Statistical Methods for Cardiovascular Researchers

Lem Moyé

Rationale: Biostatistics continues to play an essential role in contemporary cardiovascular investigations, but successful implementation of biostatistical methods can be complex.

Objective: To present the rationale behind statistical applications and to review useful tools for cardiology research.

Methods and Results: Prospective declaration of the research question, clear methodology, and study execution that adheres to the protocol together and serve as the critical foundation of a research endeavor. Both parametric and distribution-free measures of central tendency and dispersion are presented. T testing, ANOVA, and regression analyses are reviewed. Survival analysis, logistic regression, and interim monitoring are also discussed. Finally, common weaknesses in statistical analyses are considered.

Conclusions: Biostatistics can be productively applied to cardiovascular research if investigators (1) develop and rely on a well-written protocol and analysis plan, (2) consult with a biostatistician when necessary, and (3) write results clearly, differentiating confirmatory from exploratory findings. (Circ Res. 2016;118:439-453. DOI: 10.1161/CIRCRESAHA.115.306305.)

Key Words: biostatistics ■ regression analysis ■ research ■ research design ■ survival analysis

The need for biostatistics in cardiovascular research is ever-present, but investigators continue to be challenged by the implementation of statistical methods. A review of the literature reveals both general awareness of this issue1–3 and multiple attempts at its correction.4–9 Aptitude in applied statistics is generated by knowledge and experience, yet few healthcare workers can devote the time required to develop the requisite statistical skill set, a circumstance that generates common statistical mistakes that are fortunately avoidable (Table 1).

The goal here is to provide an overview of biostatistical procedures available to cardiovascular researchers as they conduct their bench or translational investigations. Given the availability of formulae on the internet and in texts, the focus will not be on computation, but on (1) concepts and guidelines in approaches and (2) the strengths and weaknesses of the available tools. In the end, the reader will have reviewed the principles and tools of biostatistics required for preclinical and translational research in cardiology.

General Principles

Nature of Relationships

The mission of cardiovascular research is to discern, quantify, and ultimately classify relationships. The most important connection between the intervention and a response is causal, that is, the intervention or agent is not merely associated in time or place with the response but actually elicits that response. These causal links are critical because they give the investigator control of a relationship that can potentially exacerbate or ameliorate a disease.

However, discriminating between association and causation requires deductive skills and clear epidemiological thinking (Table 2). Several mechanisms are available to investigators that, once embedded in the research design, can improve the researcher’s capacity for discernment. First, researcher determination of the investigative agent’s assignment is the signet of experiments as opposed to observational studies where investigators simply observe relationships in place. Planning the intervention such that the potential for confounding is reduced is essential. Second, blinding the investigator and the subject to treatments ensures that the subjects’ and investigators’ belief systems do not systematically affect one group or the other. Modulating the outcome by adjusting the intervention dose or delivery demonstrates the direct control that the investigator has on the response. Finally, studying multiple end points provides useful information about the similarity of findings. Each of these research traits contributes to the integrity of the study, enabling internal consistency.

Role of Statistics

The internal consistency of a research effort has little impact if the results cannot be applied to a population at large. Generalizability allows the observed effect in a small translational research outcome to be translated into a powerful and beneficial tool that a community of physicians and public health workers can use in vulnerable populations.

The ability to generalize from small collections of subjects to large populations cannot be taken for granted. Instead, it is a property that must be cultivated. In general, 2 features must be present to generalize (1) a prospective analysis plan and (2) the random selection of samples.

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Critical to the generalizability of the research results is the prospective design of a research plan and selection of the population to be sampled. The scientific question must be posed, the methodology set in place, the instruments for analysis chosen, and the end points selected before the samples are obtained. These procedures permit the correct population to be sampled and the sample set to be the appropriate size. Such analyses are termed as prospective analyses because they are conceived before the study is executed.

Of course, these are not the only analyses to be conducted; sometimes the most exciting evaluations are those that cannot be anticipated. Evaluations that were not contemplated during the design of the study are called retrospective, hypothesis generating, or exploratory analyses. The principal difficulty with exploratory analyses is that their absence of a prospective plan decouples the research results from applicability to the larger population. The exploratory question, for example, “Were the research findings different between male and female mice?” is an important question, but one that cannot be effectively addressed if an inadequate number of female mice were sampled. Therefore, with no plans for their execution, hypothesis generating evaluations offer only indirect and unsatisfactory answers to questions that the investigator did not think to ask prospectively. Confirmatory analyses, that is, those that answer prospectively asked questions, are the product of a well-conceived protocol.

Both confirmatory and exploratory analyses can play important roles in sample-based research, as long as the investigator reports each finding with the appropriate perspective and caveats. Well-reported research starts with the prospectively declared research question, followed by the method used to provide the answer and then the answer itself. Only after these findings are presented in the article, should the exploratory evaluations be displayed, perhaps preceded and followed by a note on the inability to generalize their intriguing results.

Random Subject Selection
The second feature required for generalizability is the random selection of subjects. Simple random sampling is the process by which each subject in the sample set has the same probability of being selected from the research sample. This property is required to produce a representative sample of the population to which one wishes to generalize. Appropriate sample selection is key to the researcher’s ability to generalize from the sample to the population from which the sample was obtained. Simple random sampling also adds the feature of statistical independence, that is, a property of observations in which knowledge of the outcome of 1 subject does not inform the investigators of the outcomes for others.

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the null hypothesis is false or (2) a population in which the null hypothesis is true yielded a sample that, through chance alone, suggested that the null hypothesis is wrong. The latter is a sampling error. If the probability of this sampling error is low, then the investigator can reject the null hypothesis with confidence. The probability of this sampling error is the \( P \) value.

**Power and Sample Size**

The preceding section developed the thought process required to manage sampling error in the face of positive results. However, should the sample reveal a result that is consistent with the null hypothesis, there are 2 possible explanations. One is that the null hypothesis is correct. However, a second is that a population in which the finding is positive produced by chance a research sample with a null result. This is considered a type II error. The type II error subtracted from one is the power of a study. Study power is the probability that a population in which the finding should be positive generates a sample that yields the same positive finding. The minimum power for research efforts in cardiology is typically 0.80 (80%). The investigators’ goal is to conduct all hypothesis tests in a high-power environment.

Thus, regardless of the findings of a study, the investigator must manage sampling error. Should the result be positive, then the investigator would need a type I error less than the protocol-declared maximum (traditionally, no \( \geq 0.05 \)). However, should the analysis of the sample lead to a null result, then the investigator should be concerned about a type II error. Essentially, it is beneficial to the investigator if the research is designed to minimize both the type I and the type II errors. This is typically managed by the sample size computation.

There are many sample size formulae. However, each essentially computes the minimum sample size required for the study of a fixed effect, an estimate of variability, and an acceptable level of type I and type II errors. Investigators will be served well by sample sizes that (1) have realistic estimates of variability that are based on the population chosen for study and (2) are based on an effect size that would serve as a finding of importance. In addition, the type I error should be prospectively specified, with the issue of multiplicity considered.

**\( P \) Value**

Although the concept of the \( P \) value is simple, one would be hard pressed to identify another probability that has generated nearly as much controversy. \( P \) values have been lauded as the most effective agent in bringing efficiency to the scientific investigation process, and also derided as—next to atomic weapons—the worst invention of the 20th century. Its difficulty lies in the twin problems of misinterpretation and over-reliance.

History provides an interesting education on the misuse of \( P \) values.\textsuperscript{10,11–24} At times, \( P \) values have been confused with causation.\textsuperscript{25} Perhaps the best advice to researchers is the admonition of Bradford Hill, the founder of modern clinical trials who, when talking about statistical hypothesis testing, said that this tool is “…like fire—an excellent servant and a bad master.”\textsuperscript{26} \( P \) values in and of themselves only provide information on the level of a sampling error that may suggest an alternative explanation of results. The best advice to the investigator is to design the research effort with great care, and to not just rely on the \( P \) value, but also to jointly consider the research design, effect size, SD, and confidence interval when attempting to integrate their research findings into the broader scientific context.

**Multiple Testing**

Research efforts typically generate >1 statistical hypothesis test and, therefore, >1 \( P \) value. Because the \( P \) value reflects the probability of a sampling error event, its repeated generation in the same experiment is analogous to flipping a coin—the more one repeats the experiment (in this case, carrying out additional analyses), the more likely one is to get at least 1 \( P \) value that is below the threshold. When investigators do not correct for this phenomenon and report results as positive simply because the \( P \) value is below the 0.05 threshold, then they are reporting results that are not related to any fundamental property of the population, but that instead misrepresent the population. This produces an unacceptably high false reporting rate.

The overall type I error rate is sometimes designated as the family-wise error rate. It is the probability of at least 1 type I error occurring among the multiple tests that the investigator has conducted, and this probability increases with the number of tests. To combat this probability inflation, the investigators can adjust the maximally acceptable \( P \) value for a positive result (typically set at 0.05) downward. This can be as simple as dividing the \( P \) value by the number of tests that are carried out, as is done in the Bonferroni approximation.\textsuperscript{27} Other procedures are available for the appropriate adjustment.\textsuperscript{28–30}

**One-Sided Testing**

Statistical hypothesis testing, firmly embedded in the conduct of the scientific method, holds an essential and powerful assumption—that the investigator does not know the answer to the question before it is put to the test. The researchers may have an intuition, perhaps even a conviction that they know the result of the study they are designing, but, despite its motivating power, this belief does not supplant data-based knowledge.

Many arguments about 1-sided testing have arisen in the literature, and the attractiveness of these tests is difficult to resist. However, the belief that a study will be beneficial is not an acceptable justification to design a study that will demonstrate benefit only. Experience\textsuperscript{31} has demonstrated that investigators cannot rely on their beliefs about treatment effectiveness when deciding on the sidedness of statistical hypothesis testing. The best solution is to allocate the type I error prospectively, and to use this level in 2 tails, prospectively looking for both benefit and harm.

**Quality Control Procedures**

Low-quality observations and missing data reduce the precision of statistical estimations and degrade the ability of the investigator to accurately answer their chosen scientific question. A periodic review of the data, beginning at the start of data collection and continuing with interim inspections of the incoming data are critical. Identifying mistakes early in the
generation of the data set permits the identification of problems so that the likelihood of error reoccurrence can be reduced (Table 3).

Inspecting the data values themselves is both straightforward and essential. If the data set is too large for the verification of each individual data point, then a random selection of observations can be chosen to compare against the source data, permitting an assessment of data accuracy, and allowing an estimation of the error rate. Examining the minimum and maximum values can alert the investigator to the appearance of physiologically impossible data points. If the researcher is willing to invest time in these straightforward efforts, data correction, published errata, and article retractions can be avoided.

**Missing Data**

Missing data occur for a variety of reasons. The unanticipated difficulty in taking critical measurements, inability to follow subjects for the duration of the study, electronic data storage failure, and dissolution of the research team can each lead to data losses. Fortunately, the effects can be minimized if not completely avoided with pre-emptive action.

One of the most important strategies that the investigator can use is to ensure that subjects (be they human or animal), and that they comprise the subjects who have the disease process of interest, are likely to continue to the end of the study and unlikely to have events that make it difficult to assess the effect of the intervention (eg, competing risks). Use of equipment in the best working order minimizes the likelihood that data cannot be collected. Once collected, data sets should be backed up on secure computer servers in multiple locations, ensuring that critical data sets are not lost because of misplacement, accidental deletion, or natural calamities (eg, floods or widespread power outages). Developing and maintaining solid professional relationships with coinvestigators, as well as establishing and adhering to prospective and clear authorship guidelines, reduces the chances that a disgruntled coworker will abscond with the data.

However, even with adequate protective steps, some data collection may be impossible to complete. In this case, the researchers should report the extent of missing data and the reasons for their absence. Deaths or other losses must be specifically reported to the research community. There are several statistical procedures available to manage this issue. Ad hoc tools that fill in missing values to permit standard software to evaluate complete data (eg, last observations carried forward) should be avoided. This is because (1) the method of data replacement is arbitrary and may be based on fallacious assumptions and (2) their use allows an underrepresentation of the variability of statistical estimates, for example, sample means, proportions, and relative risks. Imputation procedures that permit >1 set of replacements for the missing data, and therefore multiple data sets for analysis, offer improvements over the older observation carried forward approaches. In some circumstances, the occurrence of a significant clinical event (eg, death) precludes the collection of subsequent data. Score functions that are based on the Wilcoxon test or obviate the need to complete missing data that results from this circumstance.

### Table 3. Data Quality Control

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Review and rid the data set of errors before analysis</td>
</tr>
<tr>
<td>2.</td>
<td>Require that core laboratories that generate the data inspect and certify the data before analysis</td>
</tr>
<tr>
<td>3.</td>
<td>Remove data points generated by technically flawed processes</td>
</tr>
<tr>
<td>4.</td>
<td>Do not remove a data point simply because it is inconvenient and not well understood</td>
</tr>
<tr>
<td>5.</td>
<td>Identify and mitigate sources of missing data during the study; report level of missing data and all deaths</td>
</tr>
</tbody>
</table>

### Data Analysis

**Reporting Discrete Data**

Whether the data are discrete or continuous, the investigator must adequately characterize their distribution. This obligation is best met by describing their central tendency and dispersion; how this is carried out, however, depends on the character of the data (Table 4).

Investigators count dichotomous events (eg, deaths). They also categorize discrete events (eg, the distribution of the proportions of patients with different Canadian classifications for heart failure). They can lucidly display dichotomous data by providing the number of events (numerator) and the size of the sample from which the events were drawn (denominator). When faced with polychotomous data, researchers report the frequencies of each of the germane categories, although it can also be appropriate to provide means and SDs.

However, it is also important to measure the effect size (Table 5). The absolute difference between 2 proportions reflects a simple percentage change. It is the simplest reflection of how far apart the percentages lie. Researchers can take advantage of the flexibility of dichotomous end points in the estimation of event rates. For example, if the investigators are interested in assessing the first occurrence of an event, for example, the heart failure hospitalization rate >1 year, then the proportion of such cases is the incidence rate.

If the investigators are interested in comparing the incidence rates of the control group \( I_0 \) and a treatment group \( I_1 \), they can compute the relative risk \( R = I_1 / I_0 \). Although the incidence rate (which is a proportion over time) is between 0 and 1, the relative risk can be any positive number. Because the incidence rate reflects new cases, it is the estimate most sensitive to the effect of a treatment being tested. In cohort studies, in which subjects are followed prospectively over time and background events can be excluded at baseline, incidence rates, and relative risks are the preferred estimates. In this case, these estimates could be used to determine the expected number of

### Table 4. Central Tendency and Dispersion Estimates

<table>
<thead>
<tr>
<th>Type of Data</th>
<th>Central Tendency</th>
<th>Dispersion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dichotomous</td>
<td>Proportion</td>
<td>Frequency</td>
</tr>
<tr>
<td>Continuous</td>
<td>Mean, median, mode</td>
<td>Interquartile range</td>
</tr>
</tbody>
</table>

For dichotomous data

Report: Central Tendency: Proportion of subjects

For polychotomous data

Report: Central Tendency: mean, median, and mode

Dispersion: Display actual frequency distribution, SD, and range

For continuous data

Report: Central tendency: Mean, median, and mode

Dispersion: Interquartile range, SD, and confidence interval for the mean
individuals that would need to be treated to prevent 1 event. Odds ratios are useful for comparing differences in prevalence.

**Reporting Continuous Data—Normal or Distribution-Free Approaches**

Reporting continuous data requires the joint consideration of central tendency and dispersion. Continuous data that are skewed or flat are not adequately summarized by the sample mean and SD, and when reported in this fashion, they are commonly misleading. In this case, investigators are advised to use procedures that are not reliant on the underlying probability distribution. Reporting continuous data requires the joint consideration of central tendency and dispersion. Continuous data that are skewed or flat are not adequately summarized by the sample mean and SD, and when reported in this fashion, they are commonly misleading. In this case, investigators are advised to use procedures that are not reliant on the underlying probability distribution.37 These distribution-free (also termed nonparametric) procedures are based on percentiles of the data that describe the distribution’s central tendency (eg, the median) and dispersion (eg, the minimum, 25th percentile, 75th percentile, and maximum). Graphical representations are helpful in presenting continuous data; chief among these is the box plot (Figure 1).38,39

The Shapiro–Wilks test40 is useful to assess whether data are normally distributed, and the Kolmogorov–Smirnov test, although prone to low power, permits the investigator to compare the probability distribution of the sample data with that of a known probability distribution41 or that of another sample set. The best guide is to observe the data and select an approach based on their distribution. Should the community of researchers expect an approach that is different than that suggested by the data, then, in addition to the procedure that the investigators think the data requires, the investigators can take the approach suggested by the community and compare results from the 2 analyses.

When the investigators think that reporting means and their measures of dispersion are appropriate, they should consider the choice of SD versus the SE. The SE is simply the SD of a sample mean, and it is smaller than the SD. Which of these measures should be reported depends on the circumstances. If the investigators are interested in reporting the location and dispersion of individual observations in their sample set, then using the SD permits the reader to assess the variability among observations. However, if the purpose is to show the value and variability not of the observations, but of the means, then the SE should be used. However, once the investigators choose one of these measures of variability, they should be clear as to which they are providing.

**Hypothesis Testing Tools**

**Hypothesis Testing for Dichotomous and Polychotomous Data**

A list of statistical tests to assess common statistical hypotheses is available in Table 6. Investigators can test the null hypothesis for dichotomous data using Fisher exact test.

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**Table 5. Effect Size Measurements for Dichotomous Data**

<table>
<thead>
<tr>
<th>Example: 18 of 25 mice die post infarction in the control group, whereas 10 of 30 die in the gene therapy group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Absolute differences: $p_2 - p_1 = 18/25 - 10/30 = 0.720 - 0.334 = 0.386$</td>
</tr>
<tr>
<td>Relative risk: $p_2 / p_1 = 0.334/0.720 = 0.46$</td>
</tr>
<tr>
<td>Odds ratio: $p_2 / (1-p_2) / (p_1 / (1-p_1)) = (0.334/0.666)/(0.720/0.280) = 0.195$</td>
</tr>
<tr>
<td>Percent reduction (efficacy): $(p_1 - p_2) / p_1 = 0.720 - 0.334/0.720 = 0.386/0.720 = 0.54$</td>
</tr>
<tr>
<td>Number needed to treat: $1/(p_1 - p_2) = 1/0.720 - 0.334 = 1/0.386 = 2.59$</td>
</tr>
</tbody>
</table>

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**Figure 1. Example of a box plot.**

### Distribution-Free Hypothesis Testing

#### Results—Continuous Data
Many distribution-free tests are available to investigators (Figure 2). When investigators examine their data and find that the distribution-free approach is best, they can conduct a 1-sample assessment using the 1-sample Wilcoxon signed-rank test that determines the population median based on the sample. The investigators should anticipate the median value for the population under the null hypothesis. This test requires all the data, not just the sample size, mean, and variance. Its output is the sample median and \( P \) value.

Two-sample testing can be conducted using the 2-sample Wilcoxon signed-rank test if there is a natural pairing of the data. As with the 1-sample test, all of the data are required, as the data are ordered and the ranks of each pair of data points are compared. The test addresses whether the medians of the 2 populations are different, and the output is the 2 medians and the \( P \) value. The investigators can also conduct a 2-sample Wilcoxon rank sum test (distribution-free) when the data consists of 2 unpaired samples. In this case, each of the observations from 1 sample is compared to each of the observations from the other sample.

Examples of these data include the Canadian Classification Score for Angina or the New York Association Heart Failure Score. In these cases, they can use Fisher exact test as well because it is more accurate than \( \chi^2 \) testing, particularly when there is a small number of events in some variable categories.

### Table 6. Statistical Methods

<table>
<thead>
<tr>
<th>Question to be Answered</th>
<th>Procedure</th>
<th>Advises on Application</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-sample testing</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Normality assumption valid</td>
<td>1-sample ( t ) test</td>
<td>Requires the sample size, mean, and SD</td>
</tr>
<tr>
<td>Is the population mean different than expected?</td>
<td>1-sample Wilcoxon-signed rank sum test</td>
<td>Requires the entire sample</td>
</tr>
<tr>
<td>Distribution free</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Is the median of this distribution different than expected?</td>
<td>1-sample Wilcoxon-signed rank sum test</td>
<td>Requires the entire sample</td>
</tr>
<tr>
<td>2-Sample testing</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Normality assumption valid</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Do the means of 2 paired samples differ?</td>
<td>Paired ( t ) test</td>
<td>Test variance first. Use equivaraint solution for test on means if variances equal. Use unequal variance solution otherwise</td>
</tr>
<tr>
<td>Do the 2 independent samples have different means?</td>
<td>Unpaired ( t ) test</td>
<td></td>
</tr>
<tr>
<td>Distribution free</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Are the elements of 2 sequences mutually independent?</td>
<td>Spearman Rank Correlation Test</td>
<td>Requires entire data set; computes correlation based on ranks</td>
</tr>
<tr>
<td>Are the paired samples from populations with equal medians?</td>
<td>Two sample signed-rank test</td>
<td>A paired difference test requiring all of the data</td>
</tr>
<tr>
<td>Are the unpaired samples from populations with equal medians?</td>
<td>Wilcoxon Rank Sum test (Mann–Whitney ( U ) test)</td>
<td>Requires entire data set; computes a sum of signed ranks of the observations</td>
</tr>
<tr>
<td>&gt;2 Groups</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Normality assumption valid</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Do the predictor variables explain the response variable?</td>
<td>Regression analysis</td>
<td>Requires entire data set; adjusts for confounders; chunk testing is available; provides effect sizes and ( P ) values</td>
</tr>
<tr>
<td>Are there differences among the treatment group means?</td>
<td>ANOVA</td>
<td>Requires entire data set; adjusts for confounders; blocking can be used; provides effect sizes and ( P ) values</td>
</tr>
<tr>
<td>Are their differences between different groups with multiple response variables?</td>
<td>Multivariate analysis of variance</td>
<td>Requires entire data set; produces table related to ANOVA table; effects for individual response variable must be analyzed separately</td>
</tr>
<tr>
<td>Distribution free</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Are samples from each group from the same distribution?</td>
<td>Kruskal–Wallis test</td>
<td>Requires entire data set; provides results analogous to a 1-way ANOVA</td>
</tr>
<tr>
<td>Survival analysis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Is there a difference in the time to event between the groups?</td>
<td>Kaplan–Meier (log rank)</td>
<td>Requires for each subject the censoring category and time to event or end of follow-up; can manage more than 2 groups</td>
</tr>
<tr>
<td>Can predictors explain the difference in time to event?</td>
<td>Cox proportional hazard analysis</td>
<td>Requires the entire data set; provides effect sizes as well as ( P ) values. Adjusts for confounders</td>
</tr>
</tbody>
</table>
follow is to (1) transform the data, (2) carry out an analysis using parametric procedures, and then (3) apply the inverse transforms to convert the estimates of the mean and confidence intervals into the data with the original distribution.

Continuous Data Under a Normal Distribution

The investigators can avail themselves of several statistical tools when assessing the impact of an intervention on a continuous response variable that follows a normal distribution (Figure 3). When reporting data that is normally distributed, the investigators should report the effect size, the SE (which is the SD of the effect size), and the 95% confidence interval. The most useful statistical test for the investigator is the $t$ test. There are essentially 3 types of $t$ tests: (1) 1-sample, (2) 2-sample, paired, and (3) 2-sample, unpaired. The latter can be conducted when either the variances of the 2 groups are considered equal or are believed to be unequal. In each case, all that is required is the sample size, mean, and SD of the measures of interest from each group.

If investigators have response measures from $n$ subjects and wish to test whether the population from which they selected their sample has a particular mean value, they would conduct a 1-sample test based on the mean response variable value $X$ and the SD $S$. The output of this analysis is the mean, 95% confidence interval for the population mean, and $P$ value.

If investigators have an interest in tracking changes in a response variable over time, then each subject will have a measure at baseline $X_b$ and a measure at follow-up $X_f$. To assess the change in response, the investigators can conduct a paired $t$ test because each subject has a baseline and follow-up measurement, with the baseline measurement providing information about the follow-up measurement. This information provided by one measure about the other permits a smaller estimate of the variance, and therefore a more powerful hypothesis test. Here, the output is the sample means and SDs for each of the 2 groups, the 95% confidence interval for the difference in means, and the $P$ value.

If investigators have 2 groups of subjects that differ in a particular characteristic and they are interested in testing whether the means of the 2 populations are different, then the 2-sample $t$ test would serve the researchers well. As before, the output of the analysis is the sample means and SDs for each of the 2 groups, the 95% confidence interval for the difference in means, and the $P$ value. However, the exact computation for the $P$ value depends on whether the variances in the 2 samples can be considered equal. Typically, statistical
software provides the results for both the scenario where the investigators assume the variances are equal (and can therefore be pooled across the 2 groups) and also when the variances are assumed to be different. A separate hypothesis test can be conducted to assess the equality of variance assumption.

There is sometimes confusion about the best analysis to perform in a study where there is a control and a response group and each subject has a baseline and follow-up measurement. It is common for investigators to use 2 paired t tests to determine if (1) the change in the response of the control group is statistically significant and (2) the change in the response of the active group is statistically significant; however, this approach is suboptimal. A better approach to use is to compute for each group the mean change from baseline to follow-up and SE, and then to compare these mean changes using a 2-sample unpaired t test. However, investigators are well advised to avoid t testing when the sample size of each group is extremely small (eg, n=3) and effect sizes are anticipated to be small or moderate. In this case, the researchers are better off with a 1-sample sign test or 2-sample Wilcoxon rank sum test. In addition, the investigators should keep in mind that one of the principal reasons to conduct hypothesis testing is to reliably determine whether results can be generalized from the sample to the larger population, a reliability that is undermined by the small size of the sample.

Analysis of Variance

ANOVA and regression modeling focus on building relationships between the response variable of interest and a collection of predictor variables (Figure 4). Investigators can use these procedures to assess whether the values of predictor variables change the response variable’s mean.

The fundamental concept behind ANOVA is that subjects treated differently will differ more than subjects treated the same. Even though it is the arithmetic means of the response variables that are examined across the predictor variables, the analysis is principally focused on converting unexplained variability to explained variability. If a variable is completely unexplained, then the investigator does not know why one subject’s measure is different than that of another. Identifying predictor variables provides an explanation for these differences, thereby reducing the unexplained variability. The inclusion of each new predictor variable increases the percent of the total variability that is explained, an increase that is reflected in an increased $R^2$.

Researchers can randomize subjects to one type of variable but with multiple levels (eg, a single variable diet that comprises 3 different and distinct diets). They are interested in assessing whether there are any differences among these levels of the response variable. Because there are $>2$ groups in this type of analysis, a single t test is inappropriate. Although the investigators could conduct several different t tests, the multiple testing issue must be addressed to avoid type I error inflation. Alternatively, an ANOVA test could be conducted to assess the degree to which the supplements explain differences in the rats’ left ventricular ejection fraction (LVEF) measurements. Because supplements are essentially the same class of predictor variable, the analysis is a 1-way ANOVA. The F-statistic provides the overall test of significance.

Having conducted the ANOVA, investigators are commonly interested in assessing which group had the greatest effect on the response variable. This is the multiple testing problem, and the investigators are faced with comparing pairs of interventions (commonly referred to as contrasts). Dunnett test is useful when the investigator is interested in testing several different interventions against a single control, and it is relatively easy to implement. Scheffé method constructs confidence intervals simultaneously for multiple hypothesis tests, and it is helpful in ANOVA. Tukey test permits the

![Flowchart of parametric analyses for ANOVA.](http://circres.ahajournals.org/)}
investigator to identify differences in means that have passed statistical testing, while correcting for multiple comparisons. These are important tools available to the investigator that should be considered during the design of a research protocol.

A distribution-free analog to the 1-way ANOVA that is available to investigators is the Kruskal–Wallis procedure. It can be used to compare the distributions of ≥2 samples in a 1-way ANOVA.

Factorial Designs
The ANOVA model just described provides a structured set of procedures to determine if a set of >2 conditions has an effect on the response variable. However, research designs are increasingly complex, allowing deeper analyses of the relationship between a response variable and a predictor variable. In particular, factorial designs and blocking permit the investigator to examine the relationship between multiple predictor variables and the response variable, and at the same time decrease the response variable’s unknown variability. For this procedure, data on the response variable and each of the predictor variables must be available at the subject level.

A factorial design examines the impact of multiple dichotomous or polychotomous explainer variables on the response variable. In this case, the investigator is interested in each relationship. Consider an example where an investigator is interested in assessing the effects of 2 factors on LVEF recovery post infarction. In each pig, the researcher assesses the change in LVEF over time. One factor is the provision of granulocyte colony-stimulating factor (G-CSF) 1 week before the beginning of the infarction. The second factor is the presence of mesenchymal stem cells (MSCs), which are delivered immediately post infarction. The investigators randomly choose whether the pigs receive G-CSF alone, MSC alone, both treatments or a placebo. This 2-by-2 design permits an assessment of the overall effect of each of the G-CSF and MSC effects. It also evaluates whether the effect of MSC cells will be more pronounced in the presence of G-CSF than in its absence. This is traditionally known as an interaction, but is more helpfully described as an effect modifier, that is, the effect of MSC is modified by the presence of G-CSF. The estimates of the main effects of G-CSF and MSC, as well as an effect modification, give this design a unique name: factorial design. Three and higher dimension factorial designs are also possible. From a variability reduction perspective, this is an efficient design. The investigators can assess the influences of G-CSF and MSC, both separately or together, to reduce the unexplained variability in LVEF.

Blocking Designs
Blocking designs, as with factorial designs, evaluate the relationship between multiple dichotomous or polychotomous explainer variables on the response variable. However, unlike the factorial design, where the researcher is interested in each relationship, in the blocking design, the researcher is only interested in the relationship between some of the predictor variables and the response variable. Variables that have a relationship with the response variables, but in which the investigator has little interest, are used to explain (or block) and eliminate variability in the response variable.

Using the previous example as a foundation, consider now the investigator who is interested in assessing the effects of G-CSF on the change in LVEF. Here, either G-CSF or placebo is randomly provided to pigs immediately post infarction. However, the second factor for this experiment is not MSC, but is instead the size of the myocardial infarction.

The similarity of this design (G-CSF and myocardial infarction size as explainer variables) with the previous example (G-CSF and MSC as the explainer variables) is both helpful and misleading. Each design has 2 dichotomous explainer variables, and contains data in each of 4 cells. Therefore, mathematically, they can be analyzed the same way. However, there is a major difference. In the factorial design, each of the factors, G-CSF and MSC, were assigned randomly and the investigators were interested in the effects of each. In the second design, only G-CSF was randomly allocated. Infarct size could not be predetermined and assigned, and could very well be a surrogate for other variables, (eg, condition of the LV before the myocardial infarction occurred or factor precipitating the myocardial infarction). In fact, the investigators are not particularly interested in assessing the effect of myocardial infarction on \(\Delta \text{LVEF}\).

However, this established relationship between infarct size and \(\Delta \text{LVEF}\) is used to the investigator’s advantage. Because the infarct size–\(\Delta \text{LVEF}\) relationship is well understood, the investigators know that myocardial infarct size will substantially reduce the unknown variability of \(\Delta \text{LVEF}\). This reduction in unknown variability is akin to reducing the surrounding noise, a procedure that will make it easier to detect the relationship of G-CSF to \(\Delta \text{LVEF}\). In this case, the infarct size is a blocking variable, that is, it is of no real interest itself, but it reduces the background noise, making it easy to detect the G-CSF–\(\Delta \text{LVEF}\) signal.

Repeated Measures Testing
The repeated measures design refers to research in which subjects have multiple measurements over time. These designs are useful when investigators are interested in assessing the effect of time on the distribution of observations. The simplest repeated measures analysis is a paired \(t\) test, in which there are 2 measurements per subject. In these designs, time is considered to be the within subjects variable. These designs can be expanded to have observations at many time points and simultaneously contain “between subjects” measurements (eg, control versus treatment). Crossover designs are adaptations of repeated measure designs.

An efficiency introduced by the repeated measures design is the ability to directly assess the relationship between the response variables over time. Although the researcher’s knowledge of the first subject’s baseline response measurement tells them nothing about the next subject’s measurement (the sine qua non of independence), knowing subject one’s baseline response can inform the investigator about that individual’s responses in the future. If these responses are linearly dependent (ie, the fact that subject one’s baseline response is above the average baseline level makes it more likely that subject one’s response in the future will be above average) then a positive correlation is present. Just as in the ANOVA discussed above, the \(F\)-statistic determines the statistical significance of the time effect.
However, it is important for the investigators to recognize 2 critical assumptions in classical repeated measures design. The first is that this evaluation assumes an underlying normal distribution for the response variable. If the response variable is not normally distributed, or is not linear over time, then researchers can consider transformations of the data. In addition, investigators can carry out the Friedman test to assess differences over time when the response variables are ordinal, or when they are continuous but violate the normality assumption.

A second assumption in classical repeated measures designs is the compound symmetry assumption (ie, there is equal correlation between the data points over all times). Modern computing programs permit the investigator to use the precise correlation structure in the data set to be evaluated. In addition, the researcher can adjust the overall degrees of freedom used in the evaluations with the Greenhouse–Geisser, Huynh–Feldt, and lower bound procedures.

Multivariate Analysis of Variance

Investigators can turn to multivariate analyses when they are interested in assessing the effects of multiple predictor variables on the differences in means of multiple response variables. This procedure uses the correlation between multiple response variables to reduce the unexplained variability. Essentially, the multiple response variables are converted into 1 multidimensional response variable, and the statistical analysis assesses the effects of the predictor variables on this response variable vector.

For example, if the investigators are interested in assessing the effect of Ckit+ cells, MSC cells, and the combination of these 2 cell types on LVEF post myocardial infarction, they could assess these cell effects by ANOVA. However, deciding on a single response variable that determines LV function can be a challenge. Alternatively, they could measure LVEF, LVESVI (left ventricular end systolic volume index), LVEDVI (left ventricular end diastolic volume index), and infarct size as response variables and conduct 1 multivariate analysis of variance for these 4 variables.

The multivariate analysis generates an assessment of the relationship between the 2 variables (Table 7). If the R² from this analysis is a sufficient process, producing 1 P value for all of these evaluations, and it is most useful when the result is null. If the investigators conducted an ANOVA on each of the 4 response variables, the multiplicity problem would be daunting (4 hypothesis tests per ANOVA and 4 ANOVAs). However with multivariate analysis of variance, should at least 1 relationship be statistically significant, the investigator must conduct a series of contrasts to determine where in the system the significant relationship lies. Also, like ANOVA, multivariate analysis of variance hypothesis testing is based on the underlying normality of the individual response variables. When the observations do not follow a normal distribution, the investigator can transform the data to normality. If this is not possible, then permutation tests can be useful.

### Regression Analyses and Adjusting for Influences in Normal Data

Relationship building plays a pivotal role in understanding the true nature of the relationship between the response variable and the predictor variable. As pointed out earlier, ANOVA is useful when the response variable is continuous and the predictor variables are dichotomous or polychotomous in character. However, commonly, the predictor variables are continuous as well. In this case, the investigator can carry out regression analyses.

These analyses require the entire data set. When there is only 1 predictor variable, the regression analysis assesses the strength of the association between the response and the predictor variable, and the R² value indicates the percent of the total variability of the response variable that is explained by the predictor variable. When >1 predictor variable is included in the regression model, the analysis indicates the strength of the association between the response and each of the predictor variables, adjusted for the effect of the other predictor variables. P values are produced for each variable in the regression analysis.

As an example, suppose that a team of investigators is examining the change in LVEF over time in human subjects with heart failure and ongoing ischemia. These patients had their baseline CD34 phenotypes assessed. The investigators are interested in understanding the relationship between the response variable ∆LVEF and the predictor variable CD34. A simple regression analysis demonstrates the strength of the relationship between the 2 variables (Table 7). If the R² from this

<table>
<thead>
<tr>
<th>Variable</th>
<th>DF</th>
<th>Parameter Estimate</th>
<th>SE</th>
<th>t Value</th>
<th>Pr &gt;</th>
<th>R²</th>
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<td>0.50789</td>
<td>2.14</td>
<td>0.0356</td>
<td></td>
</tr>
</tbody>
</table>

LVEF indicates left ventricular ejection fraction.
model is 7.9%, then this indicates that 7.9% of the total variability of ΔLVEF is because of CD34, leaving most of the variability (100×(1−0.079)=92.1%) in this simple model to be is explained by factors other than the baseline CD34 phenotype.

Using this same table, the investigators see that there is a quantitative relationship between ΔLVEF and CD34. It can be written as follows:

\[ ΔLVEF = 1.28 \times CD34 - 3.28 \]

They can reject the null hypothesis that there is no relationship between CD34 and ΔLVEF, and can explain the effect size from this model as follows: if there are 2 patients, and one has a CD34 level one unit higher than the other, then, on average, the ΔLVEF of the patient with the higher CD34 level will be 1.28 absolute LVEF units greater than the other. The \( P \) value associated with this relationship is 0.012. Thus, the investigators would conclude that (1) there is a statistically significant relationship between ΔLVEF and CD34 and (2) the relationship still leaves most of the variability of ΔLVEF unexplained.

Now suppose that the investigator expected that ΔLVEF might also be explained by age and analyzes this new variable. The result of that analysis shows an \( R^2=6.01\% \), coefficient=-0.127, and \( P=0.027 \). From this analysis, the investigators have learned that just as there was a statistically significant relationship between ΔLVEF and CD34, there is also a statistically significant relationship between ΔLVEF and age. In the latter case, the relationship is \( ΔLVEF = -0.127 \times \text{age} + 8.53 \).

These results suggest that older subjects having a lower ΔLVEF level. However, identifying these 2 relationships begs the question: Of the 2 variables, CD34 and age, which is more closely related to ΔLVEF? This requires the research team to be familiar with the concept of statistical adjustment.

**Adjustment**

The notion of adjustment is different for researchers. To some, adjustment is synonymous with control, or in this circumstance, keeping age constant to assess the effect of baseline CD34 levels on ΔLVEF. This would be achieved by (1) choosing an age and (2) carrying out a regression analysis relating ΔLVEF to CD34 for individuals only of that age. This is difficult to carry out, of course, because there are typically not enough individuals with the same age (or in an age stratum) that would permit a useful assessment of the ΔLVEF–CD34 relationship.

To biostatisticians, adjustment means something different then control. It means identifying, isolating, and then removing the influence of the adjusted variable on the relationship between the 2 other variables. This is the process of controlling for confounding, or understanding the way in which 2 predictor variables are correlated and, therefore, confound or conflate the relationship that each of them has with LVEF.

Regression analysis achieves this adjustment by (1) identifying and isolating the relationship between ΔLVEF and age, removing this effect from ΔLVEF, then (2) identifying the relationship between age and CD34 and removing this effect from CD34, and (3) evaluating the relationship between what is left of the CD34 effect on ΔLVEF completes the adjusted analysis. This process of isolating, identifying, and removing effects is precisely what multiple regression analysis achieves. Specifically, the result of this complex sequence of evaluations is the output of a regression analysis that relates ΔLVEF to both CD34 and age when they are both in the same model.

From the table, with both variables in the model, the total variability has increased to 11.2%, which is greater than that of either CD34 by itself or age by itself. The overall model keeps the same directionality of the relationship, that is, ΔLVEF is decreased by age, and it is larger in subjects with larger CD34 measures. However, the significance of the CD34 effect remains after the adjustment (i.e., after identifying, isolating, and removing the effect of age). Note, however, that the statistical significance of the ΔLVEF–age relationship is lost after adjusting for CD34. Thus, the age-adjusted CD34 relationship with ΔLVEF remains significant, whereas the CD34–adjusted age relationship with ΔLVEF does not.

Investigators should be aware of some cautions concerning regression analyses. With small data sets, the regression model is likely to be overspecified, providing so close a fit to the data set that the results are of little value for the population at large. In this circumstance, partitioning the data into a data set from which the model was created, and a second (and sometimes third) data set on which the model is tested increases the validity of the model.

Statistical software is adept at building complex models that can explain a substantial proportion of the variability in dependent variable values. However, although a small number of predictor variables may have important impacts on the dependent variable, other predictor variables that are in the model, while also having a measurable impact on the dependent variable, are less influential. Therefore, when investigators face a model with a substantial number of predictor variables, they may be able to simplify the message conveyed to the research community by focusing on the 2 or 3 predictor variables that have the greatest impact.

An alternative approach to regression model building is to test an ensemble or group of effects for several predictor variables simultaneously. In this case, the investigators consider a model with and without this group of predictor variables. These chunk tests are straightforward, but require the investigators to know how to bundle subsets of predictor variables so that group-variable effects can be more easily interpreted.

**Survival Analysis and Logistic Regression**

Survival analysis with Cox regression and logistic regression are regression procedures in which the response variable is a dichotomous variable or the combination of a response variable and a continuous variable (time to event).

Researchers can take advantage of the flexibility of dichotomous end points in the estimation of event rates. For example, if the investigators are interested in assessing the first occurrence of an event, for example, the heart failure hospitalization rate \( \geq 1 \) year, then the proportion of such cases is the incidence rate discussed earlier.

Investigators use survival analyses when they want to examine and compare the survival of groups of individuals...
Survival is the time to an event (e.g., time to first hospitalization or time to death). The quantity of interest is the average time to event, and this is commonly compared between a control group and treatment group. It is a straightforward process when the investigators know the survival time of each individual. However, this knowledge is frequently unavailable for all subjects. Kaplan–Meier life tables, log rank testing, and Cox regression analyses are most helpful to investigators in this case.

In analyses where subjects are followed for a prescribed period, investigator knowledge of the fate of each subject can be limited, and this limitation has an impact on the analysis. In a mortality study, for example, if a subject dies during the trial and the death date is known, then the investigators can use the survival time of that individual in their analysis; this subject is uncensored and the duration of survival is known. However, subjects may disappear from the study but not necessarily be dead. In these cases, the researcher only knows when the subject was last seen in the study, but does not know their current event status after they removed themselves from the research (some of the more eyebrow-raising reasons why subjects do not finish studies are that they are fleeing the law, have left their spouse for a surreptitious partner, are active traders in illegal narcotics businesses, are in prison, or are members of an organized crime network). All that is known is that the individual survived up to the time of last contact. This survival is considered censored. These subjects are removed from follow-up after their last visit, and their removal is censored. Finally, subjects can reach the end of the study without an event occurring at all, denoted as administrative censoring. These different occurrences can all affect the estimates of mean survival times in the study.

Kaplan–Meier estimates rank the data by survival time in the study, and, then, using the censoring mechanisms, compute the number of subjects at risk of death in a time period, and, from that, determine the proportion of those at risk of death who actually die. This is assembled for the duration of the study to compute estimates of the overall survival rates. Cox hazard regression permits the investigator to conduct regression analyses on time to event/censored data. Here, there are 2 response variables: (1) whether the subject’s final status is censored or uncensored and (2) the individual’s follow-up time in the study. Predictor variables were reviewed above. Cox regression produces as its output the relative risk associated with the predictor variable (through an exponentiation of the product of the coefficient and the predictor variable) and the P value associated with that relative risk.

Alternatively, proportions can be used to estimate prevalence, for example, the proportion of the population that has ever been hospitalized for heart failure. In retrospective studies, where the investigator starts with the number of cases and cannot differentiate incident cases from those that have been present in the population for a while, the investigator must rely on prevalence measures. If the investigators have only the prevalences for the control and intervention groups, \( p \) and \( p_c \), they can compute the prevalence ratio \( \frac{p}{p_c} \). Although this is related to the incidence ratio, it can be difficult for an intervention to have an impact on this ratio if the background rate (which is not affected by the intervention) is large.

In an assessment of prevalence, investigators can compute a more helpful quantity, the odds, which, for which the control group would be \( O = \frac{p}{1−p_c} \). This can take any value >0, and with this value, it becomes easier to think of a regression model that would be based on it. To compare 2 groups, the odds ratio \( \frac{O_1}{O_2} = \frac{(p_1/1−p_1)}{(p_2/1−p_2)} \) can be used. Odds ratios greater than one indicate that the treatment is more closely associated with the event than the control, and odds ratios <1 indicate that the odds of the event are lower for the treatment than for the control group.

Another advantage of odds ratios is that investigators can produce them from logistic regression analyses. The response variable in a logistic regression is most commonly a dichotomous variable. The predictor variables can be either continuous or discrete. In addition, as was the case for regression analysis, logistic regression can accommodate multiple predictor variables. The exponentiated coefficient of a dichotomous predictor variable from a logistic regression analysis is an odds ratio. This procedure, therefore, permits the investigators to model the odds ratio as a function of covariates, and produces analysis tables that would be familiar to investigators who understand how to interpret this value from a regression analysis.

### Combined End Points

An adaptation available to investigators in clinical trials is a combined end point. In this case, the investigators combine multiple end points into 1 single but complex event. Researchers will face complications in the design of these end point structures, so the design must be based on an appropriate rationale.

There are advantages and disadvantages to this approach. When each component end point of the combined end point is dichotomous, the combined end point has a higher frequency of occurrence that will decrease the trial’s sample size. This can also decrease the cost of the trial. However, there are complications that come with the combined end point. If the combined end point has a nonfatal component, data collection can present a new administrative burden because it now requires an assessment to determine if the nonfatal component for any subject (e.g., hospitalization for heart failure) meets the study criterion. This increases the cost and complexity of the study.

In addition, the use of combined end points can complicate the interpretation. The most straightforward conclusion to the study would be if all components of the combined end point are influenced in the same direction by the intervention. Any discordance in the findings of the component end points vitiates the impact of the intervention and undermines any argument that the treatment was effective.

However, even with these difficulties, combined end points have become an important adaptation in clinical trials. They are most useful if 4 principals are followed (Table 8). Both the combined end point and each of its component end points must be clinically relevant and prospectively specified in detail. Each component of the combined end point must be carefully chosen to add coherence to the combined end point, and should be measured with the same scrupulous attention to detail. The analysis of the therapy’s effect on the combined end point should be accompanied by a tabulation of the effects of the intervention on each component end point.

In addition, investigators should avoid the temptation to alter the composite end point and its prospectively declared type I error
Table 8. Principles of Combined End Points

| Principle of Prospective Deployment: Both the combined end point and each of its component end points must be clinically relevant and prospectively specified in detail |
| Principle of Coherence: Each component of the combined end point must be carefully chosen to add coherence to the combined end point |
| Principle of Precision: Each of the component end points must be measured with the same scrupulous attention to detail |
| Principle of Full Disclosure: The analysis of the effect of therapy on the combined end point should be accompanied by a tabulation of the effect of the therapy for each of the component end points |

structure during the course of the study. This tempting midcourse correction can lead to studies that are difficult to interpret. In addition, changing components of the combined end point based on an interim result is often unsuccessful because such interim findings are commonly driven by sampling error. Therefore, investigators should adhere closely to the prospective analysis plan.

Subgroup Analyses

A subgroup analysis is an evaluation of a treatment effect in a fraction of the subjects in the overall study. These subjects share a particular characteristic (eg, only males, or only patients with phenotype CD34 levels below a specific value). There are several justifiable reasons for investigators to conduct subgroup analyses. However, they should know that these analyses are also fraught with difficulty in interpretation.

Commonly, the investigator simply and innocently wishes to determine whether the effect seen in the overall study applies to a specific demographic or comorbid group. This search for homogeneity of effect can be instigated by the investigators of the study. This evaluation can also be requested by journal reviewers and editors, or the entity that sponsors the research. Even the US Food and Drug Administration conducts subgroup analyses on data presented to it. However, investigators should adhere closely to the prospective analysis plan.

First, if a study is to be ended prematurely, the data must be sufficiently compelling to warrant an early termination. Second, the early conclusion will be most persuasive if all patients who could take the medicine and respond are compared with those who did not. Not even the prospective declaration of an “as treated” analysis can cure this problem. The third, lack of adequate power, and fourth, P value multiplicity, are statistical, and undermine the generalizability of these within-subgroup effects.

Subgroups analyses have received substantial attention in clinical trial methodologies. However, investigators are best advised to treat provocative subgroup findings like “fool’s gold.” Caveat emptor.

Interim Monitoring and Other Adaptive Procedures

Up until this point in the review, the requirement for prospective planning in research has been emphasized. The intervention to be studied, the characteristics of the study population, the number of study arms, the number of end points, and the sample size of the study should all be identified prospectively. Then, once established, the protocol should be followed in detail. The advantage of this procedure is that the results of the study are more likely to be generalizable than with ad hoc analyses.

However, even though this ability to apply the study’s results to the population at large is ideal, it is also inflexible. Might there not be some advantage to other approaches to research? The need for protocol resilience is clearest when there is an ethical reason to end the study early because of early therapeutic triumph or early catastrophe. Because investigators carrying out the trial must remain blind to therapy assignment, a special oversight board with access to unblinded data (the Data Safety and Monitoring Board) is charged with reviewing the data well in advance of the completion of subject follow-up.

In so doing, these groups are required to consider action based on compelling but incomplete data. Several quantitative tools, called group sequential procedures (procedures that examine data from groups of subjects sequentially), were specifically designed to assist in these complex determinations. Although the mathematics are somewhat complicated here, the concepts are simple.

First, if a study is to be ended prematurely, the data must be more persuasive than the minimal acceptable efficacy, had the trial been permitted to proceed to its conclusion. Second, the earlier the study is stopped, the more persuasive these findings must be. In addition, because data are examined repeatedly (although more data are added for each interim examination), there is a price one pays for multiple examinations of the data (and multiple P values). Although this price is modest, an adjustment must be made.

However, perhaps the most important part of these interim examinations is nonstatistical. Issues such as internal consistency (do primary and secondary end point align?), external consistency (have these results been seen in other studies?), biological plausibility, and data coherence must also show that the interim data support a unified conclusion. All these factors must be considered together for the decision to prematurely end a study.

Now, it is not uncommon for the Data Safety and Monitoring Board to examine not just end point data for signs of early efficacy or early harm, but to examine the event rate
in the placebo group of a clinical trial to recommend either increasing or decreasing the follow-up period based on the observed rate. If the end point is continuous, then its SD can be appraised in real time. Each of these procedures is designed to test the assumptions of the underlying sample size computation, adapting the computation to the actual data. Multi-armed clinical trials, for example, studies designed to evaluate the effects of multiple doses of the agent in question, can have the sample sizes in specific arms increased or decreased, depending on the effect seen in that treatment arm. Each of these possibilities must be described in detail in the protocol, but once specified, they can be implemented without prejudice to the trial. The Food and Drug Administration has recently produced a draft guideline on adaptive clinical trial design.62

The principal motivation for these rules is the conservation of resources. Subjects, workers, materials, and financial resources are rapidly consumed in research. In an era of increasing resource consciousness, careful planning for potential midcourse corrections that would save on trial cost and maintain ethical standards of the study is a laudable goal. However, the investigator must remember that (1) they are engaged in sample-based research, (2) the process of drawing generalizable conclusions from sample-based research is challenging, and (3) the smaller the sample size that a researcher is working with, the more likely one is to be misled.

Bayesian Analyses
Bayesian approaches are becoming increasingly common in biostatistics, and in some circumstances, can represent a viable alternative to hypothesis testing. The investigator who uses a Bayesian analysis has an alternative to the classic P value. Essentially, the investigator begins with a probability distribution of the parameter of interest. The data are then used to update this probability, essentially converting the previous probability distribution into a posterior probability. This permits the investigator to address specific questions of direct relevance, for example, “What is the probability that the effect size of the study is greater than zero?” In addition, using a loss function permits the researchers to compute an estimate of the treatment effect known as the Bayes estimate. One must take care, however, in choosing the previous distribution and loss function. Adaptive designs (ie, allowing interim evaluations of efficacy to determine the ratio of treatment to control group subjects to reduce the overall sample size of the study) are becoming more common.

Common Mistakes to Avoid
Most epidemiological and biostatistics mistakes in cardiovascular research derive from 1 flaw—an inadequately prepared protocol. In its absence, the researcher is left to flounder in a rushing river of administrative, recruitment, and logistical issues that threaten to swamp any modern research effort.

Authors of a good protocol recognize at once the information needed to conduct the research effort successfully. The scientific question is essential. From this, end points are derived that define the instruments for measurement that will be used, followed by a determination of the number of treatment groups, and then a calculation of the required sample size and duration of the study. Dwelling on these concerns early simplifies the execution of the study.

When investigators have a solid, carefully considered, and well-written protocol, they are best advised to adhere to it. There will be many temptations to change. Other studies with interesting findings will be reported, leading investigators to new questions they may wish to pursue. New measurement tools will be discovered. Regulatory agencies may change direction. It is rare occurrence that these interceding events strengthen the study; instead, they threaten its vitiation. Investigators should maintain their commitment to the protocol in these circumstances. They can certainly report exploratory analyses, but these analyses must be acknowledged as such and not be permitted to overshadow the findings of the primary end points, whether they are positive or negative.

In addition, investigators should acknowledge and obtain any epidemiological or biostatistical support that they need. The statistical analyst should keep the statistical analysis as simple as possible. The focus of the investigator’s query is to answer a cardiovascular question, and the study should be designed and the analysis conducted as simple as possible. The computations should be in line with the standards and expectations of the cardiology community. If a new computation is required, the investigator should precede this by using the standard computation, so that investigators can compare the 2 and come to a conclusion about the added value of the new approach. In this type of research, the statistics should be like the solid foundation of a house: all but invisible to the naked eye, but providing the solid basis from which to determine the exposure–disease relationship.

Investigators who are mindful of the methodologies depicted here can produce research results that permit the community to assimilate their conclusions. There is no doubt that there will be disappointments along the way, as data have no obligation to provide support for a researcher’s incorrect ideas. Data do, however, provide a window to the true state of nature, and through that, illumination.

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