is somewhat questionable, since repetitions of the experiment conducted at nearly the same point in time might exhibit some dependence. We will hope that any such dependence is minor and treat these observations as if they were independent.

The population mean μ can be thought of as the "true" time it would take light to make the "trip" as indicated for this experiment if this time could be measured very precisely. We can also view the population mean μ as the long run average of the times which would be obtained if this experiment was repeated over and over. We must keep in mind the fact that Newcomb's method of measurement may introduce a systematic bias which would make the "true" time μ differ from the actual time it would take for light to travel a distance of $7442 = 2 \times 3721$ meters.

Notice that the population mean time μ is actually defined by the process of measurement being used. That is, we can think of μ as the long run average time that we would observe if we replicated Newcomb's experiment a large times. We might wonder how this population mean time μ relates to the "true time" it would take for light to travel a distance of 7442 meters. We don't know exactly what this "true time" is, but we can use a generally accepted, modern measurement of the speed of light to obtain a hypothesized "true time." Stigler (*op. cit.*) used the modern estimate of the speed of light in a vacuum of 299,792.5 km/sec adjusted to give the speed of light in air and converted to a time as measured by Newcomb to obtain a hypothesized "true time" of 33.02. Therefore, our present goal is to determine how the population mean time μ relates to the "true time" $\mu_0 = 33.02$.

We do not have sufficient a priori information to specify a directional hypothesis; therefore, we will test the null hypothesis $H_0: \mu = 33.02$ versus the research hypothesis $H_1: \mu \neq 33.02$ to determine whether the population mean time μ is equal to or different from the hypothesized "true value" of 33.02. For Newcomb's n = 64 measurements the sample mean is $\overline{X} = 27.75$, the sample standard error of the mean is $\widehat{SE}(\overline{X}) = .6354$, and the calculated t statistic is

$$T_{calc} = \left| \frac{27.75 - 33.02}{.6354} \right| = 8.2940.$$

The *P*-value is less than .0001 indicating that observing a sample mean as far away from the hypothesized "true value" 33.02 (in either direction) as Newcomb did would be extremely unlikely if in fact $\mu = 33.02$. We can conclude that the population mean μ corresponding to Newcomb's experiment is almost certainly not equal to the hypothesized value of 33.02. The confidence interval, (26.4803, 29.0197), we computed above suggests that the mean μ

11.4 Testing a hypothesis about a normal population mean 265

 toc

that Newcomb was estimating is less than the hypothesized value 33.02 and in this sense his methodology overestimated the speed of light.

11.5 Problems

The problem sets and solutions can be found on the book web page at https://www.ucs.louisiana.edu/~jcb0773/.

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12 Inference for Two Means

12.1 Introduction

In Chapter 11 we considered inferential methods for the location of the center of the population distribution of a single continuous variable. We will now consider extensions of these methods to provide inferential methods for comparing the locations of the population distributions of two continuous variables. More specifically, we will consider methods for making inferences about the difference $\mu_1 - \mu_2$ between two population means μ_1 and μ_2 .

First consider a situation where the only difference between the population distributions of two continuous variables, X_1 and X_2 , is their location on the number line. In other words, suppose that the density curve for X_2 is identical to the density curve for X_1 except for its location on the number line. We will refer to this assumption about the population distributions of X_1 and X_2 as the **shift assumption**, since this assumption implies that the density curve for X_2 can be obtained by shifting the density curve for X_1 to the right or to the left along the number line. Under this shift assumption the difference, $\mu_1 - \mu_2$, between the two population means completely characterizes the difference between the two population distributions. Notice that under this shift assumption, if there is a positive constant d for which $\mu_1 - \mu_2 = d$ (*i.e.*, $\mu_1 = \mu_2 + d$), indicating that the density curve for X_1 is located d units to the right of the density curve for X_2 , then the difference $M_1 - M_2$ between the population medians, M_1 and M_2 , is also d (*i.e.*, $M_1 - M_2 = d$ and $M_1 = M_2 + d$). Therefore, under this shift assumption, a comparison of the locations of the two distributions based on the difference between the population means is equivalent to a comparison based on the population medians in the sense that the differences between each of these pairs of parameters is the same.

When the shift assumption is not valid, *i.e.*, when the two population distributions differ in aspects other than a simple shift in location, we must decide which parameter, say the population mean or the population median, is appropriate as a quantification of the location of each distribution and to quantify the difference between the locations of the two distributions. In other words, in the general situation when the shift assumption is not valid the difference between the population means and the difference between the population medians will be different and neither of these differences will completely describe the difference between the two distributions. For example, if the distribution of X_1 is skewed right and the distribution of X_2 is skewed left, it is possible for the population means to

toc

toc

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be equal while the population medians are different. Hence, when the shift assumption is not valid we must be careful about how we interpret an inference about the difference between any two particular location parameters, such as the population means, since the distributions differ in aspects other than a simple shift in location.

We will restrict our attention to methods that are appropriate when the data comprise two independent random samples; one random sample (the X_1 values) from a population with population mean μ_1 ; and, one random sample (the X_2 values) from a population with population mean μ_2 . The assumption that these random samples are independent basically means that the method used to select the random sample from the first population is not influenced by the method used to select the random sample from the second population, and *vice versa*.

12.2 Comparing the means of two normal populations toc

In this section we will assume that the population distribution of X_1 is a normal distribution with population mean μ_1 and population standard deviation σ_1 and the population distribution of X_2 is a normal distribution with population mean μ_2 and population standard deviation σ_2 . We will discuss methods for making inferences comparing the locations of these normal distributions as quantified by the difference, $\mu_1 - \mu_2$, between the two population means. As stated in the introduction, we will assume that the data comprise two independent random samples. Let n_1 denote the size of the random sample (of X_1 values) from the normal population with population mean μ_1 and let n_2 denote the size of the random sample (of X_2 values) from the normal population mean μ_1 and the sample mean \overline{X}_1 is the obvious estimate of the corresponding population mean μ_2 . Similarly, the difference, $\overline{X}_1 - \overline{X}_2$, between these two sample means is the obvious estimate of the corresponding difference, $\mu_1 - \mu_2$, between the population means. To describe the behavior of $\overline{X}_1 - \overline{X}_2$ as an estimator of $\mu_1 - \mu_2$ we need to know some properties of its sampling distribution.

Some properties of the sampling distribution of $\overline{\mathbf{X}}_1 - \overline{\mathbf{X}}_2$.

Let \overline{X}_1 denote the sample mean of a random sample of size n_1 from a distribution with population mean μ_1 and population standard deviation σ_1 and let \overline{X}_2 denote the sample mean of a random sample of size n_2 from a distribution with population mean μ_2 and population standard deviation σ_2 . Assume that these two random samples are independent. The sampling distribution of the difference, $\overline{X}_1 - \overline{X}_2$, between these two sample means has the following properties. The first two properties are valid in general and do not depend on the assumption of normal distributions.

1. The mean of the sampling distribution of $\overline{X}_1 - \overline{X}_2$ is the difference, $\mu_1 - \mu_2$, between the corresponding population means. Therefore, just as the sample means \overline{X}_1 and \overline{X}_2 are unbiased as estimators of μ_1 and μ_2 , respectively, the sample mean difference $\overline{X}_1 - \overline{X}_2$ is unbiased as an estimator of the population mean difference $\mu_1 - \mu_2$.

2. The population standard error of $\overline{X}_1 - \overline{X}_2$ (the standard deviation of the sampling distribution of $\overline{X}_1 - \overline{X}_2$) is

$$\operatorname{SE}(\overline{X}_1 - \overline{X}_2) = \sqrt{\frac{\sigma_1^2}{n_1} + \frac{\sigma_2^2}{n_2}}$$

This expression indicates how the variability of $\overline{X}_1 - \overline{X}_2$ depends on the sample sizes and population standard deviations. Notice that the population variance, $\operatorname{var}(\overline{X}_1 - \overline{X}_2)$ (the square of $\operatorname{SE}(\overline{X}_1 - \overline{X}_2)$), is equal to the sum of the population variance of \overline{X}_1 and the population variance of \overline{X}_2 . This property is a consequence of our assumption that the random samples are independent; and, this expression for the standard error of the difference between two sample means is not appropriate if the random samples are not independent.

3. If the random samples from which the sample means \overline{X}_1 and \overline{X}_2 are computed are random samples from **normal distributions** with population means and population standard deviations as given above, then in addition to the two properties above, the sampling distribution of $\overline{X}_1 - \overline{X}_2$ is a **normal distribution** with population mean $\mu_1 - \mu_2$ and population standard deviation SE $(\overline{X}_1 - \overline{X}_2)$.

The choice of the appropriate inferential methods for comparing the two normal population means μ_1 and μ_2 depends on the relationship between the two unknown, population standard deviations σ_1 and σ_2 . In particular, the choice of the appropriate estimate of the

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population standard error of $\overline{X}_1 - \overline{X}_2$ depends on whether the two population standard deviations σ_1 and σ_2 are equal.

Strictly speaking, the inferential methods based on the Student's t distribution described below are only appropriate when the data constitute independent random samples from normal populations (distributions). However, these methods are known to be generally reasonable even when the underlying populations are not exactly normal populations, provided the underlying population distributions are reasonably symmetric and the true density curves have a more or less normal (bell–shaped) appearance. We can use descriptive methods to look for evidence of possible nonnormality, provided the sample sizes are reasonably large. As in the one mean situation of Chapter 11, the most easily detected and serious evidence of nonnormality you should look for is evidence of extreme skewness or evidence of extreme outlying observations. If there is evidence of extreme skewness or extremly outlying observations, then inferential methods based on the Student's t distribution should not be used.

12.2a Inference when the two standard deviations are equal toc

A normal distribution is completely determined by its mean and standard deviation; therefore, in the present context of comparing two normal populations the shift assumption is equivalent to the assumption that the two population standard deviations σ_1 and σ_2 are equal. In other words, if we assume that $\sigma_1 = \sigma_2$, then the only difference between the two normal populations we are comparing is that the normal density curve for X_1 is centered at μ_1 and the normal density curve for X_2 is centered at μ_2 .

When the two population standard deviations are equal we can simplify the expression for the population standard error of $\overline{X}_1 - \overline{X}_2$. If we let $\sigma = \sigma_1 = \sigma_2$ denote the common value of the two population standard deviations, then the population standard error of $\overline{X}_1 - \overline{X}_2$ is

$$\operatorname{SE}(\overline{X}_1 - \overline{X}_2) = \sqrt{\sigma^2 \left(\frac{1}{n_1} + \frac{1}{n_2}\right)} = \sigma \sqrt{\frac{1}{n_1} + \frac{1}{n_2}}.$$

An appropriate estimator of the common standard deviation σ is the **pooled sample** standard deviation which we will denote by S_p . Recall that the sample standard deviation S_X for a single sample of n values of the variable X is the square root of the "average" of the squared deviations of the observed values of X from the sample mean \overline{X} ,

$$S_X = \sqrt{\frac{\sum (X - \overline{X})^2}{n - 1}}.$$

In the present context, the n_1 values of X_1 have sample mean \overline{X}_1 and the n_2 values of X_2 have sample mean \overline{X}_2 ; therefore, the sum of squared deviations in the formula for S_X is replaced by the sum of two such sums of squared deviations, one for each sample. The divisor n-1 in the formula for S_X is replaced by the total number of observations $n_1 + n_2$ decreased by 2, *i.e.*, the one sample divisor n-1 is replaced by the two sample divisor $(n_1-1)+(n_2-1)=n_1+n_2-2$. The resulting formula for the **pooled sample standard deviation** is

$$S_p = \sqrt{\frac{\sum (X_1 - \overline{X}_1)^2 + \sum (X_2 - \overline{X}_2)^2}{n_1 + n_2 - 2}},$$

This pooled sample standard deviation can also be expressed in terms of the two sample standard deviations S_1 and S_2 as shown below

$$S_p = \sqrt{\frac{(n_1 - 1)S_1^2 + (n_2 - 1)S_2^2}{n_1 + n_2 - 2}}.$$

Strictly speaking, the inferential methods described in this subsection are only valid when the two population standard deviations σ_1 and σ_2 are equal. However, in practice these methods still perform reasonably well provided the two population standard deviations σ_1 and σ_2 are reasonably close to being equal and the two sample sizes n_1 and n_2 are reasonably similar. (This assumption is more critical when the sample sizes are very dissimilar.) A common rule of thumb for assessing the assumption of equal standard deviations says that the assumption of a common population standard deviation is reasonable if the ratio of the sample standard deviations is between 1/2 and 2.

When $\sigma_1 = \sigma_2$, the appropriate **sample standard error** of $\overline{X}_1 - \overline{X}_2$, based on the pooled sample standard deviation S_p , is

$$\widehat{\operatorname{SE}}(\overline{X}_1 - \overline{X}_2) = S_p \sqrt{\frac{1}{n_1} + \frac{1}{n_2}},$$

and the corresponding 95% margin of error of $\overline{X}_1 - \overline{X}_2$ is

$$\operatorname{ME}(\overline{X}_1 - \overline{X}_2) = k\widehat{\operatorname{SE}}(\overline{X}_1 - \overline{X}_2) = kS_p \sqrt{\frac{1}{n_1} + \frac{1}{n_2}},$$

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where k is the 97.5 percentile of the Student's t distribution with $n_1 + n_2 - 2$ degrees of freedom. Thus the interval from $(\overline{X}_1 - \overline{X}_2) - ME(\overline{X}_1 - \overline{X}_2)$ to $(\overline{X}_1 - \overline{X}_2) + ME(\overline{X}_1 - \overline{X}_2)$ is a 95% confidence interval for $\mu_1 - \mu_2$.

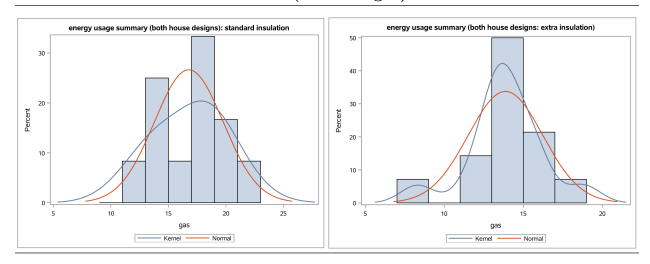
Example 12.1. Energy consumption. The data used in this example are part of data set 93 in Hand, Daly, Lunn, McConway, and Ostrowski (1994), A Handbook of Small Data Sets, Chapman and Hall, London. The original source is two reports issued in 1983 and 1984 by the Open University. A large–scale experiment on energy consumption was conducted in the early 1980's in the Pennyland district of Milton Keynes. A housing development of about 180 houses was built. About half of the houses had a standard level of roof and wall insulation. The others had extra roof and wall insulation (these houses also had double glazing and under-floor insulation). In addition to the differences in level of insulation, many of the houses were designed with passive solar heating features, e.g., southern orientation with most of the windows on the south side. The other houses had a more traditional design. Energy consumption was monitored over several years. Table 12.1 provides the annual gas consumption (in 1000 kWh) for two independent random samples of houses. One random sample was selected from all of the houses with standard insulation (regardless of design type) and the other was selected from all of the houses with extra insulation (regardless of design type). Histograms, with fitted density curves and fitted normal curves are given in Figure 12.1, and QQ (normal probability) plots are provided in Figure 12.2.

Tabl	Table 12.1. Gas consumption data (1000 kWh) (both designs).													
standard insulation								extra insulation						
11.4	13.9	13.9	14.0	15.3	18.0		8.3	11.7	12.7	13.0	13.4	13.6	13.7	
18.0	18.1	18.9	19.0	19.0	21.7		13.7	13.8	14.6	15.3	15.6	16.0	18.8	

The histograms and fitted density curves show some evidence of slight skewness to the left in the distribution for houses with extra insulation. Both histograms appear to be unimodal and reasonably symmetric with mild outliers on both sides. Both QQ plots are reasonably linear. Thus it seems reasonable to model these data as independent random samples from normal distributions. Furthermore, the two sample standard deviations, 2.9959 and 2.3636 (these can be found in the first table in Figure 12.3), are quite similar

(ratio .7889); therefore, we can also reasonably assume that the two population standard deviations are equal.

Figure 12.1. Histograms (with smooth curve and normal fits) for gas consumption for houses with standard and extra insulation (both designs).



All of the values computed below can be found in the SAS output of Figure 12.3. Letting X_1 denote the annual gas consumption for a house with extra insulation and X_2 denote annual gas consumption for a house with standard insulation, we find that the difference in the sample means is $\overline{X}_1 - \overline{X}_2 = 13.8714 - 16.7667 = -2.8953$ (2,895.3 kWh) suggesting that, among these 180 houses, the mean annual gas consumption for a house with extra insulation μ_1 is approximately 2.8953 thousand kW hours lower than the mean annual gas consumption for a house with standard insulation μ_2 . The pooled sample standard deviation $S_p = 2.6720$, the standard error $\widehat{SE}(\overline{X}_1 - \overline{X}_2) = 1.0512$, and the margin of error multiplier k = 2.064 for the Student's t distribution with $n_1 + n_2 - 2 = 24$ degrees of freedom yield the 95% confidence interval (-5.0648, -.7258) for $\mu_1 - \mu_2$. Thus we are 95% confident that, among these 180 houses, the population mean annual gas consumption for a house with extra insulation is at least 725.8 kW hours and as much as 5,064.8 kW hours lower than the mean annual gas consumption for a house with standard insulation. Note that, technically, this inference is restricted to these 180 houses but we might conjecture that a similar difference would occur for similar houses (with standard and extra insulation) in this same area.

Figure 12.2. QQ plots for gas consumption for houses with standard and extra insulation (both designs).

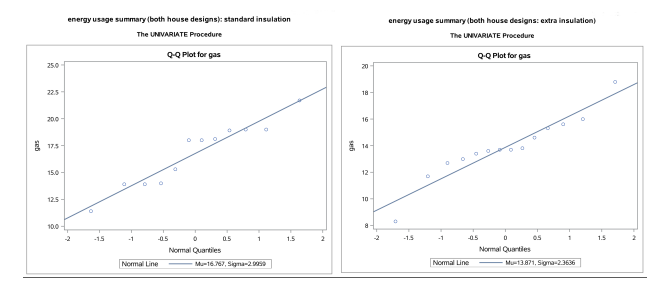


Figure 12.3. SAS output for gas consumption for houses example standard insulation versus extra insulation (both designs).

energy usage summary (both house designs)

The TTEST Procedure

Variable: gas

type	Method	N	Mean	Std Dev	Std Err	Minimum	Maximum
extra		14	13.8714	2.3636	0.6317	8.3000	18.8000
standard		12	16.7667	2.9959	0.8648	11.4000	21.7000
Diff (1-2)	Pooled		-2.8952	2.6720	1.0512		
Diff (1-2)	Satterthwaite		-2.8952		1.0710		

type	Method	Mean	95% C	% CL Mean	
extra		13.8714	12.5067	15.2362	
standard		16.7667	14.8632	18.6701	
Diff (1-2)	Pooled	-2.8952	-5.0648	-0.7257	
Diff (1-2)	Satterthwaite	-2.8952	-5.1234	-0.6670	

Method	Variances	DF	t Value	Pr > t 0.0110	
Pooled	Equal	24	-2.75		
Satterthwaite	Unequal	20.848	-2.70	0.0134	

Remark regarding directional confidence bounds. We can find an upper or lower 95% confidence bound for $\mu_1 - \mu_2$ by selecting the appropriate confidence limit from a 90% confidence interval estimate of $\mu_1 - \mu_2$.

When $\sigma_1 = \sigma_2$, we can use the **two sample Student's t test statistic**

$$T = \frac{\overline{X}_1 - \overline{X}_2}{\widehat{\operatorname{SE}}(\overline{X}_1 - \overline{X}_2)} = \frac{\overline{X}_1 - \overline{X}_2}{S_p \sqrt{\frac{1}{n_1} + \frac{1}{n_2}}},$$

based on the standard error computed using the pooled estimate of the common standard deviation, to test hypotheses relating μ_1 to μ_2 .

First consider a situation where we want to determine whether there is sufficient evidence to conclude that the population mean μ_1 exceeds the population mean μ_2 . Our research hypothesis is the contention that the population mean μ_1 exceeds the population mean μ_2 , *i.e.*, $H_1 : \mu_1 > \mu_2$. The corresponding null hypothesis is $H_0 : \mu_1 \leq \mu_2$. Values of $\overline{X}_1 - \overline{X}_2$ which are large relative to zero provide evidence in favor of $H_1 : \mu_1 > \mu_2$, since this hypothesis is equivalent to $H_1 : \mu_1 - \mu_2 > 0$, and against $H_0 : \mu_1 \leq \mu_2$. Since large values of $\overline{X}_1 - \overline{X}_2$ yield large values of the Student's *t* statistic, we will reject $H_0 : \mu_1 \leq \mu_2$ in favor of $H_1 : \mu_1 > \mu_2$ if the calculated Student's *t* statistic is sufficiently large. This decision will hinge on the size of the *P*-value, which is the probability, computed under the assumption that $\mu_1 = \mu_2$, that $\overline{X}_1 - \overline{X}_2$ is as large or larger than the value actually observed and is equal to the probability that a Student's *t* variable with $n_1 + n_2 - 2$ degrees of freedom is as large or larger than the calculated *t* value T_{calc} . Notice that this *P*-value is the area to the right of T_{calc} under the density curve of the Student's *t* distribution with $n_1 + n_2 - 2$ degrees of freedom, since values of $\overline{X}_1 - \overline{X}_2$ that are sufficiently far above zero provide evidence in favor of the research hypothesis.

The steps for performing a hypothesis test for

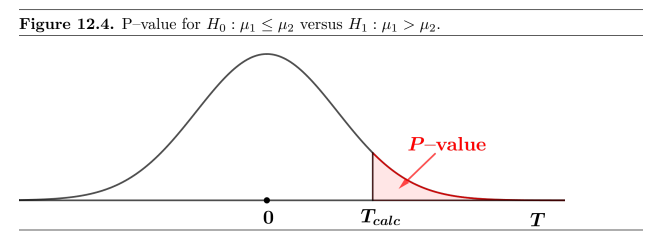
$$H_0: \mu_1 \le \mu_2$$
 versus $H_1: \mu_1 > \mu_2$

are summarized below.

1. Use a suitable calculator or computer program to find the *P*-value = $P(T \ge T_{calc})$, where *T* denotes a Student's *t* variable with $n_1 + n_2 - 2$ degrees of freedom and $T_{calc} = (\overline{X}_1 - \overline{X}_2)/\widehat{\operatorname{SE}}(\overline{X}_1 - \overline{X}_2)$ as described above. This *P*-value is the area to the right of T_{calc}

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under the density curve for the Student's t distribution with $n_1 + n_2 - 2$ degrees of freedom as shown in Figure 12.4.



2a. If the *P*-value is small enough (less than .05 for a test at the 5% level of significance), conclude that the data favor $H_1: \mu_1 > \mu_2$ over $H_0: \mu_1 \leq \mu_2$. That is, if the *P*-value is small enough, then there is sufficient evidence to conclude that the first population mean μ_1 is greater than the second population mean μ_2 .

2b. If the *P*-value is not small enough (is not less than .05 for a test at the 5% level of significance), conclude that the data do not favor $H_1: \mu_1 > \mu_2$ over $H_0: \mu_1 \leq \mu_2$. That is, if the *P*-value is not small enough, then there is not sufficient evidence to conclude that the first population mean μ_1 is greater than the second population mean μ_2 .

The procedure for testing the null hypothesis $H_0: \mu_1 \leq \mu_2$ versus the research hypothesis $H_1: \mu_1 > \mu_2$ given above is readily modified for testing the null hypothesis $H_0: \mu_1 \geq \mu_2$ versus the research hypothesis $H_1: \mu_1 < \mu_2$. The essential modification is to change the direction of the inequality in the definition of the P-value. Consider a situation where the research hypothesis specifies that the population mean μ_1 is less than the population mean μ_2 . Values of $\overline{X}_1 - \overline{X}_2$ that are sufficiently far from 0 in the negative direction provide evidence in favor of the research hypothesis $H_1: \mu_1 < \mu_2$ and against the null hypothesis $H_0: \mu_1 \geq \mu_2$. Therefore, the appropriate P-value is the probability of observing a value of $\overline{X}_1 - \overline{X}_2$ as small or smaller than the value actually observed. As before, the P-value is computed under the assumption that $\mu_1 = \mu_2$. The calculated t statistic T_{calc} is defined as before; however, in this situation the P-value is the area to the left of T_{calc} under the density curve of the Student's t distribution with $n_1 + n_2 - 2$ degrees of freedom, since

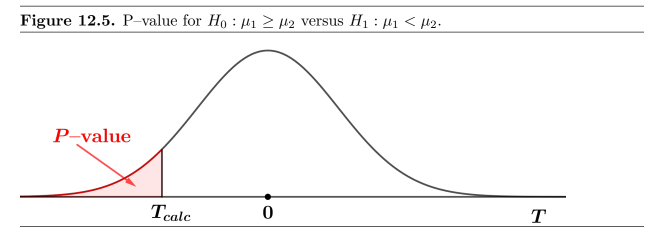
values of $\overline{X}_1 - \overline{X}_2$ that are sufficiently far below zero provide evidence in favor of the research hypothesis.

The steps for performing a hypothesis test for

$$H_0: \mu_1 \ge \mu_2$$
 versus $H_1: \mu_1 < \mu_2$

are summarized below.

1. Use a suitable calculator or computer program to find the *P*-value = $P(T \leq T_{calc})$, where *T* denotes a Student's *t* variable with $n_1 + n_2 - 2$ degrees of freedom and $T_{calc} = (\overline{X}_1 - \overline{X}_2)/\widehat{SE}(\overline{X}_1 - \overline{X}_2)$ as before. This *P*-value is the area to the left of T_{calc} under the density curve for the Student's *t* distribution with $n_1 + n_2 - 2$ degrees of freedom as shown in Figure 12.5.



2a. If the *P*-value is small enough (less than .05 for a test at the 5% level of significance), conclude that the data favor $H_1: \mu_1 < \mu_2$ over $H_0: \mu_1 \ge \mu_2$. That is, if the *P*-value is small enough, then there is sufficient evidence to conclude that the first population mean μ_1 is less than the second population mean μ_2 .

2b. If the *P*-value is not small enough (is not less than .05 for a test at the 5% level of significance), conclude that the data do not favor $H_1 : \mu_1 < \mu_2$ over $H_0 : \mu_1 \ge \mu_2$. That is, if the *P*-value is not small enough, then there is not sufficient evidence to conclude that the first population mean μ_1 is less than the second population mean μ_2 .

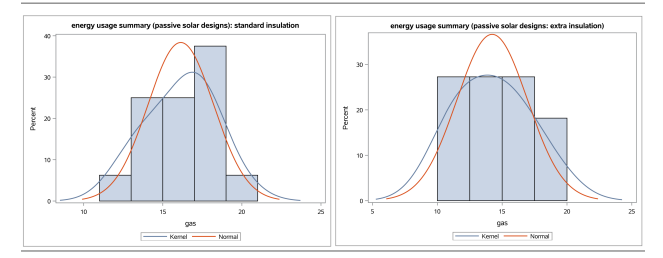
Example 12.1. Energy consumption — revisited. We will now consider the reduction in energy consumption due to extra insulation when the population is restricted to the houses

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among the 180 houses which have passive solar designs. Table 12.2 provides the annual gas consumption (in 1000 kWh) for two independent random samples of houses. One random sample was selected from all of the passive solar houses with standard insulation and the other was selected from all of the passive solar houses with extra insulation. Histograms, with fitted density curves and fitted normal curves are given in Figure 12.6, and QQ (normal probability) plots are provided in Figure 12.7.

Tabl	Table 12.2. Gas consumption data (1000 kWh) (passive solar).												
standard insulation						ext	tra ins	sulatio	on				
12.3	13.3	13.7	13.8	14.9	15.6	15.9	16.3	10.5	11.3	11.4	12.6	13.0	14.5
$\underline{16.5 \ 17.2 \ 17.5 \ 17.6 \ 17.8 \ 17.9 \ 18.0 \ 19.9 \ 15}$								15.2	15.7	15.7	17.6	19.0	

Figure 12.6. Histograms (with smooth curve and normal fits) for gas consumption for houses with standard and extra insulation (passive solar designs).



In this example both stem and leaf histograms appear to be unimodal and reasonably symmetric. The QQ plots are reasonably linear. Thus it seems reasonable to model these data as independent random samples from normal distributions. The two sample standard deviations, 2.0791 and 2.7225, are quite similar (ratio .7637); therefore, we can also reasonably assume that the two population standard deviations are equal.

Figure 12.7. QQ plots for gas consumption for houses with standard and extra insulation (passive solar designs).

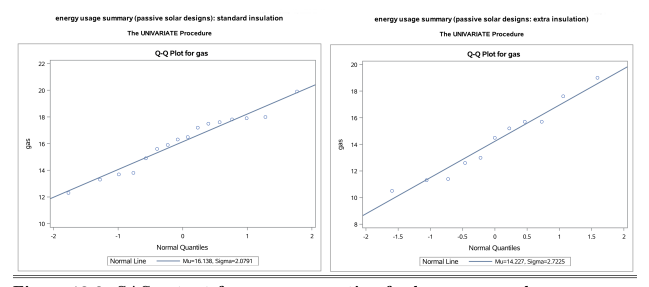


Figure 12.8. SAS output for gas consumption for houses example standard insulation versus extra insulation (passive solar designs).

energy usage summary (solar house designs)

The TTEST Procedure

			variab	ie. yas	7		
type	Method	N	Mean	Std Dev	Std Err	Minimum	Maximum
extra		11	14.2273	2.7225	0.8209	10.5000	19.0000
standard		16	16.1375	2.0791	0.5198	12.3000	19.9000
Diff (1-2)	Pooled		-1.9102	2.3576	0.9234		
Diff (1-2)	Satterthwaite		-1.9102		0.9716		

type	Method	Mean	95% CL Mean		
extra		14.2273	12.3983	16.0563	
standard		16.1375	15.0296	17.2454	
Diff (1-2)	Pooled	-1.9102	-3.8120	-0.00841	
Diff (1-2)	Satterthwaite	-1.9102	-3.9537	0.1333	

Method	Variances	DF	t Value	Pr > t
Pooled	Equal	25	-2.07	0.0491
Satterthwaite	Unequal	17.726	-1.97	0.0652

All of the values computed below can be found in the first table in Figure 12.8. Let X_1 denote the annual gas consumption for a passive solar house with extra insulation and

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let X_2 denote annual gas consumption for a passive solar house with standard insulation. Similarly, let μ_1 and μ_2 denote the respective population means for all of the passive solar houses among the 180 houses with extra and standard insulation. The obvious research hypothesis $H_1: \mu_1 < \mu_2$ states that among the passive solar houses in this development, on average, the annual gas consumption is lower for a house with extra insulation than it is for a house with standard insulation. For these data the pooled sample standard deviation is $S_p = 2.3576$, the standard error is $\widehat{SE}(\overline{X}_1 - \overline{X}_2) = .9234$, the observed value of the Student's t statistic is $T_{calc} = -2.07$ with 25 degrees of freedom, and the corresponding Pvalue is $\frac{.0491}{2} = .0245$. This *P*-value is reasonably small indicating that there is reasonably strong evidence that μ_1 is less than μ_2 . Therefore, there is reasonably strong evidence that for this population of passive solar houses, on average, the annual gas consumption for a passive solar house with extra insulation is lower than the annual gas consumption for a passive solar house with standard insulation. We can form a 95% confidence interval for $\mu_1 - \mu_2$ to get a feel for the practical importance of this result. Using the margin of error multiplier k = 2.060 for the Student's t distribution with 25 degrees of freedom yields the 95% confidence interval (-3.8120, -.00841) for $\mu_1 - \mu_2$. Thus we are 95% confident that, among this population of passive solar houses, the population mean annual gas consumption for a house with extra insulation is between 8.41 kW hours and 3.812 kW hours lower than the mean annual gas consumption for a house with standard insulation. Notice that this confidence interval estimate indicates that the difference between these means might be as small as 8.41 kW hours which is not much of a difference. Of course, the confidence interval estimate also allows that the difference in these means might be as large as 3,812 kW hours which is more impressive. In this case, technically, our inferences are restricted to all of the passive solar houses among these 180 houses.

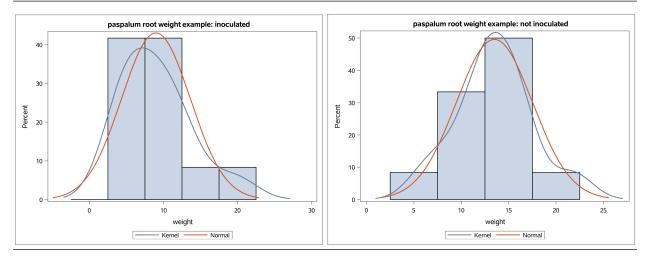
Example 12.2. Paspalum grass. This example is taken from Seber (1984), Multivariate Observations, Wiley, New York. (The data were provided by Peter Buchanan.) Paspalum grass is a weed which grows in pastures used for grazing farm animals. Scientists at the Mount Albert Research Centre in Auckland conducted a laboratory experiment to determine whether inoculation of paspalum with a fungal infection might be effective in reducing the growth of this weed. The experimenters randomly assigned 48 pots of paspalum to the 8 combinations of treatment (inoculated, not inoculated) and temperature (14, 18, 22, 26 degrees C). For our purposes we will restrict our attention to the 24 pots of plants grown under moderate temperatures (18 or 22 degrees) and we will not distinguish

between the two temperatures. Thus we have two samples of size 12. The experimenters measured several characteristics of the paspalum. The response variable we will consider is the fresh weight of the roots (in grams) of the paspalum in a pot. (In this example a pot of paspalum is a unit; the number of plants per pot is not specified.) Table 12.3 provides the fresh root weights for the 12 pots assigned to each treatment. Histograms, with fitted density curves and fitted normal curves are given in Figure 12.9, and QQ (normal probability) plots are provided in Figure 12.10.

Table 12.3.	Paspalum	\mathbf{root}	weight	(grams)	•
-------------	----------	-----------------	--------	---------	---

inoculated					no	ot ino	culate	ed			
3.9	4.3	4.9	5.2	6.5	7.6	6.2	8.7	11.0	12.2	12.3	13.1
9.6	10.0	10.1	12.3	13.6	19.7	13.6	14.5	15.4	16.4	16.7	21.8

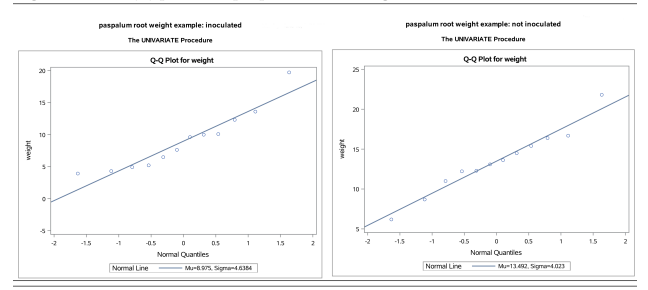
Figure 12.9. Histograms (with smooth curve and normal fits) for paspalum root weight.

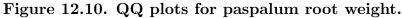


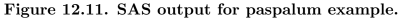
Let X_1 denote the fresh root weight for a pot of paspalum inoculated with the fungus and let X_2 denote the fresh root weight for a pot of paspalum not inoculated with the fungus. We can think of the corresponding population means μ_1 and μ_2 as the mean fresh root weights we would observe if all 48 of the pots of paspalum had been inoculated (μ_1) or not inoculated (μ_2). We want to determine whether there is sufficient evidence to claim that inoculation with this fungus retards the growth of paspalum in the sense of reducing fresh root weight. In terms of the population means the research hypothesis $H_1 : \mu_1 < \mu_2$ states that, for this collection of 48 pots of paspalum, on average, the fresh root weight would be

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smaller if the paspalum were inoculated with the fungus than it would be if the paspalum were not inoculated.







paspalum root weight example

The TTEST Procedure

		V	ariable:	weight				
treat	Method	N	Mean	Std Dev	S	d Err	Minimum	Maximum
inoculated		12	8.9750	4.6384	1	.3390	3.9000	19.7000
notinoculated		12	13.4917	4.0230	1	.1613	6.2000	21.8000
Diff (1-2)	Pooled		-4.5167	4.3416	1	.7725		
Diff (1-2)	Satterthwaite		-4.5167		1	.7725		

treat	Method	Mean	95% CL Mean		
inoculated		8.9750	6.0279	11.9221	
notinoculated		13.4917	10.9356	16.0478	
Diff (1-2)	Pooled	-4.5167	-8.1925	-0.8408	
Diff (1-2)	Satterthwaite	-4.5167	-8.1968	-0.8366	

Method	Variances	DF	t Value	Pr > t	
Pooled	Equal	22	-2.55	0.0183	
 Satterthwaite	Unequal	21.569	-2.55	0.0185	

Both of the histograms are unimodal and both show some evidence of slight skewness to the right. Each sample contains a mild outlier (19.7 for the inoculated sample and 21.8 for the not inoculated sample). Closer examination indicates that it is these outliers which give

the impression of skewness to the right. The QQ plots are reasonably linear suggesting that skewness is not a problem. Thus it seems reasonable to model these data as independent random samples from normal distributions. The two sample standard deviations, 4.6384 and 4.0230, are quite similar (ratio .8673); therefore, we can also reasonably assume that the two population standard deviations are equal.

For these data the pooled sample standard deviation is $S_p = 4.3416$, the standard error is $\widehat{SE}(\overline{X}_1 - \overline{X}_2) = 1.7725$, the observed value of the Student's *t* statistic is $T_{calc} = -2.55$ with 22 degrees of freedom, and the corresponding *P*-value is $\frac{.0183}{2} = .0092$. This *P*-value is very small indicating that there is very strong evidence that μ_1 is less than μ_2 . Therefore, there is very strong evidence that for this collection of 48 pots of paspalum, on average, the fresh root weight would be smaller if the paspalum was inoculated with the fungus than it would be if the paspalum was not inoculated.

Using the margin of error multiplier k = 2.074 for the Student's t distribution with 22 degrees of freedom yields the 95% confidence interval (-8.193, -.8410) for $\mu_1 - \mu_2$. Thus we are 95% confident that, for this collection of 48 pots of paspalum, the mean fresh root weight we would observe if all 48 of the pots of paspalum had been inoculated is between .8410 grams and 8.1930 grams smaller than the mean fresh root weight we would observe if none of the 48 of the pots of paspalum had been inoculated.

The directional hypothesis tests we discussed above are readily modified for testing a nondirectional hypothesis. To decide between the null hypothesis $H_0: \mu_1 = \mu_2$ and the research hypothesis $H_1: \mu_1 \neq \mu_2$, we need to decide whether $\overline{X}_1 - \overline{X}_2$ supports the null hypothesis by being "close to 0", or supports the research hypothesis by being "far away from 0". In this situation the P-value is the probability that $\overline{X}_1 - \overline{X}_2$ would be as far or farther away from 0 in either direction as is the value that we actually observe. In other words, the P-value is the probability that the distance $|\overline{X}_1 - \overline{X}_2|$ between the two sample means (the absolute value of the difference between \overline{X}_1 and \overline{X}_2) is as large or larger than the actual observed value of this distance. As before, the P-value is computed under the assumption that the null hypothesis is true and $\mu_1 = \mu_2$. In this situation the calculated t statistic T_{calc} is the absolute value of the t statistic that would be used for testing a directional hypothesis. That is, the calculated t statistic is

$$T_{calc} = \frac{|\overline{X}_1 - \overline{X}_2|}{\widehat{\operatorname{SE}}(\overline{X}_1 - \overline{X}_2)}$$

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In terms of this t statistic the P-value is the probability that the absolute value of a Students's t variable with $n_1 + n_2 - 2$ degrees of freedom would take on a value as large or larger than T_{calc} , computed assuming that $\mu_1 = \mu_2$. This probability is the sum of the area under the appropriate Student's t density curve to the left of $-T_{calc}$ and the area under this Student's t density curve to the right of T_{calc} . We need to add these two areas (probabilities) since we are finding the probability that $\overline{X}_1 - \overline{X}_2$ would be as far or farther away from 0 in either direction as is the value that we actually observe, when $\mu_1 = \mu_2$.

The steps for performing a hypothesis test for

$$H_0: \mu_1 = \mu_2$$
 versus $H_1: \mu_1 \neq \mu_2$

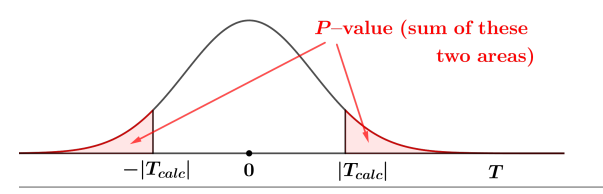
are summarized below.

1. Use a suitable calculator or computer program to find the P-value = $P(|T| \ge T_{calc}) = P(T \le -T_{calc}) + P(T \ge T_{calc})$, where T denotes a Student's t variable with $n_1 + n_2 - 2$ degrees of freedom and

$$T_{calc} = \frac{|\overline{X}_1 - \overline{X}_2|}{\widehat{\operatorname{SE}}(\overline{X}_1 - \overline{X}_2)}.$$

Notice that this calculated t value is the absolute value of the calculated t value we would use for a directional hypothesis. This P-value is the area, under the density curve for the Student's t distribution with $n_1 + n_2 - 2$ degrees of freedom, to the left of $-T_{calc}$ plus the area to the right of T_{calc} as shown in Figure 12.12.

Figure 12.12. P-value for $H_0: \mu_1 = \mu_2$ versus $H_1: \mu_1 \neq \mu_2$.



2a. If the *P*-value is small enough (less than .05 for a test at the 5% level of significance), conclude that the data favor $H_1: \mu_1 \neq \mu_2$ over $H_0: \mu_1 = \mu_2$. That is, if the *P*-value is

small enough, then there is sufficient evidence to conclude that the first population mean μ_1 and the second population mean μ_2 are different.

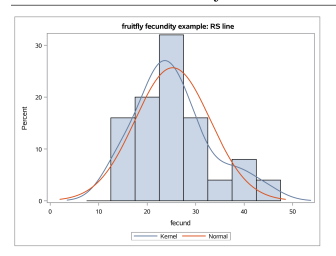
2b. If the *P*-value is not small enough (is not less than .05 for a test at the 5% level of significance), conclude that the data do not favor $H_1: \mu_1 \neq \mu_2$ over $H_0: \mu_1 = \mu_2$. That is, if the *P*-value is not small enough, then there is not sufficient evidence to conclude that the population means μ_1 and μ_2 are different.

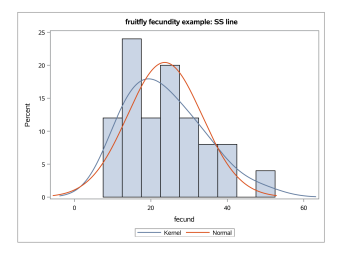
Example 12.3. Fecundity of fruitflies. Sokal, R.R. and Rohlf, F.J. (1969) Biometry, W.H. Freeman, p.232, discuss a study conducted to compare the fecundity of three genetic lines of Drosophila melanogaster. The data in Table 12.4 consist of per diem fecundities (number of eggs laid per female per day for the first 14 days of life) for 25 females of three lines of Drosophila melanogaster. Two of these genetic lines were selected for resistance (RS) and susceptibility (SS) to DDT, the third line is a nonselected control (NS). These data can be used to address two questions which were of interest to the investigator. We can use the data for the two selected lines (RS and SS) to determine if there is evidence that the mean fecundity differs for these selected lines. We can then use the data for the control line (NS) to compare the mean fecundity of the control line with that of the two selected lines. For the time being we will use two-sample Student's t tests to address these questions.

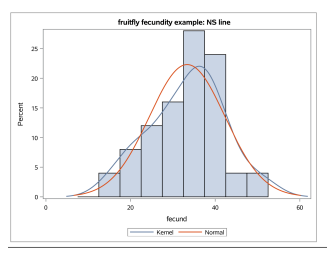
resistant RS		suscej S		nonselected NS		
12.8	22.4	38.4	23.1	35.4	22.6	
21.6	27.5	32.9	29.4	27.4	40.4	
14.8	20.3	48.5	16.0	19.3	34.4	
23.1	38.7	20.9	20.1	41.8	30.4	
34.6	26.4	11.6	23.3	20.3	14.9	
19.7	23.7	22.3	22.9	37.6	51.8	
22.6	26.1	30.2	22.5	36.9	33.8	
29.6	29.5	33.4	15.1	37.3	37.9	
16.4	38.6	26.7	31.0	28.2	29.5	
20.3	44.4	39.0	16.9	23.4	42.4	
29.3	23.2	12.8	16.1	33.7	36.6	
14.9	23.6	14.6	10.8	29.2	47.4	
27.3		12.2		41.7		

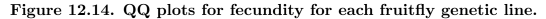
Table 12.4. Fruitfly fecundity data.

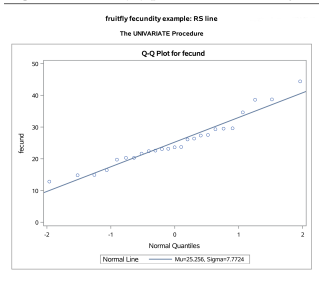
Figure 12.13. Histograms (with smooth curve and normal fits) for fecundity for each fruitfly genetic line.

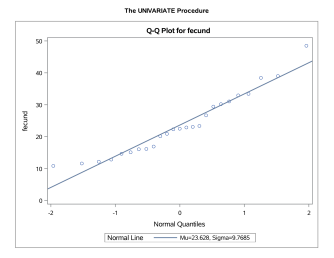












fruitfly fecundity example: SS line

fruitfly fecundity example: NS line The UNIVARIATE Procedure Q-Q Plot for fecund 60 50 40 fecund 30 20 10 -1 0 Normal Quantiles Normal Line — Mu=33.372, Sigma=8.942 -

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Let X_{RS} denote the fecundity for an RS female, X_{SS} the fecundity for an SS female, and X_{NS} the fecundity for an NS female; and let μ_{RS}, μ_{SS} , and μ_{NS} denote the corresponding population means. The first question, concerning the relationship between the population mean fecundities μ_{RS} and μ_{SS} , can be addressed via a test of $H_0: \mu_{RS} = \mu_{SS}$ versus $H_1: \mu_{RS} \neq \mu_{SS}$. Our approach to the second question will depend on our conclusion for the first. If we decide that there is no difference between the two selected line population mean fecundities ($\mu_{RS} = \mu_{SS}$), then we can combine the data for these two lines and, viewing this as a random sample from a population of selected lines with population mean μ_S , we can test for a difference between the population mean for selected lines and the population mean for the nonselected line by testing $H_0: \mu_S = \mu_{NS}$ versus $H_1: \mu_S \neq \mu_{NS}$. On the other hand, if we decide that there is a difference between the population mean fecundities for the two selected lines, then we will need to perform two tests; one for comparing μ_{RS} to μ_{NS} and another for comparing μ_{SS} to μ_{NS} .

Figure 12.15. SAS output for fruitfly eample RS line versus SS line.

fruitfly fecundity example: RS versus SS

The TTEST Procedure Variable: fecund

line	Method	N	Mean	Std Dev	Std Err	Minimum	Maximum
RS		25	25.2560	7.7724	1.5545	12.8000	44.4000
SS		25	23.6280	9.7685	1.9537	10.8000	48.5000
Diff (1-2)	Pooled		1.6280	8.8270	2.4967		
Diff (1-2)	Satterthwaite		1.6280		2.4967		

line	Method	Mean	95% CL Mean		
RS		25.2560	22.0477	28.4643	
SS		23.6280	19.5958	27.6602	
Diff (1-2)	Pooled	1.6280	-3.3919	6.6479	
Diff (1-2)	Satterthwaite	1.6280	-3.3984	6.6544	

Method	Variances	DF	t Value	Pr > t
Pooled	Equal	48	0.65	0.5175
Satterthwaite	Unequal	45.693	0.65	0.5176

Figure 12.16. SAS output for fruitfly eample NS line versus S line.

fruitfly fecundity example: NS versus S

The TTEST Procedure

Variable: fecund

line2	Method	N	Mean	Std Dev	Std Err	Minimum	Maximum
NS		25	33.3720	8.9420	1.7884	14.9000	51.8000
S		50	24.4420	8.7751	1.2410	10.8000	48.5000
Diff (1-2)	Pooled		8.9300	8.8303	2.1630		
Diff (1-2)	Satterthwaite		8.9300		2.1768		

line2	Method	Mean	95% CL Mean		
NS		33.3720	29.6809	37.0631	
s		24.4420	21.9481	26.9359	
Diff (1-2)	Pooled	8.9300	4.6192	13.2408	
Diff (1-2)	Satterthwaite	8.9300	4.5516	13.3084	

Method	Variances	DF	t Value	Pr > t
Pooled	Equal	73	4.13	<.0001
Satterthwaite	Unequal	47.305	4.10	0.0002

The histograms in Figure 12.13 indicate that the fecundity distributions for the two selected lines (RS and SS) are unimodal with some evidence of skewness to the right; and the fecundity distribution for the nonselected line (NS) is unimodal and reasonably symmetric with slight evidence of skewness to the left in the middle of the distribution. The QQ plots in Figure 12.14 are reasonably linear. Thus it seems reasonable to treat these samples as forming independent random samples from normal populations. The three sample standard deviations are reasonably similar allowing us to also assume a common population standard deviation. Therefore, we will model the three population distributions as normal distributions with respective population means μ_{RS} , μ_{SS} and μ_{NS} and with common population standard deviation. If we decide to combine the samples from the two selected lines, we will model the corresponding population distribution as a normal distribution with population mean μ_S and the same common population standard deviation as before.

The difference between the sample mean fecundities for the two selected lines $\overline{X}_{RS} - \overline{X}_{SS} =$ 1.628 is small relative to the corresponding standard error $\widehat{SE}(\overline{X}_{RS} - \overline{X}_{SS}) = 2.4967$ suggesting that there is little evidence of a difference between the population means μ_{RS} and μ_{SS} . The observed value of the Student's t statistic for testing $H_0: \mu_{RS} = \mu_{SS}$ versus

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 $H_1: \mu_{RS} \neq \mu_{SS}$ is $T_{calc} = .65$ with 48 degrees of freedom, and the corresponding P-value is .5175. This large P-value allows us to conclude that the two population mean fecundities μ_{RS} and μ_{SS} are equal. In light of this conclusion we will now combine the samples for the selected lines as described above and test $H_0: \mu_S = \mu_{NS}$ versus $H_1: \mu_S \neq \mu_{NS}$. Recall that μ_S denotes the population mean fecundity for the population of fruitflies obtained by combining the populations for the two selected lines. The difference between the sample mean fecundities for the combined population of selected lines and the nonselected line is $\overline{X}_S - \overline{X}_{NS} = -8.93$ with an associated standard error of $\widehat{SE}(\overline{X}_S - \overline{X}_{NS}) = 2.163$. The observed value of the Student's t statistic for testing $H_0: \mu_S = \mu_{NS}$ versus $H_1: \mu_S \neq \mu_{NS}$ is $T_{calc} = -4.13$ with 73 degrees of freedom and a corresponding P-value which is less than .0001. This P-value is quite small indicating that there is very strong evidence that the population mean fecundity for the selected lines μ_S is different from the population mean fecundity μ_{NS} for the nonselected line. The data clearly support the conclusion that the population mean fecundity is higher for the nonselected line, however, technically speaking, we cannot make this conclusion based on the preceding hypothesis test, since we did not have a priori reason to justify a directional hypothesis. We can however form a confidence interval for $\mu_{NS} - \mu_S$ and use it to justify this conclusion. In this example, we are 95% confident that $\mu_{NS} - \mu_S$ is between 4.6192 and 13.241. More precisely we are 95% confident that the population mean fecundity (mean number of eggs laid per day for the first 14 days of life) μ_{NS} for the nonselected line exceeds the population mean fecundity μ_S for the selected lines by at least 4.6192 eggs per day and perhaps as much as 13.241 eggs per day. Thus it appears that the population of fruitflies which are either resistant to or susceptible to DDT has lower fecundity on average than the population of fruitflies which are neither resistant nor susceptible to DDT.

Remark regarding the comparison of the difference of two means to a nonzero constant. In some situations we may have enough a priori information to specify a known constant d with the goal of comparing the difference $\mu_1 - \mu_2$ to this particular constant. For example, we might hypothesize that the first population mean μ_1 exceeds the second population mean μ_2 by more than d = 2 units, *i.e.*, $H_1 : \mu_1 - \mu_2 > 2$ or $H_1 : \mu_1 > \mu_2 + 2$. To test such a hypothesis we simply replace the difference $\overline{X}_1 - \overline{X}_2$ by the quantity $\overline{X}_1 - \overline{X}_2 - d$ in the formula for T and proceed as before. Many computer programs provide an option for testing such a hypothesis.

12.2b Inference when the two standard deviations are not equal toc

In this subsection we will describe an alternate method of inference which can be used when the population standard deviations σ_1 and σ_2 are not equal. Notice that when $\sigma_1 \neq \sigma_2$ the two normal populations are not identical when their population means, μ_1 and μ_2 , are equal. Therefore, a statement regarding the difference between two population means does not tell the whole story about the relationship between the corresponding normal populations when the population standard deviations are not equal. This does not indicate that there is anything wrong with comparing population means when the corresponding population standard deviations are unequal. However, it does indicate that the interpretation of a particular difference between two population means is somewhat different when the population standard deviations are different than it is when the population standard deviations are equal.

When the population standard deviations σ_1 and σ_2 are different, the appropriate estimator of the standard error of $\overline{X}_1 - \overline{X}_2$ is based on the two sample standard deviations S_1 and S_2 rather than the pooled sample standard deviation. That is, when $\sigma_1 \neq \sigma_2$ the appropriate sample standard error of $\overline{X}_1 - \overline{X}_2$ is

$$\widehat{\operatorname{SE}}(\overline{X}_1 - \overline{X}_2) = \sqrt{\frac{S_1^2}{n_1} + \frac{S_2^2}{n_2}},$$

where S_1 is the sample standard error for the sample from the first population (the X_1 values) and S_2 is the sample standard error for the sample from the second population (the X_2 values).

Inference about the relationship between two normal population means when $\sigma_1 \neq \sigma_2$ is based on an approximation to the sampling distribution of the quantity

$$T^* = \frac{\overline{X}_1 - \overline{X}_2}{\sqrt{\frac{S_1^2}{n_1} + \frac{S_2^2}{n_2}}}.$$

Because the details of this approximation are fairly complicated, you really need an appropriate calculator or computer program to implement this method. The test and confidence interval values in the rows labeled Satherthwaite in the SAS two–sample t–test out contain results for this method. 292 12.2b Inference when the two standard deviations are not equal

Using this method the 95% margin of error of $\overline{X}_1 - \overline{X}_2$ is

$$\mathrm{ME}(\overline{X}_1 - \overline{X}_2) = k \sqrt{\frac{S_1^2}{n_1} + \frac{S_2^2}{n_2}},$$

where k is the 97.5 percentile of a Student's t distribution with ν degrees of freedom. The relevant degrees of freedom ν is computed using a complex formula which may yield a value that is not a whole number. An approximate 95% confidence interval for $\mu_1 - \mu_2$ based on this approach is given by the values between $(\overline{X}_1 - \overline{X}_2) - \text{ME}(\overline{X}_1 - \overline{X}_2)$ and $(\overline{X}_1 - \overline{X}_2) + \text{ME}(\overline{X}_1 - \overline{X}_2)$, where the margin of error is as given above. A suitable calculator or computer program will provide the calculated value of this margin of error or the actual 95% confidence interval values.

To test a hypothesis relating μ_1 to μ_2 using this method we simply replace the Student's t statistic T by the approximate Student's t statistic T^* and compute the P-value using the appropriate degrees of freedom ν . A suitable calculator or computer program will provide the calculated value of the approximate t statistic T^*_{calc} and the associated P-value.

One way to determine whether the assumption of a common population standard deviation is reasonable is to compare the results of the confidence intervals and P-values computed assuming equal standard deviations and not assuming equal standard deviations. If the two methods yield essentially the same conclusions, then the assumption of equal standard deviations is reasonable and the methods based on the pooled estimate of the standard error are appropriate; otherwise, the methods which do not use the pooled estimate of the standard error should be used.

12.3 Problems

The problem sets and solutions can be found on the book web page at https://www.ucs.louisiana.edu/~jcb0773/.

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