Learn Six Sigma

A Lean Six Sigma Green Belt Training Guide

Featuring Examples Using Minitab v.18



Lean Sigma Corporation Michael Parker

LEARN SIX SIGMA Using Minitab v.18

A LEAN SIX SIGMA GREEN BELT TRAINING GUIDE FEATURING EXAMPLES FROM MINITAB V.18

Lean Sigma Corporation

Michael Parker

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0.0 INTRODUCTION

This book has been written to explain the topics of Lean Six Sigma and provide step by step instructions on how perform key statistical analysis techniques using Minitab. The content of this book has been updated to include instructions using Minitab version 18.

This book will provide the reader with all the necessary knowledge and techniques to become an effective Lean Six Sigma practitioner. For those who have already achieved certification this book is an excellent reference manual as well as companion for all who pass on the virtues of Lean Six Sigma.

Another valuable component to this publication is the use of numerous step by step analysis instructions for the reader to learn exactly how to perform and interpret statistical analysis techniques using Minitab. Anywhere throughout this book where you see the image of a chart on a clipboard followed by the words "Data file:" will be a place where the reader will need the necessary data file to accurately follow the exercise. This data file can be downloaded from your account in your Study Six Sigma portal.

1.0 DEFINE PHASE

1.1 SIX SIGMA OVERVIEW

1.1.1 WHAT IS SIX SIGMA?

In statistics, sigma (σ) refers to standard deviation, which is a measure of variation. You will come to learn that variation is the enemy of any quality process; it makes it much more difficult to meet a customer's expectation for a product or service. We need to understand, manage, and minimize process variation.

Six Sigma is an aspiration or goal of process performance. A Six Sigma goal is for a process average to operate approximately 6σ away from the customer's high and low specification limits. A process whose average is about 6σ away from the customer's high and low specification limits has abundant room to "float" before approaching the customer's specification limits.

Most people think of Six Sigma as a disciplined, data-driven approach to eliminating defects and solving business problems. If you break down the term, Six Sigma, the two words describe a measure of quality that strives for near perfection.

A Six Sigma process only yields 3.4 defects for every 1 million opportunities! In other words, 99.9997% of the products are defect-free, but some processes require more quality and some require less.

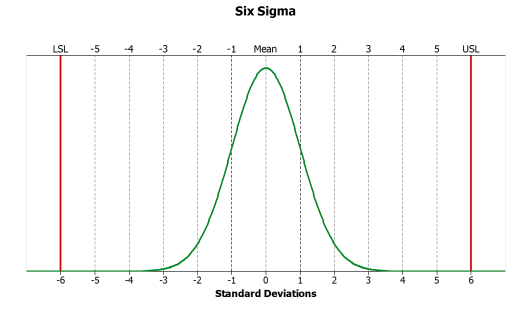


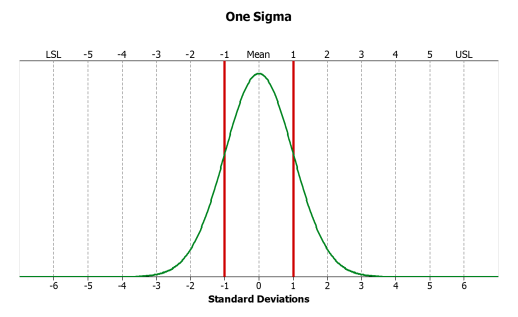
Fig. 1.1 Six Sigma Process with mean 6 standard deviations away from either specification limit.

The more variation that can be reduced in the process (by narrowing the distribution), the more easily the customer's expectations can be met.

It is important to note that a Six Sigma level of quality does not come without cost, so one must consider what level of quality is needed or acceptable and how much can be spent on resources to remove the variation.

What is Six Sigma: Sigma Level

Sigma level measures how many sigma there are between your process average and the nearest customer specification. Let us assume that your customer's upper and lower specifications limits (USL and LSL) were narrower than the width of your process spread. The USL and LSL below stay about one standard deviation away from the process average. Therefore, this process operates at *one sigma*.





In Fig. 1.1, the LSL and USL were at six standard deviations from the mean. Would that process be more or less forgiving than the one shown in Fig. 1.2?

Answer: *This* process is much less forgiving. It is a one sigma process because the USL and LSL are only one standard deviation from the mean. The area under the blue curve to the left of the LSL and to the right of the USL represents *process defects*.

A process operating at one sigma has a defect rate of approximately 70%. This means that the process will generate defect-free products only 30% of the time, so for every three units that are good, seven are defective. Obviously, a one sigma process is not desirable anywhere because its implications are high customer claims, high process waste, lost reputation, and many others.

What about processes with more than one sigma level? A higher sigma level means a lower defect rate. Let us look at the defect rates of processes at different sigma levels.

Table 1.1 shows each sigma level's corresponding defect rate and DPMO (defects per million opportunities). The higher the sigma level, the lower the defective rate and DPMO. The table shows how dramatically the quality can improve when moving from a one sigma process, to two-sigma, to three, and so on. Next, we will look at how this concept applies to some everyday processes.

Sigma Level	Defect Rate	DPMO
1	69.76%	697612
2	30.87%	308770
3	6.68%	66810
4	0.62%	6209
5	0.023%	232
6	0.00034%	3.4

Table 1.1 Sigma level, Defect Rate and DPMO

Let us look at processes operating at three-sigma, which has a defect rate of approximately 7%. What would happen if processes operated at three-sigma? According to http://www.qualityamerica.com:

- Virtually no modern computer would function.
- 10,800,000 health care claims would be mishandled each year.
- 18,900 US savings bonds would be lost every month.
- 54,000 checks would be lost each night by a single large bank.
- 4,050 invoices would be sent out incorrectly each month by a modest-sized telecommunications company.
- 540,000 erroneous call details would be recorded each day from a regional telecommunications company.
- 270 million erroneous credit card transactions would be recorded each year in the United States.

Just imagine what a three-sigma process would look like in the tech industry, in health care, in banking. Can you apply this concept to a process that you relate to? What if processes operated with 1% defect rate? There would be:

- 20,000 lost articles of mail per hr. (Implementing Six Sigma Forest W. Breyfogle III).
- Unsafe drinking water almost 15 minutes per day.
- 5,000 incorrect surgical operations per week.
- Short or long landings at most major airports each day.
- 200,000 wrong drug prescriptions each year.
- No electricity for almost seven hours per month.

Even at a 1% defect rate, some processes would be unacceptable to you and many others. Would you drink the water in a community where the local water treatment facility operates at a 1% defect rate? Would you go to a surgeon with a 1% defect rate? So, what is Six Sigma? Sigma is the measure of quality, and Six is the *goal*.

What is Six Sigma: The Methodology

Six Sigma itself is the *goal*, not the method. To achieve Six Sigma, you need to improve your process performance by:

- Minimizing the process variation so that your process has enough room to fluctuate within customer's spec limits.
- Shifting your process average so that it is centered between your customer's spec limits.

Accomplishing these two process improvements, along with stabilization and control (the ability to maintain the process improvements or level of quality over time), you can achieve Six Sigma. This way, the process becomes more capable of meeting the specification limits that are set by the customer.

The methodology prescribed to achieve Six Sigma is called DMAIC. DMAIC is a systematic and rigorous methodology that can be applied to *any* process in order to achieve Six Sigma. It consists of five phases of a project:

- 1. **D**efine
- 2. Measure
- 3. Analyze
- 4. Improve
- 5. Control

You will be heavily exposed to many concepts, tools, and examples of the DMAIC methodology through this training. The goal of this training is to teach you how to apply DMAIC and leverage the many concepts and tools within it. At the completion of the curriculum, you will become capable of applying the DMAIC methodology to improve the performance of *any* process.

1.1.2 Six Sigma History

The Six Sigma terminology was originally adopted by Bill Smith at Motorola in the late 1980s as a quality management methodology. As the "Father of Six Sigma," Bill forged the path for Six Sigma through Motorola's CEO Bob Galvin who strongly supported Bill's passion and efforts.

Bill Smith originally approached Bob Galvin with what he referred to as the "theory of latent defect." The core principle of the theory is that *variation* in manufacturing processes is the main driver for *defects*, and eliminating variation will help eliminate defects. In turn, it will eliminate the wastes associated with defects, saving money and increasing customer satisfaction. The threshold agreed to by Motorola was 3.4 defects per million opportunities. Does it sound familiar?

Starting from the late 1980s, Motorola extensively applied Six Sigma as a process management discipline throughout the company, leveraging Motorola University. In 1988, Motorola was

recognized with the prestigious Malcolm Baldrige National Quality Award for its achievements in quality improvement.

Six Sigma has been widely adopted by companies as an efficient way of improving the business performance since General Electric implemented the methodology under the leadership of Jack Welch in the 1990s. As GE connected Six Sigma results to its executive compensation and published the financial benefits of Six Sigma implementation in their annual report, Six Sigma became a highly sought-after discipline of quality. Companies across many industries followed suit. In some cases GE taught companies how to deploy the methodology, and in many cases experts from GE and other pioneer companies were heavily recruited to companies that were new to methodology.

Most Six Sigma programs cover the aspects, tools, and topics of Lean or Lean Manufacturing. The two, work hand in hand, benefitting each other. Six Sigma focuses on minimizing process variability, shifting the process average, and delivering within customer's specification limits. Lean focuses on eliminating waste and increasing efficiency.

While the term Lean was coined in the 1990s, the methodology is mainly based on the concepts of the Toyota Production System, which was developed by Taiichi Ohno, Shigeo Shingo, and Eiji Toyoda at Toyota between 1948 and 1975.

Lean and its popularity began to form and gain significant traction in the mid-1960s with the Toyota initiative TPS or Toyota Production System, originally known as the "just-in-time production system." The concepts and methodology of Lean, however, were fundamentally applied much earlier by both Ford and Boeing in the early 1900s.

The DMAIC methodology is essentially an ordered collection of concepts and tools that were not unique to Six Sigma. The concepts and tools are leveraged in a way that leads down a problem-solving path.

Despite the criticism and immaturity of Six Sigma in many aspects, its history continues to be written with every company and organization striving to improve its business performance. The track record for many of these companies after implementing Six Sigma, speaks for itself.

1.1.3 Six Sigma Approach

Six Sigma Approach: Y = f(x)

The Six Sigma approach to problem solving uses a transfer function. A *transfer function* is a mathematical expression of the relationship between the inputs and outputs of a system. The relational transfer function that is used by all Six Sigma practitioners is:

Y = f(x)

The symbol Y refers to the measure or output of a process. Y is usually your primary metric, the measure of process performance that you are trying to improve. The expression f(x) means

function of x, where x's are factors or inputs that affect the Y. Combined, the Y = f(x) statement reads "Y is a function of x," in simple terms, "My process performance is dependent on certain x's."

The objective in a Six Sigma project is to identify the critical x's that have the most influence on the output (Y) and adjust them so that the Y improves. It is important to keep in mind this concept throughout the DMAIC process. The system (or process) outputs (Y's) are a function of the inputs (x).

Example

Let us look at a simple example of a pizza delivery company that desires to meet customer expectations of on-time delivery.

Measure = on-time pizza deliveries Y = percent of on-time deliveries

f(x) would be the x's or factors that heavily influence timely deliveries

```
x1: might be traffic
x2: might be the number of deliveries per driver dispatch
x3: might be the accuracy of directions provided to the driver
x4: might be the reliability of the delivery vehicle
```

The statement Y = f(x) in this example will refer to the proven x's determined through the steps of a Six Sigma project. Most would agree that an important requirement of pizza delivery is to be fast or on-time. In this example, let us consider factors that might cause variation in Y (percent of on-time deliveries). In the DMAIC process, the goal is to determine which inputs (x's) are the main drivers for the variation in Y.

With this approach, all potential x's are evaluated throughout the DMAIC methodology. The x's should be narrowed down until the vital few x's that significantly influence on-time pizza deliveries are identified.

This approach to problem solving will take you through the process of determining all potential x's that *might* influence on-time deliveries and then determining through measurements and analysis which x's *do* influence on-time deliveries. Those significant x's become the ones used in the Y = f(x) equation.

The Y = f(x) equation is a very powerful concept and requires the ability to measure your output and quantify your inputs. Measuring process inputs and outputs is crucial to effectively determining the significant influences to any process.

1.1.4 Six Sigma Methodology

Six Sigma is a data-driven methodology for solving problems, improving, and optimizing business problems. Six Sigma follows a methodology that is conceptually rooted in the

principles of a five-phase project. Each phase has a specific purpose and specific tools and techniques that aid in achieving the phase objectives.

The five phases of DMAIC are:

- 1. **D**efine
- 2. Measure
- 3. Analyze
- 4. Improve
- 5. **C**ontrol

The goals of the five phases are:

- 1. **D**EFINE—To define what the project is setting out to do and scope the effort
- 2. **M**EASURE—To establish a baseline for the process, ensure the measurement system is reliable, and identify all possible root causes for a problem
- 3. **A**NALYZE—To narrow down all possible root causes to the critical few that are the primary drivers of the problem
- 4. IMPROVE—To develop the improvements for the process
- 5. **C**ONTROL—To implement the fix and a control plan to ensure the improvements are sustained over time

In terms of the transfer function, the five phases mean:

- 1. **D**EFINE—Understand the project Y's and how to measure them
- 2. MEASURE—Prioritize potential x's and measure x's and Y's
- 3. ANALYZE Test x–Y relationships and verify/quantify important x's
- 4. IMPROVE—Implement solutions to improve Y's and address important x's
- 5. **C**ONTROL—Monitor important x's and the Y's over time

Six Sigma Methodology: Define Phase

The goal of the *Define* phase is to establish a solid foundation and business case for a Six Sigma project. Define is arguably the most important aspect of any Six Sigma project. It sets the foundation for the project and, if it is not done well and properly thought, it is very easy for a project to go off-track.

All successful projects start with a current state challenge or problem that can be articulated in a quantifiable manner—without a baseline and a goal, it is very difficult to keep boundaries around the project. But it is not enough to just know the problem, you must quantify it and also determine the goal. Without knowing how much improvement is needed or desired, it can be very difficult to control the scope of the project.

Once problems and goals are identified and quantified, the rest of the Define phase will be about valuation, team, scope, project planning, timeline, stakeholders, Voice of the Customer, and Voice of the Business.

Define Phase Tools and Deliverables

1. *Project charter*, which establishes the:

- Business Case
- Problem Statement
- Project Objective
- Project Scope
- Project Timeline
- Project Team

The project charter is essentially a contract formed between a project team, the champion, and the stakeholders. It defines what the project is going to do, why they are going to do it, when it will be done, and by whom. It includes the business case, problem statement, project objective, scope, timeline, and team.

2. Stakeholder Assessment

Stakeholder assessment involves the following:

- High-Level Pareto Chart Analysis
- High-Level Process Map
- Voice of the Customer/ Voice of the Business and Critical to Quality Requirements Identified and Defined
- Financial Assessment

A stakeholder assessment is done to understand where there are gaps in stakeholder support and develop strategies to overcome them. High-level Pareto chart and process maps, along with Voice of the Customer, Voice of the Business, and Critical to Quality Requirements also help to develop the scope and put some guardrails around the process.

Six Sigma Methodology: Measure Phase

The goal of the *Measure* phase is to gather baseline information about the process (process performance, inputs, measurements, customer expectations etc.). This phase is necessary to determine if the measurement systems are reliable, if the process is stable, and how capable the process is of meeting the customer's specifications.

Throughout the Measure phase you will seek to achieve a few important objectives:

- Gather All Possible x's
- Assess Measurement System and Data Collection Requirements
- Validate Assumptions
- Validate Improvement Goals
- Determine Cost of Poor Quality
- Refine Process Understanding

- Determine Process Stability
- Determine Process Capability

Measure Phase Tools and Deliverables

The tools and deliverables for this phase are:

- Process Maps, SIPOC, Value Stream Maps—To visualize a process
- Failure Modes and Effects Analysis—To identify possible process failure modes and prioritize them
- Cause-and-Effect Diagram—To brainstorm possible root causes for defects
- XY Matrix—To prioritize possible root causes for defects
- Six Sigma Statistics
- Basic Statistics and Descriptive Statistics—To understand more about your process data
- Measurement Systems Analysis—To establish repeatability, reproducibility, linearity, accuracy, and stability
- Variable and/or Attribute Gage R&R
- Gage Linearity and Accuracy or Stability
- Basic Control Charts—To assess process stability
- Process Capability and Sigma Levels—To assess and quantify a process' ability to meet customer specification limits
- Data Collection Plan—To ensure that when data is collected, it is done properly

Six Sigma Methodology: Analyze Phase

The *Analyze* phase is all about establishing verified drivers. In the DMAIC methodology, the Analyze phase uses statistics and higher-order analytics to discover relationships between process performance and process inputs. In other words, this phase is where the potential root causes are narrowed down to the *critical* root causes.

Statistical tools are used to determine whether there are relationships between process performance and the potential root causes. Where the strong relationships are discovered, these become the foundation for the solution to improve the process.

Ultimately, the Analyze phase establishes a reliable hypothesis for improvement solutions. During the Analyze phase, one needs to:

- Establish the Transfer Function Y = f(x)
- Validate the List of Critical x's and Impacts
- Create a Beta Improvement Plan (e.g., pilot plan)

Analyze Phase Tools and Deliverables

The Analyze phase is about proving and validating critical x's using the appropriate and necessary analysis techniques. The tools that help to formulate a hypothesis about how much improvement can be expected in the Y, given a change in the x include:

- Hypothesis Testing (e.g., t-tests, Chi-Square)
- Parametric and Non-Parametric
- Regression (Simple Linear Regression, Multiple Linear Regression)—To establish quantitative and predictive relationships between the x's and Y's

The Analyze phase is also about establishing a set of solution hypotheses to be tested and further validated in the Improve phase.

Six Sigma Methodology: Improve Phase

The goal of the *Improve* phase is make the improvement. Improve is about designing, testing, and implementing your solution. To this point, you have defined the problem and objective of the project, brainstormed possible x's, analyzed and verified critical x's. Now it's time to make it real! During the Improve phase, the following are necessary:

- Statistically Proven Results from Active Study/Pilot
- Improvement/Implementation Plan
- Updated Stakeholder Assessment
- Revised Business Case with Return on Investment
- Risk Assessment/Updated Failure Modes and Effects Analysis
- New Process Capability and Sigma

Methodologies for the Improve phase include:

- Experiments and planned studies
- Pilots or tests designed to validate relationships and determine how much change in an input is needed to induce the desired result in the output
- Implementation plan to stimulate thoughts and planning for any necessary communications, training, and preparations to implement the process improvement

Improve Phase Tools and Deliverables

The tools and deliverables of this phase are:

- Any appropriate tool from previous phases
 - $\circ~$ An updated stakeholder assessment to ensure the right support exists to make the process change
 - $\circ~$ An updated business case to justify the change
 - An updated Failure Modes and Effects Analysis to ensure the solution is robust and proper control points are identified
 - A revisited process capability and sigma level to quantify how well the improved process will perform against customer specifications
- Design of Experiment: Full Factorial and Fractional Factorial
- Pilot or Planned Study using Hypothesis Testing and Valid Measurement Systems
- Implementation Plan

Six Sigma Methodology: Control Phase

The last of the five core phases of the DMAIC methodology is the *Control* phase. The purpose of this phase is to establish the mechanisms to ensure the process sustains improved performance and, if it does not, to establish a reaction/mitigation plan to activate by a process owner. The Control phase is a common failure point for projects. Improvements are made, but gains will only be temporary if the proper controls are not in place.

Control Phase Tools and Deliverables

The tools and deliverables of this phase are:

- Statistical Process Control (SPC/Control Charts): IMR, Xbar-S, Xbar-R, P, NP, U, C etc.— For monitoring process inputs and outputs for stability and quickly identifying when the process goes "out of control"
 - Control Plan Documents
 - Control Plan—To ensure control points are identified and accountability for monitoring them is taken
 - Training Plan—To ensure employees are properly trained to perform process changes
 - Communication Plan—To alert any stakeholders of a change
 - Audit Checklist
- Lean Control Methods—To mistake-proof a process
 - o Poka-Yoke
 - 5-S—To organize the workplace and make it more visual
 - o Kanban

1.1.5 ROLES AND RESPONSIBILITIES

The various roles in a Six Sigma program are commonly referred to as "Belts." In addition to Belts, there are also other key roles with specific responsibilities.

It is important to know the many roles in Six Sigma to understand where responsibilities lie to execute a project. Let us explore the different roles and their corresponding responsibilities in a Six Sigma program. Each of the four Six Sigma belts represents a different level of expertise in the field of Six Sigma:

- Six Sigma Master Black Belt
- Six Sigma Black Belt
- Six Sigma Green Belt
- Six Sigma Yellow Belt

Different Belt levels are associated with differing levels of expertise in Six Sigma, with Master Black Belt being the most highly trained expert, to Yellow Belt having very basic knowledge. In addition to Belts, there are other critical and complementary roles, very important to the success of Six Sigma initiatives:

- Champions
- Sponsors
- Stakeholders
- Subject Matter Experts

Roles and Responsibilities: Master Black Belt

The *Master Black Belt* (MBB) is the most experienced, educated, and capable Six Sigma expert, often thought of as a trusted advisor to high-level leaders.

A typical MBB has managed dozens of Black Belt level projects. An MBB can simultaneously lead multiple Six Sigma Belt projects, ensuring the proper and consistent application of Six Sigma across departments, while mentoring and certifying Black Belt and Green Belt candidates.

An MBB typically works with high-level operations directors, senior executives, and business managers to help with assessing and planning business strategies and tactics. This is why the MBB needs to have a strong ability to communicate with and influence leaders at all levels of an organization.

An MBB commonly advises management team on the cost of poor quality of an operation and consults on methods to improve business performance.

While an MBB is typically technically savvy with all of the concepts and tools, it is critical that he or she have the skills to communicate in a practical manner with those that are not well-versed in Six Sigma.

Typical Responsibilities of an MBB An MBB:

- Identifies and defines the portfolio of projects required to support a business strategy
- Establishes scope, goals, timelines, and milestones
- Assigns and marshals resources
- Trains and mentors Green Belts and Black Belts
- Facilitates tollgates or checkpoints for Belt candidates
- Reports-out/updates stakeholders and executives
- Establishes organization's Six Sigma strategy/roadmap
- Leads the implementation of Six Sigma

Roles and Responsibilities: Black Belt

The *Black Belt* (BB) is the most active and valuable experienced Six Sigma professional among all Six Sigma Belts. The Black Belt is the key to success among all the other Belts.

A typical BB has:

- Led multiple projects
- Trained and mentored various Green Belts candidates
- Understood how to define a problem and drive effective solution

The BB incorporates many skills that are critical to successfully and quickly implementing a process improvement initiative through the DMAIC methodology. The BB is well rounded in terms of project management, statistical analysis, financial analysis, meeting facilitation, prioritization, and a range of other value-added capabilities, which makes a BB highly valuable asset in the business world.

BBs commonly serve as the dedicated resource continuing their line management role while simultaneously achieving a BB certification.

Typical Responsibilities of a BB

A BB has the following responsibilities:

- Project Management
 - Defines projects, scope, teams etc.
 - Marshals resources
 - Establishes goals, timelines, and milestones
 - Provides reports and/or updates to stakeholders and executives
- Task Management
 - o Establishes the team's Lean Sigma roadmap
 - Plans and implements the use of Lean Sigma tools
 - Facilitates project meetings
 - o Does project management of the team's work
 - Manages progress toward objectives
- Team Management
 - \circ $\,$ Chooses or recommend team members $\,$
 - Defines ground rules for the project team
 - Coaches, mentors, and directs project team
 - Coaches other Six Sigma Belts
 - Manages the team's organizational interfaces

Roles and Responsibilities: Green Belt

The *Green Belt* (GB) is considered as a less intense version of Six Sigma professional compared to the Black Belt (BB). A GB is exposed to all the comprehensive aspects of Six Sigma with less focus on the statistical theories and some other advanced analytical methodologies such as Design of Experiment. When it comes to project management, a GB has almost the same responsibilities as a BB. In general, the GB works on less complicated and challenging business problems than a BB.

While a Black Belt's sole business responsibility could be the implementation of projects, Green Belts often implement projects in addition to other job responsibilities. Therefore, the Green Belt typically works on less complicated business problems. A Green Belt takes direction and coaching from the Black Belt.

Typical Responsibilities of a Green Belt

A GB has the following responsibilities:

- Project Management
 - Defines the project, scope, team etc.
 - Marshals resources
 - Sets goals, timelines, and milestones
 - Reports-out/updates stakeholders and executives
- Task Management
 - o Establishes the team's Lean Sigma Roadmap
 - o Plans and implements the use of Lean Sigma tools
 - Facilitates project meetings
 - o Does Project Management of the team's work
 - Manages progress toward objectives
- Team Management
 - Chooses or recommends team members
 - Defines ground rules for the project team
 - \circ $\,$ Coaches, mentors, and directs project team $\,$
 - Coaches other Six Sigma Belts
 - Manages the team's organizational interfaces

As you can see, the roles and responsibilities are identical to those of a Black Belt. The key differences are the level of complexity and the dedication that a Black Belt has toward Six Sigma projects. Again, a Green Belt typically does projects in addition to his or her normal line job responsibilities.

Roles and Responsibilities: Yellow Belt

The *Yellow Belt* (YB) understands the basic objectives and methods of a Six Sigma project. YB has an elementary understanding about what other Six Sigma Belts (GB, BB, and MBB) are doing to help them succeed.

In a Six Sigma project, YB usually serves as a subject matter expert regarding some aspects of the process or project. Supervisors, managers, directors, and sometimes executives are usually trained at the YB level. This level of training is important to have in an environment where Six Sigma projects are prevalent, such as when managing Belts.

Typical Responsibilities of a Yellow Belt A Yellow Belt:

- Helps define process scope and parameters
- Contributes to team selection process

- Assists in information and data collection
- Participates in experiential analysis sessions (Failure Modes and Effects Analysis, Process Mapping, Cause and Effect etc.)
- Assists in assessing and developing solutions
- Delivers solution implementations

Roles and Responsibilities: Champions and Sponsors

Champions and sponsors are those individuals (directors, executives, managers etc.) chartering, funding, or driving the Six Sigma projects that BBs and GBs are conducting. Champions and Sponsors' roles are to set direction for projects and ensure that the path is clear for the Belts to succeed: obtain resources and funding, overcome stakeholder issues, and knock down other barriers to progress.

Champions and sponsors need to have a basic understanding of the concepts, tools, and techniques involved in the DMAIC methodology so that they can provide proper support and direction. Champions and sponsors play critical roles in the successful deployment of Six Sigma. Strong endorsement of Six Sigma from the leadership team is critical for success.

Typical Responsibilities of a Champion or Sponsor

A Champion or Sponsor typically:

- Maintains a strategic oversight
- Establishes strategy and direction for a portfolio of projects
- Clearly defines success
- Provides resolution for issues such as resources or politics
- Establishes routine tollgates or project reviews
- Clears the path for solution implementation
- Assists in project team formation

From a project standpoint, it is important for the Champions and Sponsors to be highly engaged in the work. It is their role to provide oversight and guidance to define success for the project or projects. They resolve issues when necessary, such as resource needs, stakeholder conflicts, or political issues. They create the routines for tollgate reviews to ensure there are regular checkpoints with the project team.

Roles and Responsibilities: Stakeholders

Stakeholders are usually the recipients or beneficiaries of the success of a Six Sigma project. Generally speaking, a stakeholder is anyone who has interest in a project; stakeholders are usually the beneficiaries of the process improvement. A stakeholder can be a person, a group, or an organization that is affected by the project.

Stakeholders are individuals owning the process, function, or production/service line that a Six Sigma Belt focuses on improving the performance of.

BBs and GBs need to keep strong working relationships with stakeholders because without their support, it would be extremely difficult to make the Six Sigma project a success. A lack of stakeholder support can cause a project to fail, either through active or passive resistance, so it is very important to be aware of how stakeholders feel about the work being done so the team can proactively manage their perceptions and level of support.

Roles and Responsibilities: Subject Matter Experts

Subject Matter Experts (SMEs) are commonly known as the experts of the process or subject matter. They play critical roles to the success of a project. Six Sigma Belts should proactively look to key SMEs to round out their working project team. Based on SMEs' extensive knowledge about the process, they have the experience to identify which solutions can work and which cannot work.

SMEs who simply do not speak up can hurt the chances of the process' success. SMEs' input is critical to ensure the team is leveraging accurate information when it comes to the process or evaluating data and information.

SMEs are also the same people who prefer to keep the status quo. Six Sigma Belts may find many of them unwilling to help implement the changes. When selecting SMEs, it is important to choose people who are vocal, but who also bring ideas. SMEs that are closed-minded to change may be vocal, but can be a barrier to making the necessary improvements.

SUMMARY

Throughout this module, we have reviewed the various common roles and corresponding responsibilities in any Six Sigma program:

- Six Sigma Master Black Belt
- Six Sigma Black Belt
- Six Sigma Green Belt
- Six Sigma Yellow Belt

- Champion and Sponsors
- Stakeholders
- Subject Matter Experts

These Six Sigma belts and other roles are designed to deliver value to the business effectively and successfully. For a Six Sigma program to be effective, it is important to assign these roles to individuals who are well equipped to carry out the responsibilities.

1.2 SIX SIGMA FUNDAMENTALS

1.2.1 DEFINING A PROCESS

What is a Process Map?

A *process map* is a graphical representation of a process flow. It is necessary to document a process because process maps:

- Visually show how the business process is accomplished in a step by step manner.
- Describe how information or materials sequentially flow from one entity to the next.
- Illustrate who is responsible for steps and actions.
- Depict the input and output of each individual process step.

In the Measure phase, the project team should map the current state of the process. Teams should be sure not to fall into the trap of mapping an ideal state or what they think the process should look like.

Process Map Basic Symbols

Figure 1.3 below depicts a basic process map that was created with the four most commonly used process mapping symbols. The four symbols are:

- Oval Used for start and end points
- Square or Rectangle Used for process steps
- Diamond Used for decision points
- Arrow Used to represent flow direction

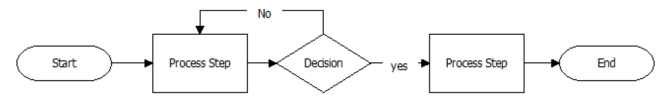
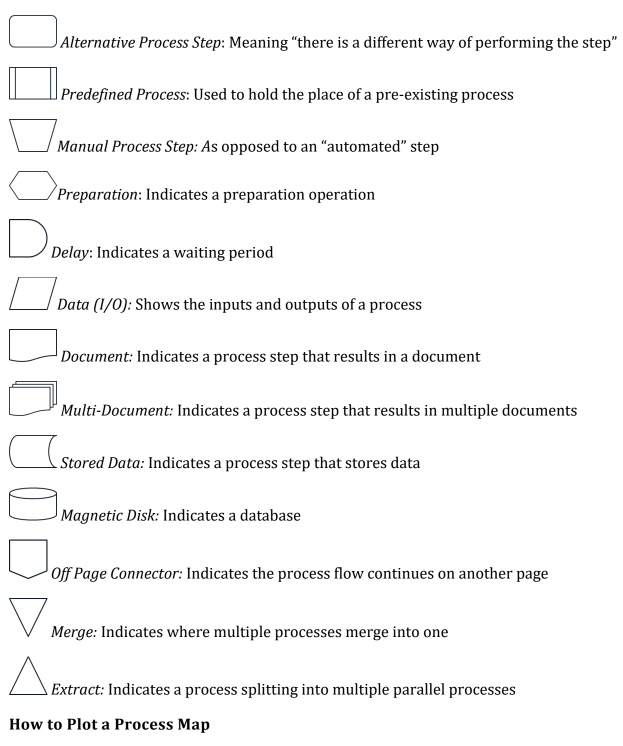


Fig. 1.3 Basic Process Mapping Symbols

Additional Process Mapping Symbols

Below are additional process mapping symbols that may be used to provide further information and detail when developing a process map. Although these symbols are not as common as those shown in figure 1.3, they are universally understood to represent process information as described:



Process Mapping Step 1: Define boundaries of the process you want to map.

- A process map can depict the flow of an entire process or a segment of it.
- You need to identify and define the beginning and ending points of the process before starting to plot.
- Use operational definitions.

The first step in plotting a process map is deciding where the beginning and ending points are for the process you are trying to depict. You might be trying to map a whole process that begins and ends with the customer, or you may be trying to map out a segment of it.

An *operational definition* is an exact description—in other words, it is useful in removing ambiguity and reduces the chances of getting disparate results from a process if performed by different people.

Process Mapping Step 2: Define and sort the process steps with the flow. There are a few ways to do this:

- Consult with process owners and subject matter experts or observe the process in action to understand how the process is performed.
- Record the process steps and sort them according to the order of their occurrence.

Usually, it is a good idea to do both; sometimes hearing what is supposed to happen might be different than what you observe, this often indicating a problem.

Process Mapping Step 3: Fill the step information into the appropriate process symbols and plot the diagram.

- In the team meeting of process mapping, place the sticky notes with different colors on a white board to flexibly adjust the under-construction process map.
- The flow lines are plotted directly on the white board. For the decision step, rotate the sticky note by 45°.
- When the map is completed on the white board, record the map using Excel, PowerPoint, or Visio.

Mapping out the process with sticky notes on a white board is a good practice to do as a team because the map is easy to adjust until finished. Use a dry-erase marker to draw the flow lines between the sticky notes.

Process Mapping Step 4: Identify and record the inputs/outputs and their corresponding specifications for each process step.

- The process map helps in understanding and documenting Y=f(x) of a process where Y represents the outputs and x represents the inputs.
- The inputs of each process step can be controllable or non-controllable, standardized operational procedure, or noise. They are the source of variation in the process and

need to be analyzed qualitatively and quantitatively in order to identify the vital few inputs that have significant effect on the outcome of the process.

• The outputs of each process step can be products, information, services, etc. They are the little Y's within the process.

Ideally, the specifications should be recorded for each step so it is understood what the inputs and outputs should look like to ensure process efficiency and quality.

Process Mapping Step 5: Evaluate the process map and adjust it if needed.

- If the process is too complicated to be covered in one single process map, you may create additional detailed sub-process maps for further information.
- Number the process steps in the order of their occurrence for clarity.

Functional Process Map or "Swim Lane"

To illustrate the responsibility of different organizations involved in the process, use a functional process map. This type of map is also commonly referred to as a swim lane map. The value of this type of map is that it adds the dimension of accountability.

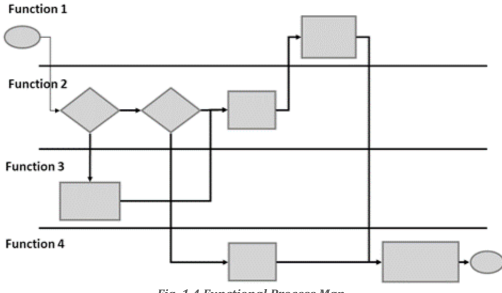


Fig. 1.4 Functional Process Map

Figure 1.4 is a functional process map or "swim lane" diagram which depicts the responsibilities for each step in the process. Swim-lane diagrams can be drawn vertically or horizontally. The swim lanes can show organizations, roles, or functions within a business.

High Level Process Map

Most high-level process maps are also referred to as *flow charts*. The key to a high-level process map is to over-simplify the process being depicted so that it can be understood in its

most generic form. As a general rule, high-level process maps should be four-six steps and no more.

Below, figure 1.5a depicts an oversimplified version of a high-level process map for cooking a 10 lb. prime rib for a dozen holiday guests. As mentioned, high level maps are basic and the challenge is summarizing a process well enough to depict it in 4 – 6 steps.

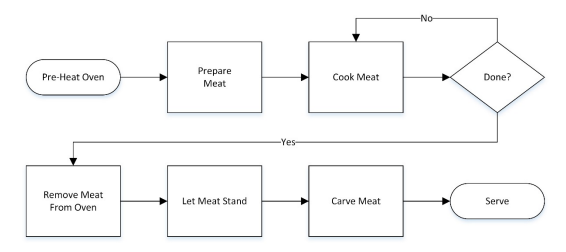


Fig. 1.5a High Level Process Map (cooking prime rib)

Detailed Process Map

Detailed process maps or multi-level maps take the high-level map much further. Detailed maps can be two, three, or more levels deeper than your high-level process map.

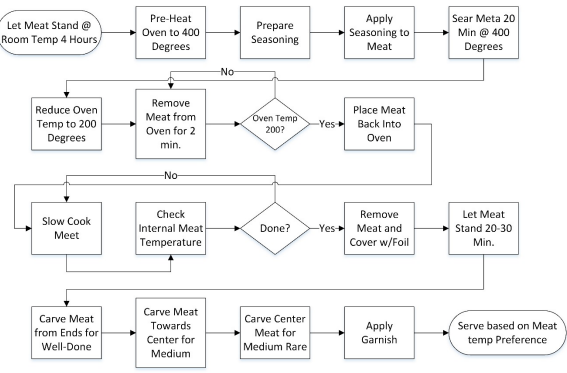


Fig. 1.5b Detailed Process Map (cooking prime rib)

Figure 1.5b is a more detailed representation of our high-level map shown in figure 1.5a. This would be considered a level 2 process map. A good guideline to use to help create the second level is that for each high-level step break it down into another two to four steps each (no more). This is usually a helpful way to index a process, so when you want to understand more about a specific part of a process later, you can find it by starting at the high level and digging down into the process steps you are interested in. Repeat this process (level 3, level 4 etc.) until reaching the desired level of detail. Some detailed maps are two or three levels deep, others can be five or six levels deep. Obviously, the deeper the levels, the more complex and the more burdensome to create and maintain. However, more detail reveals far more information.

What is SIPOC?

A SIPOC (Suppliers, Inputs, Process, Outputs, and Customers) is a high-level visualization tool to help identify and link the different components in a process. SIPOCs summarize the inputs and outputs of a process.

SIPOCs are usually applied in the Measure phase in order to better understand the current state of the process and define the scope of the project. At the outset of a process improvement effort it is very helpful to provide an overview to people who are unfamiliar or need to become reacquainted with a process.

Key Components of a SIPOC

- **S**uppliers: The vendors who provide the raw material, services, and information. Customers can also be suppliers sometimes.
- Input: The raw materials, information, equipment, services, people, environment involved in the process
- **P**rocess: The high-level sequence of actions and decisions that results in the services or products delivered to the customers
- **O**utput: The services or products delivered to the customers and any other outcomes of the process
- **C**ustomers: The end users or recipients of the services or products. Customers can be external or internal to an organization.

How to Plot a SIPOC Diagram

Creating a SIPOC - The first method:

- 1. Create a template that can contain the information of the five key components in a clear way.
- 2. Plot a high-level process map that covers five steps at maximum.
- 3. Identify the outputs of the process.

- 4. Identify the receipt of the process.
- 5. Brainstorm the inputs required to run each process step.
- 6. Identify the suppliers who provide the inputs.

In the first method of creating a SIPOC, begin with the process steps (five steps at most), determine the outputs, identify who the recipients of the process, then work to the left, identifying the inputs and then the suppliers for those inputs.

Creating a SIPOC - The second method:

- 1. Create a template that can contain the information of the five key components in a clear way.
- 2. Identify the receipt of the process.
- 3. Identify the outputs of the process.
- 4. Plot a high-level process map that covers five steps at maximum.
- 5. Brainstorm the inputs required to run each process step.
- 6. Identify the suppliers who provide the inputs.

In the second method, start with the customer (or the recipient of the process), work your way back by identifying the process outputs, then the high-level process map (five steps maximum), brainstorm the inputs to the process, then identify the suppliers for those inputs.

Creating a SIPOC Diagram

Step 1: Vertically List High-Level Process

If you followed the general rules for a high-level process map, then you should have no more than four to six steps for your process. List those steps in a vertical manner in the middle section labeled "Process" as depicted below.

SUPPLIERS	INPUTS	PROCESS	OUTPUTS	CUSTOMERS
		Start		
		Step 1		
		Step 2		
		Step 3		
		Last Step		

Fig. 1.6 SIPOC High Level Process Map

Step 2: List Process Outputs

In step two, for each process step list any outputs that are generated from the corresponding process step. Outputs can be finished goods, documents, reports, raw materials etc.

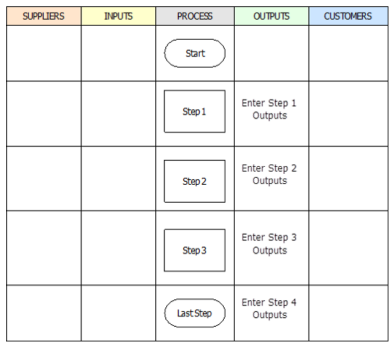


Fig. 1.7 SIPOC Outputs

Step 3: List Output Customers

In step 3, list the customers that receive the outputs from each process step. Customers can be internal or external. The important thing to remember is who is the output being delivered to.

SUPPLIERS	INPUTS	PROCESS	OUTPUTS	CUSTOMERS
		Start		
		Step 1	Enter Step 1 Outputs	Enter Step 1 Customers
		Step 2	Enter Step 2 Outputs	Enter Step 2 Customers
		Step 3	Enter Step 3 Outputs	Enter Step 3 Customers
		Last Step	Enter Step 4 Outputs	Enter Step 4 Customers

Fig. 1.8 SIPOC Output Customers

Step 4: List Process Inputs

For the inputs, in step 4 list any inputs that are necessary for any or all process steps. Inputs can be raw materials, information, finished goods from other processes etc.

SUPPLIERS	INPUTS	PROCESS	OUTPUTS	CUSTOMERS
		Start		
	Enter Step 1 Inputs	Step 1	Enter Step 1 Outputs	Enter Step 1 Customers
	Enter Step 2 Inputs	Step 2	Enter Step 2 Outputs	Enter Step 2 Customers
	Enter Step 3 Inputs	Step 3	Enter Step 3 Outputs	Enter Step 3 Customers
	Enter Step 4 Inputs	Last Step	Enter Step 4 Outputs	Enter Step 4 Customers

Fig. 1.9 SIPOC Process Inputs

Step 5: List Suppliers of Inputs

The last step is to identify and list suppliers of the inputs. Who provides the materials or information as inputs to the process? Where do the inputs come from?

SUPPLIERS	INPUTS	PROCESS	OUTPUTS	CUSTOMERS
		Start		
Enter Step 1	Enter Step 1	Step 1	Enter Step 1	Enter Step 1
Suppliers	Inputs		Outputs	Customers
Enter Step 2	Enter Step 2	Step 2	Enter Step 2	Enter Step 2
Suppliers	Inputs		Outputs	Customers
Enter Step 3	Enter Step 3	Step 3	Enter Step 3	Enter Step 3
Suppliers	Inputs		Outputs	Customers
Enter Step 4	Enter Step 4	Last Step	Enter Step 4	Enter Step 4
Suppliers	Inputs		Outputs	Customers

Fig. 1.10 SIPOC Suppliers of Inputs

SIPOC Benefits

A SIPOC has the following benefits, SIPOCs help to:

- Visually communicate project scope
- Identify key inputs and outputs of a process
- Identify key suppliers and customers of a process
- Verify:
 - Inputs match outputs for upstream processes
 - Outputs match inputs for downstream processes.
 - This type of mapping is effective for identifying opportunities for improvement of your process.
- If you have completed your high-level process map, follow the outlined steps to create a process map of **S**uppliers, **I**nputs, **P**rocess, **O**utputs, and **C**ustomer.

SIPOC is more powerful than a simple high-level process map because it conveys the inputs and outputs of a process, *as well as* the suppliers and customers.

What is Value Stream Mapping?

Value stream mapping is a method to visualize and analyze the path of how information and raw materials are transformed into products or services customers receive. It is used to identify, measure, and decrease the non-value-adding steps in the current process.

Non-Value-Added Activities

Non-value-adding activities are activities in a process that, from the customers' perspective, do not add any other value to the products or services customers demand. Examples of non-value-adding activities are:

- Rework—It should have been done right the first time, so a second time certainly does not add value
- Overproduction—Work that is done before it is needed, and could change, leading to rework
- Excess transportation—Moving the product around does not add value
- Excess stock—Similar to overproduction
- Waiting—Wasted time
- Unnecessary motion—Wasted energy

Not all non-value-adding activities are unnecessary. Sometimes, non-value added activities are required. For example, an inspection that is done for regulatory compliance purposes is necessary but does not add value to the product or service the customer receives.

Keys to Plotting a Value Stream Map

Plot the entire high-level process flow from when the customer places the order to when the customer receives the products or services in the end. A value stream map requires more detailed information for each step than the standard process map:

- Cycle time
- Available time
- Demand & Takt Time
- Inventory
- Working time
- Wait time

Basic Value Stream Map Layout

- Value Added Activities
- Non-Value Added (NVA)
- Preparation type
- Scrap rate
- Rework rate
- Number of operators

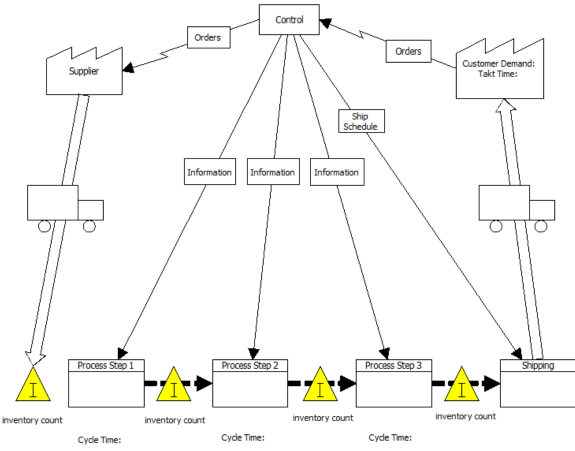


Fig. 1.11 Value Stream Map Layout

Figure 1.11 demonstrates a generic process so that you can see the simple view of the sawtooth box in the upper right where the customer initiates orders. Orders flow to a control point and which are distributed to supplier orders and throughout the production process of four sub-processes which eventually end with the custom receiving the goods.

Value stream maps are useful to determine:

- What the current process looks like
- Where the process starts and where it ends
- How value flows through the process
- The steps which are value added in the process
- The steps which are non-value added in the process
- What the relationship is between information flow and material flow
- Average cycle time throughout the process
- Inventory levels throughout the process
- Sources of the waste
- Areas in need of the most improvement

The true value of the value stream map is not the map itself but the information gathered and assessed to create the map. This information is vital to ascertaining where inefficiencies and opportunities exist.

Additional Mapping Technique

We will now touch on an additional mapping technique that can come in very handy during a process understanding session; the thought process map.

Thought Process Mapping

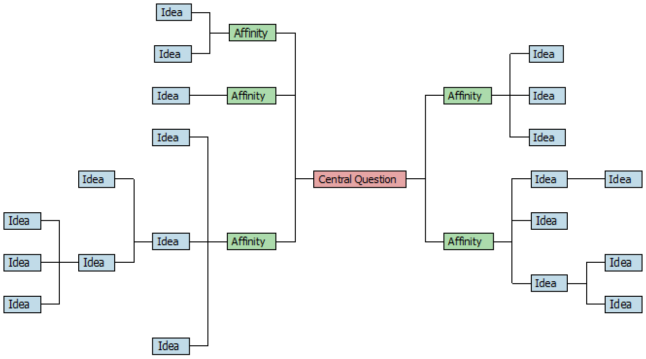


Fig. 1.12 Thought Process Map Example

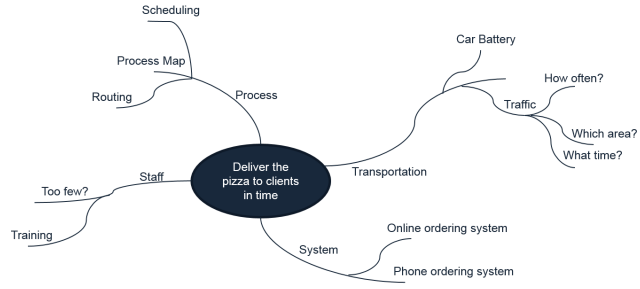
A *thought process map* is a graphical tool to help brainstorm, organize, and visualize information, ideas, questions, or thoughts regarding reaching a project goal. It helps a project team to understand where they have been, where they still need to go, what questions have been answered, how were they answered, and what is yet to be answered. "Thought Maps" are a popular tool used at any phase of a project to:

- Identify knowns and unknowns
- Communicate assumptions and risks
- Discover potential problems and solutions
- Identify resources, information, and actions required to meet the goal
- Present relationship of thoughts

How to Plot a Thought Process Map

1. Define the project goal.

- 2. Brainstorm knowns and unknowns about the project.
- 3. Brainstorm questions and group the unknowns.
- 4. Sequence the questions below the project goal and link related questions.
- 5. Identify tools or methods that would be used to answer the questions.
- 6. Repeat steps 3 to 5 as the project continues until the goal is met.



Thought Process Map Example

Fig. 1.13 Thought Process Map Example

This is an example of a thought process map. In this case, the main theme is on-time pizza delivery. You can see several themes and concepts explored. Thought process maps serve well to introduce questions and ideas while also being great tools to leave "trails" of where you have been so that later in a project when someone makes a suggestion that has already been evaluated you can refer to the thought map as evidence.

1.2.2 VOICE OF THE CUSTOMER AND CRITICAL TO QUALITY

Voice of the Customer

VOC stands for Voice of the Customer. Voice of the customer is a term used for a data-driven plan to discover customer wants and needs. It is *always* important to understand the VOC when designing and improving any Six Sigma process.

There are also other "voices" that need to be heard when conducting projects. The three primary forms are:

- VOC: Voice of the Customer
- VOB: Voice of the Business What does the business require of a process?

• VOA: Voice of the Associate or Voice of the Employee - What does the employee require of a process (e.g., safety, simplicity)?

Gathering VOC

Gathering VOC should be performed methodically. The two most popular methods of collecting VOC are indirect and direct.

1. Indirect data collection for VOC involves passive information exchange, meaning the VOC was not expressly collected for the purposes of the project, but was gathered through byproducts of the process, such as:

- Warranty claims
- Customer complaints/compliments
- Service calls
- Sales reports

All the above are effective ways to understand how the customer feels about a product or service.

2. Indirect methods are less effective, sometimes dated, require heavy interpretation, and are also more difficult to confirm because the data was not collected to understand VOC specific to a business or process problem. Direct data collection methods for VOC are active and planned customer engagements, such as:

- Conducting interviews
- Conducting customer surveys
- Conducting market research
- Hosting focus groups

Direct methods are typically more effective for several reasons:

- Less need to interpret meaning
- Researchers can go a little deeper when interacting with customers and take a direction that might not have been expected or planned
- Customers are aware of their participation and will respond better upon follow-up
- Researchers can properly plan engagements (questions, sample size, information collection techniques etc.)

There are professional organizations that can perform customer data collection on behalf of a company. They commonly plan carefully by designing the questions, determining the sample size, and considering what technique is most appropriate. If there is a need to get very specific customer feedback, a survey is appropriate. However, if the need is more exploratory, interviews or focus groups are more appropriate.

Gathering VOC

- Gathering VOC requires consideration of many factors such as product or service types, customer segments, manufacturing methods or facilities etc. All this information will influence the sampling strategy because different factors might indicate different customer needs across customer segments, or perhaps the customers' feedback may differ by product type or manufacturing method.
- Consider which factors are important and build a sample size plan around them. Also, consider response rates and adjust the initial sample strategy to ensure adequate input is received.
- Once a sampling plan is in place, collect data via the direct and indirect methods discussed earlier.
- After gathering VOC, it will be necessary to translate it into something meaningful: Critical to Quality.

Critical to Quality

CTQs or *Critical to Quality* are generated from VOC feedback. Because VOC is usually vague, emotional, or simply a generalization about products or services, it is important to translate it into something that is applicable to a process: CTQs, which are the quantifiable, measurable, and meaningful translations of VOC.

To create CTQs, the VOC must be organized first into like groupings. One effective way to organize VOC is to group or bucket it using an *affinity diagram*. Affinity diagrams are ideal for large amounts of soft data resulting from brainstorming sessions or surveys.

Affinity Diagram: Building a CTQ Tree

Steps for conducting an affinity diagram exercise:

- Clearly define the question or focus of the exercise. ("Why are associates late for work?").
- Record all participant responses on note cards or sticky notes. (This is the sloppy part, record everything!).
- 3. Lay out all note cards or post the sticky notes onto a wall.
- 4. Look for and identify common themes.
- 5. Begin moving the notes into the themes until all responses are allocated.
- 6. Re-evaluate and adjust.

This is an outline of building an affinity diagram (or CTQ tree); it is typically done with a group of people. The group must agree on what question they are trying to answer, or what the focus for the VOC is. It is good to do this exercise in a group because people bring different perspectives and might understand and organize VOC differently than a single person would.

Affinity Diagram: Building a CTQ Tree Step by Step

In summary, to build an affinity diagram:

- Define the question or focus.
- Record responses on note cards or sticky notes.
- Display all note cards or sticky notes on a wall if necessary.

Why are Associates late to work?

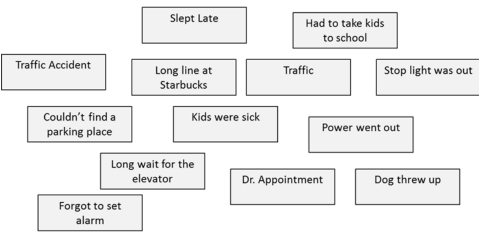


Fig. 1.14 Random reasons prior to an affinity diagram

In figure 1.14 the focus question is "Why are associates late to work?" You can see all of the responses randomly distributed across the page.

• Look for and identify common themes within the responses.

Why are Associates late to work?

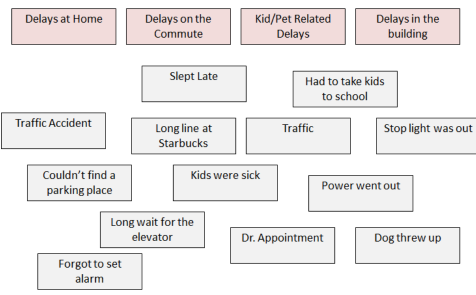


Fig. 1.15 Common themes

In figure 1.15, you can see that some common themes have emerged (delays at home, delays on the commute, kids/pet related delays, or delays in the building).

- Group note cards or sticky notes into themes until all responses are allocated.
- Re-evaluate and make final adjustments.

Delays at Home	Delays on the Commute	Kid/Pet Related Delays	Delays in the building
Slept Late	Traffic Accident	Had to take kids to school	Long line at Starbucks
Power went out	Stop light was out	Kids were sick	Couldn't find a parking place
Dog threw up	Traffic	Dr. Appointment	Long wait for the elevator
Forgot to set alarm			

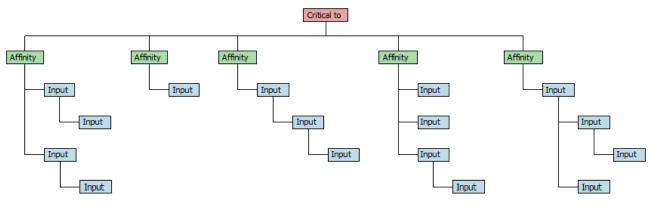
Why are Associates late to work?

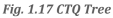


Finally, in figure 1.16 you can now see that the individual responses are lined up under each theme. See any that might be misplaced? How about "dog threw up?" Perhaps "delays at home" and "kid/pet-related delays" do not need to be broken down separately. These are the kinds of things to consider when evaluating and adjusting.

CTQ Tree

The next figure shows an example of a generic CTQ tree transposed from a white board to a software package (*generated with Minitab's Quality Companion 3*)





Kano

Another VOC categorization technique is the *Kano*. The Kano model was developed by Noriaki Kano in the 1980s and it is a graphic tool that further categorizes VOC and CTQs into three distinct groups:

- 1. Must Haves
- 2. Performance Attributes
- 3. Delighters

The Kano helps to:

- Identify CTQs that add incremental value vs. those that are simply requirements
- Understand that having more is not necessarily better

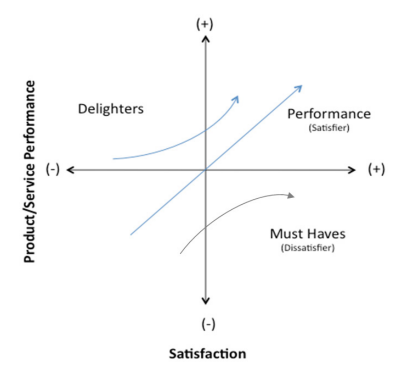


Fig. 1.18 Kano Diagram

It is important to understand what CTQs are non-negotiable (*must haves*) versus those that are nice to have. "Must have" attributes are often taken for granted when they are fulfilled, but result in dissatisfaction when they are not fulfilled. The curve for a "must have" will drop in satisfaction as you move to the left where the need is not fulfilled, but flattens out to a neutral satisfaction level as you move to the right.

A *delighter* is the opposite of the must have—if it is not fulfilled, the customer will not be dissatisfied, but as it is fulfilled, the customer's satisfaction increases. These are things the customer did not realize they wanted or needed, until they have experienced them. Eventually, delighters will become expected, turning them into *performance attributes* or must haves.

Examples:

- How would you classify the steering wheel in a car?
- Low prices at a discount store?
- What about the touch screen on an iPhone (think back to when it came out)?

A performance attribute (or one-dimensional attribute) will increase satisfaction level as the need is fulfilled, and will decrease satisfaction when it is not fulfilled.

Validating VOC and CTQs

Once the process of collecting VOC is complete, and CTQs have been determined, the next step is to *validate* what has been developed. Confirming can be accomplished by conducting surveys through one or more of the following methods:

- Group sessions
- One-on-one meetings
- Phone interviews
- Electronic means (chat, email, social media etc.)
- Physical mail

Consider your confirming audience and try to avoid factors that may influence or bias responses such as inconvenience or overly burdensome time commitments. For example, an overly extensive customer survey might end up getting a rushed response and, therefore, may not represent the true views of an audience.

Translating CTQs to Requirements

To be able to incorporate a CTQ into the design of a process, the next step is to *translate* CTQs to specific process requirements. A *requirements tree* translates CTQs to meaningful and measurable requirements for production processes and products.

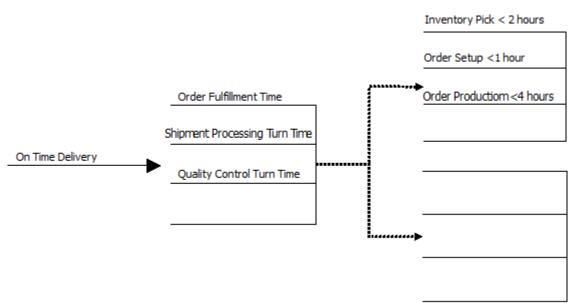


Fig. 1.19 CTQ Requirements Tree

In figure 1.19, the actual feedback from the customer (VOC) might have been something like, "Deliver my product when you say you're going to deliver it". This statement is then translated to something your business can measure such as "on-time delivery." To achieve on time delivery and ensure that you're meeting your customers' expectations, on time delivery must be translated to the very specific process requirements that your business must achieve to meet on time delivery requirements.

To do so, one must then consider what aspects of the process contribute to the time that elapses between when the customer places an order and when it is delivered. The customer's order must be fulfilled, how does that process occur and what are the necessary performance requirements? Also, the order must be inspected by quality control and processed for shipping. What are the performance requirements for those process steps?

The requirements tree help to translate a CTQ into process requirements that can be measured and attained to meet customer expectations.

1.2.3 QUALITY FUNCTION DEPLOYMENT

History of Quality Function Deployment

The QFD (Quality Function Deployment) was developed in 1972 by Shigeru Mizuno (1910– 1989) and Yoji Akao (b. 1928) from the TQM practices of a Japanese company, Mitsubishi Heavy Industries Ltd. QFD aims to design products that assure customer satisfaction and value—the first time and every time.

The QFD framework can be used for translating actual customer statements and needs (the voice of the customer) into actions and designs to build and deliver a quality product. QFD is considered a valuable tool fundamental to a business' success, hence it is widely applied.

What is QFD?

- *QFD* is a construction methodology and quantification tool used to identify and measure customer's requirements and transform them into meaningful and measurable parameters. QFD uses planning matrices—it is also called "The House of Quality."
- QFD helps to prioritize actions to advance process or product to meet customer's anticipations.
- QFD is an excellent tool for contact between cross-functional groups.

Purpose of QFD

The quality function deployment has many purposes, some of the most important being:

- Market analysis to establish needs and expectations
- Examination of competitors' abilities
- Identification of key factors for success
- Translation of key factors into product and process characteristics

QFD focuses on customer requirements, prioritizes resources, and uses competitive information effectively.

Phases of QFD

There are four key phases of QFD:

- Phase I: Product Planning Including the "House of Quality" (Requirements Engineering Life Cycle)
- Phase II: Product Design (Design Life Cycles)
- Phase III: Process Planning (Implementation Life Cycle)
- Phase IV: Process Control (Testing Life Cycle)

How to build a House of Quality

The steps to building a House of Quality are:

• Determine Customer Requirements ("What's" from VOC/CTQ)

 \wedge

- Technical Specifications/Design Requirements ("How's")
- Develop Relationship Matrix ("What's" and "How's")
- Prioritize Customer Requirements
- Conduct Competitive Assessments
- Develop Interrelationship ("How's")
- Prioritize Design Requirements

	Correlation Matrix [How vs. How]		
	Technical Specifications [How]	Why	/
Customer Requirements [What]	Relatinship Matrix [What vs How]	Customer Importance Rating	Customer Market Evaluation (What vs. Why)
How Much	Degree of Technical Difficulty Overall Importance Ratings Technical Competitive Evaluations Target Goals		

Fig. 1.20 House of Quality

This is the structure of a House of Quality. You can see that it contains the following elements, sometimes referred to as rooms:

- Customer Requirements or the "What's" derived from VOC and CTQs
- Technical specifications or Design Requirements (the "How's")
- Relationship Matrix (the Roof or "What's" and "How's")
- Prioritized Customer Requirements
- Competitive Assessment
- Interrelationships or Correlation Matrix (the "How's")
- Prioritized Design Requirements

How to Build a House of Quality Step by Step Step 1: Determine Customer Requirements

Identify the important customer requirements. These are the "What's" and are typically determined through the VOC/CTQ process. Use the results from your requirements tree diagram as inputs for the customer requirements in your HOQ.

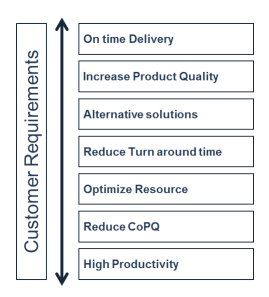


Fig. 1.21 Customer Requirements

Customers drive the development of the product, not the designers. Hence, it is necessary to determine all customer requirements. Customers will provide the requirements in their own language; you will need to categorize the requirements based on importance. General customer requirements are:

- Human factors
- Physical requirements
- Reliability

Step 2: Technical Specifications

- Life-cycle concerns
- Manufacturing requirement

- Potential choices for product features
- Voice of Designers or Engineers
- Each "What" item must be refined to "How's"

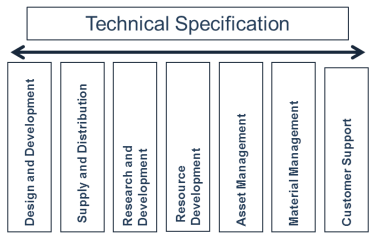


Fig. 1.22 Technical Specifications

The goal is to develop a set of technical specifications from the customer's requirements. Designers and engineers then take the customer requirements and determine technical parameters that need to be designed into the product or process.

Step 3: Develop Relationship Matrix ("What's" and "How's")

This is the center portion of the house. Each cell represents how each technical specification relates to each customer requirement.

	R	elationship between	←	Te	chnica	I Spec	cificati	on	\rightarrow
Т		omer Requirements and chnical Specifications WHATs vs. HOWs) +9 Strong) +3 Medium \ +1 Weak	Design and Development	Supply and Distribution	Research and Development	Resource Development	Asset Management	Material Management	Customer Support
പ്		Ontime Delivery	\odot	0		Ο	Δ	\odot	
ment		Increase Product Quality	Ο	$oldsymbol{O}$		0	Δ	\odot	
Requirements		Alternative solutions	$oldsymbol{O}$	Δ	Δ	$oldsymbol{O}$		0	Ο
		Reduce Turn around time	\triangle	\odot		Ο	$oldsymbol{O}$		$oldsymbol{O}$
Customer		Optimize Resource		Δ					Δ
Cust		Reduce CoPQ		$oldsymbol{O}$			0	Δ	0
Ļ		High Productivity	Δ					Δ	$oldsymbol{O}$

Fig. 1.23 Relationship Matrix

The relationship matrix represents the strength of the relationship between each technical specification and each customer requirement. Use symbolic notations for depicting strong, medium, and weak relationships, such as assigning weights of 1–3–9.

Step 4: Prioritize Customer Requirements

This is the right portion of the house. Each cell represents customer requirements based on relative importance to customers and perceptions of competitive performance.

	Design and Development	Supply and Distribution	Research and Development	Resource Development	Asset Management	Material Management	Customer Support	Customer Preferences
Ontime Delivery	$oldsymbol{O}$	0		0	Δ	$oldsymbol{O}$		3
Increase Product Quality	Ο	\odot		Ο	Δ	\odot		5
Alternative solutions	\odot	Δ	Δ	\odot		Ο	Ο	4
Reduce Turn around time	Δ	$oldsymbol{O}$		0	\odot		$oldsymbol{O}$	2
Optimize Resource		Δ					Δ	3
Reduce CoPQ		$oldsymbol{O}$			Ο	Δ	Ο	4
High Productivity	Δ					Δ	$oldsymbol{eta}$	1

Fig. 1.24 Prioritize Customer Requirements

Prioritize the customer requirements based on importance rating, target value, scale-up factor, sales point.

Step 5: Competitive Assessments

This is the extreme right portion of the house. Comparison of the organization's product to its competitor's products.

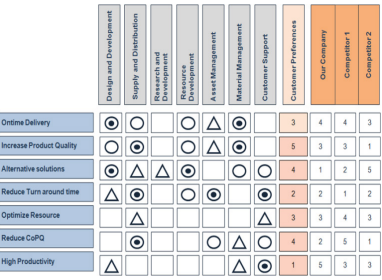


Fig. 1.25 Competitive Assessment

Use the competitive assessment as an opportunity to compare your product/process requirements and specifications with your competitors' products/process. The customer can be used to estimate 1–5 ratings on all products/processes.

This is the top portion of the house. It identifies the correlation between "how" items. Some have support (positive) correlations and other may have conflict or negative correlations.

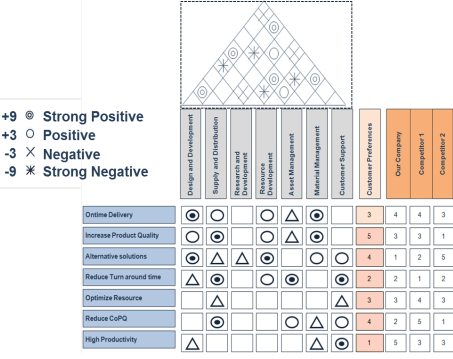


Fig. 1.26 Correlation Matrix

The purpose of the correlation matrix is to identify any inconsistency within the design or technical specifications.

Step 7: Prioritize Design Requirements

- Overall Importance Ratings—Function of relationship ratings and customer prioritization ratings
- Technical Difficulty Assessment—Like customer market competitive evaluations but conducted by the technical team

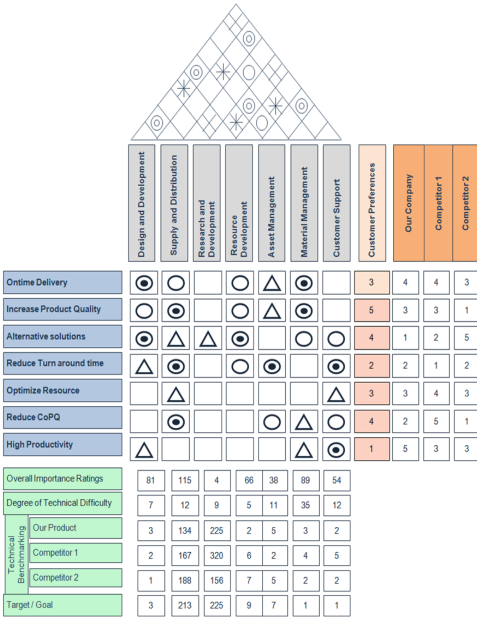


Fig. 1.27 Prioritize Design Requirements

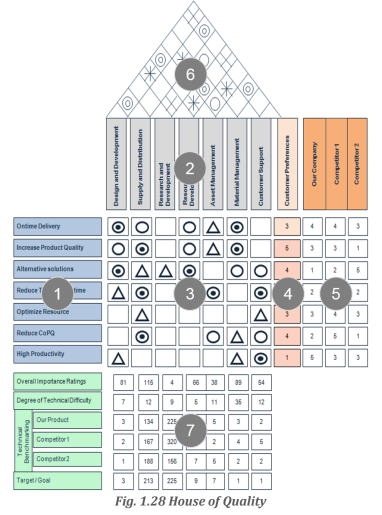
- Overall Importance
- Technical Specification

- Competitive Evaluation
- Target Goals

Each of these prioritization categories helps to establish the feasibility and realization of each "How" item. Target Goals for example can help identify how much is good enough to satisfy the customer. Technical specifications & competitive evaluation should be performed by technical teams. Generally, 1 to 5 ratings should be used. They should be clearly stated in a measurable way.

Figure 1.28 depicts a full House of Quality with each "room" labeled. In summary, the rooms are:

- 1. Customer Requirements
- 2. Technical Specifications
- 3. Relationship Matrix
- 4. Prioritized Customer Requirements
- 5. Competitive Assessment
- 6. Correlation Matrix
- 7. Prioritized Design Requirements



Pros of QFD

- Focuses the design of the product or process on satisfying customer's needs and wants.
- Improves the contact channels between customers, advertising, research and improvement, quality and production departments, which sustains better decision making.
- Reduces the new product development project period and cost.

Cons of QFD

- The relationship matrix can be too obscure with many process inputs and/or many customer constraints.
- It can be very complicated and difficult to implement without experience.

• If throughout the process new ideas, specifications, or requirements are not discovered, you run the risk of losing team members' trust in the process.

QFD Summary

When used properly, the quality function deployment is an extremely valuable approach to product/process design. There are many benefits of QFD that can only be realized when each step of the process is completed thoroughly:

- Logical way of obtaining information and presenting it
- Smallest product development cycle
- Considerably condensed start-up costs
- Fewer engineering alterations
- Reduced chance of supervision during design process
- Collaborating environment
- Preserving everything in characters

1.2.4 Cost of Poor Quality

The *Cost of Poor Quality* (COPQ) is the expense incurred due to waste, inefficiencies, and defects. It is the cost absorbed when a process operates at anything less than optimal: the cost of waste, resource inefficiency, and defects (e.g., repairs, rework, service calls, scrap, warranty claims, and write-offs). COPQ can be staggering when all process inefficiencies are revealed.

Experts have estimated COPQ to be between 5% and 30% of gross sales for most companies. Consider a company that does approximately \$100 million in sales per week, and has a COPQ of 20%. That means that one day out of a five-day work week is completely devoted to producing scrap!

Understanding COPQ and where to look for it will help uncover process inefficiencies, defects, and hidden factories within your business. Consider for instance the hidden factories—Do you have procedures for rework, or workarounds? They might seem "hidden" at first, because they have come to be accepted as the normal process for doing things. There are many different types of non-value added activities that can drive waste.

Cost of Poor Quality

There are seven common forms of waste that are often referred to as the *seven deadly muda*. *Muda* is a Japanese word meaning "futility, uselessness, idleness, superfluity, waste, wastage, wastefulness."

Technically, there are more than seven forms of waste but if you can remember these you will capture over 90% of your waste. The seven deadly muda, as defined by Taiichi Ohno (the Toyota Production System), are:

- 1. Defects
- 2. Overproduction—Producing more than what your customers are demanding
- 3. Over-Processing—Unnecessary time spent (e.g., relying on inspections instead of designing a process to eliminate problems)
- 4. Inventory—Things awaiting further processing or consumption
- 5. Motion—Extra, unnecessary movement of employees
- 6. Transportation—Unnecessary movement of goods
- 7. Waiting—For an upstream process to delivery, for a machine to finish processing, or for an interrupted worker to get back to work

The seven deadly muda are very important to understand. They are the best way to identify the COPQ. The presence of any muda causes many other forms of inefficiencies and hidden factories to manifest themselves. The next section we will discuss how to determine the cost of poor quality related to the seven muda.

COPQ: Costs Related to Production

Costs related to *production* are the direct costs of the presence of muda. These forms of COPQ are usually understood and easily observable. They are in fact the seven deadly muda themselves:

- 1. Defects—The cost of a defect itself (i.e., the lost labor and materials)
- 2. Overproduction—Resources spent producing more of something than what is needed, along with the material cost. This is producing "just-in-case" instead of "just-in-time" and creates excessive lead times, results in high storage costs, and makes it difficult to detect defects. Over-producing hides problems in production and takes a lot of courage to produce only what is needed (which will uncover where the issues are).
- 3. Over-Processing—Resources spent inspecting and fixing instead of getting right the first time
- 4. Inventory—Work in process on the floor is a symptom of over-processing and waiting
- 5. Motion—Unnecessary time spent by employees to perform the process; consider motion like bending, stretching, walking, lifting, and reaching
- 6. Transportation—Adds no value to the product and even increases risk of damage. This can be a hard cost to overcome if facilities are built in a way that requires transportation from one process to another.
- 7. Waiting—Usually because material flow is poor, production runs are too long, and distances between work centers are too long

COPQ: Costs Related to Prevention

Costs related to the *prevention* of muda are those associated with trying to reduce or eliminate any of the seven deadly muda before they occur:

• Costs for error proofing methods or devices

- Costs for process improvement and quality programs
- Costs for training and certifications, etc.

Any costs directly associated with the prevention of waste and defects should be included in the COPQ calculation.

COPQ: Costs Related to Detection

Costs related to the *detection* of muda are those associated with trying to find or observe any of the seven deadly muda after they occur. Any costs directly associated with the detection of waste and defects should be included in the COPQ calculation. Detection is not as effective as prevention, but sometimes is the only option. It can keep additional costs from being incurred, like continuing to process a defective unit. A few detection examples are:

- Costs for sampling
- Costs for quality control check points
- Costs for inspection costs
- Costs for cycle counts or inventory accuracy inspections, etc.

COPQ: Costs Related to Obligation

Costs related to *obligation* are those associated with addressing the muda that reach a customer. Any costs directly associated with customer obligations should be included in the COPQ calculation. Examples include:

- Repair costs
- Warranty costs
- Replacement costs
- Customer returns and customer service overhead, etc.

COPQ: Types of Cost

There are two types of costs to be considered when determining COPQ:

- Hard Costs—Tangible costs that can be traced to the income statement (e.g., costs of resources or materials)
- Soft Costs—Intangible costs: avoidance, opportunity costs, lost revenue, cost avoidance (the difference between what is actually spent and what *could* have been spent had no action been taken; think of this as slowing the rate of cost increases), opportunity cost (value of lost time that could have been used producing something of value)

Calculating the COPQ

- 1. Determine the types of waste that are present in your process.
- 2. Estimate the frequency of waste that occurs.
- 3. Estimate the cost per event, item, or time frame, whichever is appropriate.

4. Multiply the costs.

1.2.5 PARETO CHARTS AND ANALYSIS

Pareto Principle

The *Pareto principle* is commonly known as the "law of the vital few" or "80:20 rule." It means that the majority (approximately 80%) of effects come from a few (approximately 20%) of the causes. It is a very powerful tool for scoping process improvement efforts because it tells us what the "few" causes are that drive most the effects (or in many cases defects).

This principle was first introduced in early 1900s and has been applied as a rule of thumb in various areas. Business-management consultant <u>Joseph M. Juran</u> suggested the principle and named it after Italian economist <u>Vilfredo Pareto</u>, who observed in 1906 that 80% of the land in Italy was owned by 20% of the population; he developed the principle by observing that 20% of the pea pods in his garden contained 80% of the peas. Examples of applying the Pareto principle:

- 80% of the defects of a process come from 20% of the causes
- 80% of sales come from 20% of customers

The Pareto principle helps us to focus on the vital few items that have the most significant impact. In concept, it also helps us to prioritize potential improvement efforts. Since this 80:20 rule was originally based upon the works of Wilfried Fritz Pareto (or Vilfredo Pareto), all Pareto chart and principle references should be capitalized because Pareto refers to a person (proper noun). Mr. Pareto is also credited for many works associated with the 80:20, some more loosely than others: Pareto's Law, Pareto efficiency, Pareto distribution etc.

Pareto Charts

A *Pareto chart* is a chart of descending bars with an ascending cumulative line on the top. *Sum or Count:* The descending bars on a Pareto chart may be set on a scale that represents the total of all bars or relative to the biggest bucket, depending on the software you are using.

Percent to Total: A Pareto chart shows the percentage to the total for individual bars.

Cumulative Percentage: A Pareto chart also shows the cumulative percentage of each additional bar. The data points of all cumulative percentages are connected into an ascending line on the top of all bars. Not all software packages display Pareto charts the same way. Some show the percent of total for each individual bar, and others show the cumulative percentage.

Pareto Charts Case Study

Next, we will use Minitab to run Pareto charts on the same data set. The following table shows the count of defective products by team. Input the tabled data below into your software program and follow the instructions over the next few pages to run Pareto charts.

Count	Category
2	Team 1
12	Team 2
4	Team 3
22	Team 4
2	Team 5
2	Team 6

Create a Pareto Chart in Minitab

Steps to generate a Pareto chart using Minitab:

- 1. Open Minitab and type or copy the data from table 1.2 into the Minitab worksheet.
- 2. Click on Stat \rightarrow Quality Tools \rightarrow Pareto Chart. A new window with the title "Pareto Chart" pops up.
- 3. Select "Category" into the "Defects or attribute data in" box.
- 4. Select "Count" into the box "Frequency in."
- 5. Click "OK."

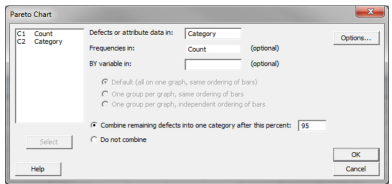


Fig. 1.29 Pareto Chart Dialog Box

The Pareto chart output is created in a newly-generated window. Figure 1.30 depicts the Pareto chart generated by Minitab. It represents the count of defective products by team. The bars are descending on a scale with the peak at 50, which is a rounded approximation of the total count of all defective products for all teams. The table below the chart contains counts, individual percentages, and cumulative percentages. Lastly, cumulative percentages are represented by the data points driving the line that spans across the top of the graphic.

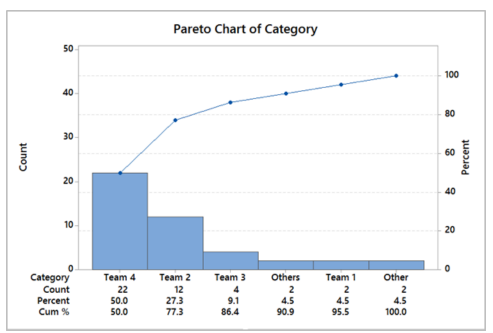


Fig. 1.30 Minitab Pareto Chart Output

With this data and analysis, we were looking to determine how many defective products (i.e. effects) were created by each team (i.e. causes). As you can see, Minitab sorted the data in decreasing order. The team with the most defective products (team 4) was placed on the left. The total number of defects is 44 which you can estimate by glancing at where the cumulative line ends (the cumulative percentage line (red line) shows what each team, cumulatively is contributing as a percentage to the total number of defects). It's also notable that nearly 80% of the defects are contributed by nearly 20% of the causes. Thus, the Pareto principle holds true.

Pareto Analysis

The *Pareto analysis* is used to identify root causes by using multiple Pareto charts. In Pareto analysis, we drill down into the bigger buckets of defects and identify the root causes of defects that contribute heavily to total defects. This drill-down approach often effectively solves a significant portion of the problem. Next you will see an example of three-level Pareto analysis.

- The second-level Pareto is a Pareto chart that is a subset of the tallest bar on the first Pareto. Therefore, the chart will represent categorized defects of Team 4.
- The third-level Pareto will then be a subset of the tallest bar of the second-level Pareto.

Pareto Analysis: First Level

The first-level Pareto was already demonstrated in figure 1.30. It showed the count of defective items by team. In that first level Pareto we concluded that Team 4 accounted for 50% of the total number of defects. The next level Pareto will be a second level and will only show the defective items from Team 4 categorized in a logical manner or in a way that the data allows.

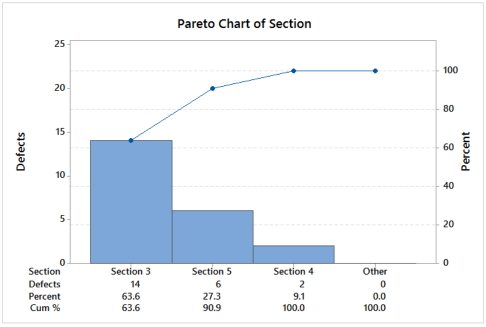


Fig. 1.31 Second Level Pareto (defects by section)

The second-level Pareto shows the count of the defective items by section for only Team 4 (22 defects). You can see that teams are broken down to sections and it appears that section 3 accounts for 63% of the total defects for team 4.

Pareto Analysis: Third Level

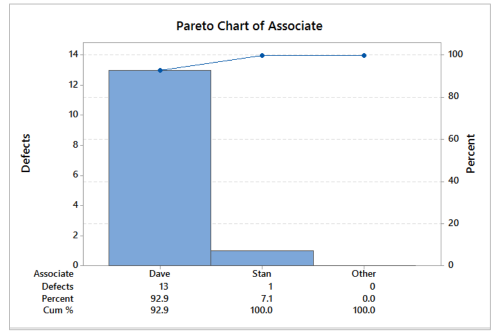


Fig. 1.32 Third Level Pareto (defects by associate)

The third-level Pareto shows the count of defective items by associate for only Section 3 of Team 4. The total defects by associate sum to the total for Section 3 and Dave accounts for 93% of Section 3's defects.

Pareto Analysis: Conclusion

Our analysis clearly shows the root of the problem being associate Dave who has 13 of the 14 defects in Section 3 of Team 4. Through this multi-level Pareto analysis, instead of focusing on all teams to reduce the number of defects, we discovered that we can focus on a single individual who is contributing 13 defects out of a total of 44 which is 30% of the total for all teams. Determining what Dave might be doing differently or wrong and solving that problem could fix about 30% of the entire defective products (13/44). This type of analysis significantly narrowed the focus necessary for reducing defective products.

1.3 SIX SIGMA PROJECTS

1.3.1 SIX SIGMA METRICS

There are many Six Sigma metrics and/or measures of performance used by Six Sigma practitioners. In addition to the ones we will cover here, several others (Sigma level, C_p, C_{pk}, P_p, P_{pk}, takt time, cycle time, utilization etc.) will be covered in other modules throughout this training. The Six Sigma metrics of interest in the *Define* phase are:

- Defects per Unit
- Defects per Million Opportunities
- Yield
- Rolled Throughput Yield

Defects per Unit

DPU, Defects per Unit, is the basis for calculating DPMO and RTY, which we will cover in the next few pages. DPU is found by dividing total defects by total units.

$$DPU = \frac{D}{U}$$

For example, if you have a process step that produces an average of 65 defects for every 598 units, then your DPU = 65/598 = 0.109 or 10.9%.

Defects per Million Opportunities

DPMO, Defects per Million Opportunities, is one of the few important Six Sigma metrics that you should get comfortable with if you are associated with Six Sigma. Remember 3.4 defects per million opportunities? In order to understand DPMO it is best if you first understand both the nomenclature and the nuances such as the difference between defect and defective.

Nomenclature:

- Defects = D
- Unit = U
- Opportunity to have a defect = 0

To properly discuss DPMO, we must first explore the differences between "defects" and "defective."

Defective

Defective suggests that the value or function of the entire unit or product has been compromised. Defective items will always have at least one defect. Typically, however, it takes multiple defects and/or critical defects to cause an item to be defective.

Defect

A defect is an error, mistake, flaw, fault, or some type of imperfection that reduces the value of a product or unit. A single defect may or may not render the product or unit "defective" depending on the specifications of the customer.

To summarize in simple terms, defect means that part of a unit is bad. Defective means that the whole unit is bad. Now let us turn our attention to defining "opportunities" so that we can fully understand Defects per Million Opportunities (DPMO).

Opportunities

Opportunities are the total number of possible defects. Therefore, if a unit has six possible defects, then each unit produced is equal to six defect opportunities. If we produce 100 units, then there are 600 defect opportunities (100 units × 6 opportunities/unit).

Calculating Defects per Million Opportunities

The equation is:

$$DPMO = \frac{D}{U \times O} \times 1,000,000$$

For example, let us assume there are six defect opportunities per unit and there is an average of 4 defects every 100 units.

Opportunities = 6 × 100 = 600 Defect rate = 4/600 DPMO = 4/600 × 1,000,000 = 6,667

What is the reason or significance of 1,000,000? Converting defect rates to a "per million" value becomes necessary when the performance of your process approaches Six Sigma. When this happens, the number of defects shrink to virtually nothing. In fact, if you recall from section "1.1 What is Six Sigma", six sigma is equivalent to 3.4 defects per million opportunities.

By using 1,000,000 opportunities as the barometer we have the resolution in our measurement to count defects all the way up to Six Sigma.

Rolled Throughput Yield

Rolled Throughput Yield (RTY) is a process performance measure that provides insight into the cumulative effects of an entire process. RTY measures the yield for each of several process steps and provides the probability that a unit will come through that process defect-free.

RTY allows us to expose the "hidden factory" by providing visibility into the yield of each process step. This helps us identify the poorest performing process steps and gives us clues into where to look to find the most impactful process improvement opportunities.

Calculating RTY

RTY is found by multiplying the yields of each process step. Let us take the five-step process below and calculate the RTY (the yield or percent of units that get through without a defect) using the multiplication method mentioned above.

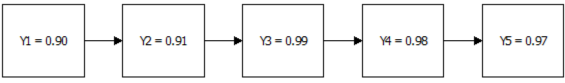


Fig. 1.33 Five Yields for a 5-step process

The RTY calculation for figure 1.33 is: RTY = 0.90 × 0.91 × 0.99 × 0.98 × 0.97 = 0.77

Therefore, RTY = 77%. In this example, 77% of units get through all five process steps without a defect. You may have noticed that to calculate RTY we must determine the yield for each process step. Before we get into calculating yield, there are a few abbreviations that need to be declared.

- Defects = D
- Unit = U
- Defects per Unit = DPU
- Yield = Y
- e = 2.71828 (mathematical constant)

Calculating Yield

The *yield* of a process step is the success rate of that step or the probability that the process step produces no defects. To calculate the yield, we need to know the DPU and then we can apply it to the yield equation below.

$$Y = e^{-dpu}$$

For example, let us assume a process step has a DPU of 0.109 (65/598).

Y = 2.718 ^ -0.109 = 0.8967. Rounded, Y = 90%.

In this example, the DPU (defects per unit) is 0.109 (65 defects/598 units). Plugging in the DPU, and using the mathematical constant for e (2.718), the result is 0.8967, or 90% when rounded. Below in table 1.3, is the above process yield data that we used in the earlier RTY calculation. This table allows us to see the DPU and yield of each step as well as the RTY for the whole process.

Process Step	Defects	Units	DPU	Yield	RTY
1	65	598	0.10870	0.89701	0.90
2	48	533	0.09006	0.91389	0.82
3	5	485	0.01031	0.98974	0.81
4	10	480	0.02083	0.97938	0.79
5	15	471	0.02972	0.97072	0.77

Table 1.3 DPU, Yield and RTY from figure 1.33

The rightmost column of table 1.3 shows the RTY cumulated through all the process steps. Which steps seem to be the greatest opportunities? Notice the yield in steps 1 and 2 are the lowest.

Estimating Yield*

Instead of using the equation provided earlier, there is a simpler way to calculate yield using *yield estimation:* It is possible to "estimate" yield by taking the inverse of DPU or simply subtracting DPU from 1.

Yield Estimation = 1 – DPU

Consider the DPU's in table 1.3 and follow the 1-DPU method for estimating yield. You should have something similar to the following equations:

- Yield Estimate for process step 1: 1 0.10870 = 0.90
- Yield Estimate for process step 2: 1 0.09006 = 0.91
- Yield Estimate for process step 3: 1 0.01031 = 0.99
- Yield Estimate for process step 4: 1 0.02083 = 0.98
- Yield Estimate for process step 5: 1 0.02972 = 0.97

Now, let's calculate RTY using the Yield Estimation Method:

 $RTY = 0.90 \times 0.91 \times 0.99 \times 0.98 \times 0.97 = 0.77 = 77\%$

Assuming the maximum DPU is 1^{*}, so to get a yield just subtract DPU from 1. As you can see, by calculating yield this way we get the same result for RTY. The previous method's calculated result was also 77%. This may not be the most exact approach but for business professionals who make decisions based on sound analysis, the estimation method will typically suffice.

*This method assumes max DPU is 1. Although this is not technically true, in practice its rare to find processes with DPU greater than 1. If it is the case this method will not work.

1.3.2 BUSINESS CASE AND CHARTER

Earlier we stated that DMAIC is a structured and rigorous methodology designed to be repeatedly applied to *any* process in order to achieve Six Sigma. We also stated that DMAIC was a methodology that refers to five phases of a project: Define, Measure, Analyze, Improve, and Control.

Given that the premise of the DMAIC methodology is project-based, we must take the necessary steps to define and initiate a project, hence the need for project charters. The project charter is what sets direction for a project.

Project Charter

The purpose of a *project charter* is to provide vital information about a project in a quick and easy-to-comprehend manner. It is a contract of sorts between project champions, sponsors, stakeholders, and the project team. It is a quick reference to tell the story of what, why, who, when, etc. To set the *right* direction for a project, it is critical to define certain guardrails to ensure the project stays on track and is successful. Project charters are used to get approval and buy-in for projects and initiatives as well as declaring:

- Scope of work
- Project teams
- Decision authorities

- Project lead
- Success measures

The key elements of project charters are:

- Title
- Project Lead
- Business Case
- Problem Statement
- Project Objective
- Primary Metric
- Secondary Metrics
- Project Scope

- Project Timeline
- Project Constraints
- Project Team
- Stakeholders
- Approvers
- Constraints
- Dependencies
- Risk

				Project Title:	593.	114	-46 A
Proje	ect Char	rter		Black Belt	Project Champion	Executive Sponsor	MBB/Mentor
Primary N	Metric				Secondary Metric		
Problem S	Statement				Business Case		
	el Project	Timeline	Constr	aints & Dependencies	Project Risks	Other Dia	agnostics
Phase	el Project Start	Timeline Finish	Constr	aints & Dependencies	Project Risks	Other Dia	agnostics
Phase Define			Constr	aints & Dependencies	Project Risks	Other Dia	agnostics
Phase Define Measure			Constr	aints & Dependencies	Project Risks	Other Dia	agnostics
Phase Define Measure Analyze			Constr	aints & Dependencies	Project Risks	Other Dia	agnostics
Phase Define Measure Analyze Improve			Constr	aints & Dependencies	Project Risks	Other Dia	agnostics
Phase Define Measure Analyze Improve Control	Start	Finish	Constr				
Phase Define Measure Analyze Improve Control Approval	Start	Finish		Stakeholders & Ad	visors	Project Team & SME	i's
Phase Define Measure Analyze Improve Control	Start	Finish					
Phase Define Measure Analyze Improve Control Approval	Start	Finish		Stakeholders & Ad	visors	Project Team & SME	i's
Phase Define Measure Analyze Improve Control Approval	Start	Finish		Stakeholders & Ad	visors	Project Team & SME	i's
Phase Define Measure Analyze Improve Control Approval	Start	Finish		Stakeholders & Ad	visors	Project Team & SME	i's
Phase Define Measure Analyze Improve Control Approval	Start	Finish		Stakeholders & Ad	visors	Project Team & SME	i's

Fig. 1.34 Project Charter Template

Figure 1.34 above shows an example of a project charter with all the key elements. Because of its formatting, it's easy to share, easy to read, and is useful for quickly communicating to people who may not be familiar with the project.

Project Charter: Key Elements

Title: Projects should have a name, title, or some reference that identifies them. Branding can be an important ingredient in the success of a project so be sure your project has a reference name or title. This is how your project will be known to others, and keeping it top of mind is obviously part of ensuring its success.

Leader: All projects need a declared leader or someone who is responsible for project's **RACI** stands for **R**esponsible, **A**ccountable, **C**onsulted, and Informed. RACI identifies the people that play those roles. Every project must have declared leaders indicating who is responsible and who is accountable.

Business Case: A business case is the quantifiable reason why the project is important. It is important for the business case to be quantifiable. Stakeholders have to know what success looks like. Cost of poor quality, as described earlier, is a great approach to quantifying the business case.

Business cases help shed light on problems. They explain why a business should care. A business case clearly articulates why it is necessary to do the project and what is the benefit to the customer, to employees, or to the shareholders.

A problem by itself is not enough to articulate why a project needs to be done. The business case turns the problem into the reason a business should care: What is the cost of the problem, what is the impact to customers, or what is the lost revenue opportunity? Business cases must be quantified and stated succinctly. COPQ is a key method of quantification for any business case.

Problem Statement and Objective: A properly written problem statement has an objective statement woven into it. There should be no question as to the current state or the goal.

A gap should be declared, the gap being the difference between the present state and the goal state. The project objective should be to close the gap or reduce the gap by some reasonable amount. Valuation or COPQ is the monetary value assigned to the gap.

Lastly, a well-written problem statement refers to a timeline expected to be met. A wellwritten problem statement includes all of the following:

- Declaration of the current state of a problem in terms of a given measure
- A goal for the measure, which establishes a gap from the current state
- An objective that states how much of the gap the project aims to close
- The monetary value of the objective (cost of poor quality)
- A timeline that the project will meet

Project Charter: Problem Statement Examples

Currently, process defect rates are 17% with a goal of 2%. This represents a gap of 15%, costing the business \$7.4 million dollars. The goal of this project is to reduce this gap by 50% before November 2010 putting process defect rates at 9.5% and saving \$3.7MM.

Process cycle time has averaged 64 minutes since Q1 2009. However, production requirements put the cycle time goals at 48 minutes. This 16-minute gap is estimated to cost the business \$296,000. The goal of this project is to reduce cycle time by 16 minutes by Q4 2010 and capture all \$296,000 cost savings. Can you identify the key elements of a problem statement?

Metrics: A measure of success is an absolute for any project.

- Metrics give clarity to the purpose of the work.
- Metrics establish how the initiative will be judged.
- Metrics establish a baseline or starting point.

For all Six Sigma projects, metrics are mandatory! How does one know if a project does what it sets out to do? It is critical that *every* project choose the right metrics to determine success. Project metrics are the center point for the work to ensure that the direction and decisions made through the course of the project are based on data. The metrics are what stakeholders will judge the success of the project on.

Primary Metric: The *primary metric* is a generic term for a Six Sigma project's most important measure of success. Choosing the primary metric should not be taken lightly. A Six Sigma expert should play a significant role in determining the primary metric to ensure it meets the characteristics of a good one. The primary metric is defined by the BB, GB, MBB, or Champion.

A primary metric is an absolute must for any project and it should not be taken lightly. Here are a few characteristics of good primary metrics. Primary metrics should be:

- Tied to the problem statement to ensure that what is being measured is a direct indication of whether the problem is being solved or not
- Measurable
- Expressed with an equation, meaning that the primary metric, Y, can be expressed as a function of the x's
- Aligned to business objectives, meaning that the primary metric should be correlated to business results
- Tracked at the proper frequency (hourly, daily, weekly, monthly etc.); in other words, they should be something that can be tracked frequently enough so that action can be taken quickly when the metric change
- Expressed pictorially over time with a run chart, time series, or control chart
- Validated with an MSA

The primary metric is the reason for your work, the success indicator, and your beacon. The primary metric is of utmost importance and should be improved, *but* not at the expense of your secondary metric. Do not trade one problem for another. Solutions need to be balanced. Be sure that you do not create an issue somewhere else when you implement a solution to improve your primary metric. That is why *secondary metrics* are important.

Secondary Metric: The *secondary metric* is the thing you do not want sacrificed on behalf of a primary improvement. A secondary metric is one that makes sure problems are not just "changing forms" or "moving around." The secondary metric keeps us honest and ensures we are not sacrificing too much for our primary metric.

If your primary metric is a cost or speed metric, then your *secondary metric* should probably be some quality measure. For example, if you were accountable for saving energy in an office building and your primary metric was energy consumption then you *could* shut off all the lights and the HVAC system and save tons of energy . . . except that your secondary metrics are probably comfort and functionality of the work environment.

When you think about your primary metric, think about the really easy and obvious ways to improve the primary metric. Chances are that you can easily improve it by sacrificing something else. The elements of a good project charters include:

- Scope Statement—Defined by high-level process map
- Stakeholders Identified—Who is affected by the project

- Approval Authorities Identified—Who makes the final call
- Review Committees Defined—Who is on the review team
- Risks and Dependencies Highlighted—Identify risks and critical path items
- Project Team Declared—Declare team members
- Project Timeline Estimated—Set high-level timeline expectations

1.3.3 Project Team Selection

Six Sigma project team selection is the cornerstone of a successful Six Sigma project. Team selection is obviously not something that is unique to a Six Sigma project. Team selection is important in any kind of project. You want to choose a group of people that complement each other, share similar goals and objectives, and are not afraid to hold each other accountable for achieving the project's objectives.

Teams and Team Success

A *team* is a group of people who share complementary skills and experience.

- A team will be dedicated to consistent objectives.
- Winning teams share similar and coordinated goals.
- Teams often execute common methods or approaches.
- Team members hold each other accountable for achieving shared goals.

What Makes a Team Successful?

There should not be any surprises about what makes a team successful. These characteristics are all important for a team to be successful:

- Shared goals
- Commitment
- Leadership
- Respect

- Effective communication
- Autonomy
- Diverse knowledge & skills
- Adequate resources

The keys to team success are:

- Agreed focus on the goal or the problem at hand
 - Focus on problems that have meaning to the business
 - Focus on solvable problems within the scope of influence; a successful team does not seek unattainable solutions
- Team Selection
 - Selected teammates have proper skills and knowledge
 - Adequately engaged management
 - Appropriate support and guidance from their direct leader
- Successful teams use reliable methods

- Follow the prescribed DMAIC methodology
- Manage data, information, and statistical evidence
- Successful teams always have "exceeds" rated players. Winning teams typically have unusually high standards
 - Have greater expectations of themselves and each other
 - Do not settle for average or even above average results

Teams should be able to agree on what the focus is. The team members all need to understand how the problem translates to something meaningful for the business. They also should be realistic about what they can and should seek to accomplish.

The people that are selected need to have "skin in the game." Not only do they need to be knowledgeable and have the needed skills, but they (and their management) need to be motivated to seek the solution.

The team should use the facts and data to lead the way. Be disciplined to follow the DMAIC approach and be aware of when a gut feeling or emotion is driving their behavior. Of course, teams should be built from the top performers whenever possible.

Principles of Team Selection

- Select team members based on:
 - Skills required to achieve the objective
 - Experience (subject matter expertise)
 - Availability and willingness to participate
 - Team size (usually four to eight members)
 - Don't go at it alone!
 - Don't get too many cooks in the kitchen!
 - Members' ability to navigate
 - \circ The process
 - \circ The company
 - The political landscape
- Be sure to consider the inputs of others
 - Heed advice
 - Seek guidance

A team member should be selected based on the skills and subject matter expertise. For example, if you are focusing on a process improvement, find a top performing employee that performs the process on a regular basis. A team member needs to have capacity to participate on a project. Therefore, it is important to pick people with shared goals. If they have a shared goal, it is likely that they can find the time. The team size needs to be optimal. Include enough people to generate consensus, but not too many people because they can never reach consensus, and this can paralyze a project. Be open minded, listen to others, be inclusive.

Project Team Development

All teams experience the following four stages of development. It is helpful to understand these phases so that you can anticipate what your team is going to experience. The four stages of team development process are:

- 1. Forming
- 2. Storming
- 3. Norming
- 4. Performing

Teammates seek something different at each stage:

- 1. In the forming stage they seek inclusion
- 2. In the storming stage they seek direction and guidance
- 3. In the norming stage they seek agreement
- 4. In the performing stage they seek results

Patterns of a team in the Forming stage:

- Roles and responsibilities are unclear
- Process and procedures are ignored
- Scope and parameter setting is loosely attempted
- Discussions are vague and frustrating
- There is a high dependence on leadership for guidance

In the Forming stage, people are trying to figure out where they fit in; the team is trying to figure out where the social boundaries are with each other; they are all trying to "get to know" each other.

Patterns of a team in the Storming stage:

- Attempts to skip the research and jump to solutions
- Impatience for some team members regarding lack of progress
- Arguments about decisions and actions of the team
- Team members establish their position
- Subgroups or small teams form
- Power struggles exist and resistance is present

After the delay of the forming stage, urgency can set in and they start to seek progress. Team members even seek progress by sacrificing the discipline. Patterns of a team in the Norming stage are:

- Agreement and consensus start to form
- Roles and responsibilities are accepted
- Team members' engagement increases
- Social relationships begin to form
- The leader becomes more enabling and shares authority

In the Norming stage the team is settling into place and into roles with responsibilities. Once roles are clear, engagement increases with comfort level, and the team starts to form relationships with each other. Patterns of a team in the Performing stage are:

- Team is directionally aware and agrees on objectives
- Team is autonomous
- Disagreements are resolved within the team
- Team forms above average expectations of performance

In the Performing stage, the team is aligned and directed. The members are autonomous in that they can resolve issues within and stay on the path to progress. The team is united to exceed expectations of performance.

Well-structured and energized project teams are the essential components of any successful Six Sigma project. To have better chances of executing the project successfully, you will need to understand and effectively manage the team development process. Even with the most perfectly defined project, a well-structured team can be the difference between success and failure.

1.3.4 Project Risk Management

Risk

Risk is defined as a future event that *can* impact the task/project if it occurs. A broad definition of project risk is an uncertainty that can have a negative *or* positive effect on meeting project objectives. Positive risks are risks that result in good things happening—sometimes called opportunities.

What is Project Risk Management?

The main purpose of *risk management* is to foresee potential risks that may inhibit the project deliverables from being delivered on time, within budget, and at the appropriate level of quality, and then to mitigate these risks by creating, implementing, and monitoring

contingency plans. Risk management is concerned with identifying, assessing, and monitoring project risks before they develop into issues and impact the project.

Risk analysis helps to identify and manage potential problems that could impact key business initiatives or project goals.

Three Basic Parameters of Risk Analysis

- 1. Risk Assessment: The process of identifying and evaluating risks, whether in absolute or relative terms
- 2. Risk Management: Project risk management is the effort of responding to risks throughout the life of a project and in the interest of meeting project goals and objectives
- 3. Risk Communication: Communication plays a vital role in the risk analysis process because it leads to a good understanding of risk assessment and management decisions

Why is Risk Analysis Necessary?

What can happen if you omit the risk analysis?

- Vulnerabilities cannot be detected
- Mitigation plans are introduced without proper justification
- Customer dissatisfaction
- Not meeting project goals
- Remake the whole system
- Huge cost and time loss

Project Risk Analysis Steps

The project risk analysis process consists of the following steps that evolve through the life cycle of a project.

- 1. Risk Identification: Identify risks and risk categories, group risks, and define ownership.
- 2. Risk Assessment: Evaluate and estimate the possible impacts and interactions of risks.
- 3. Response Planning: Define mitigation and reaction plans
- 4. Mitigation Actions: Implement action plans and integrate them into the project
- 5. Tracking and Reporting: Provide visibility to all risks
- 6. Closing: Close the identified risk

1. Risk Identification

The first action of risk management is the identification of individual events that the project may encounter during its lifecycle. The identification step comprises:

- Identify the risks
- Categorize the risks

- Match the identified risks to categories
- Define ownership for managing the risks

The process of identification, matching, and assigning ownership may involve the full project team in a brainstorming exercise or a workshop. The team should be encouraged to participate in risk identification by actively reporting risks as they arise, and to make suggestions as to the controls needed to rectify the situation.

Source of Risk:

Identification of risk sources provides a basis for systematically examining changing situations over time to uncover circumstances that impact the ability of the project to meet its objectives.

Establishing categories for risks provides a mechanism for collecting and organizing risks as well as ensuring appropriate scrutiny and management attention for those risks that can have more serious consequences on meeting project objectives.

Source of Risk	Description					
Human Resources	The risks originated from human resources (e.g. resource availability, skills, training etc.)					
Physical Resources	The risks originated from physical resources (e.g., hardware or software, availability of the required number at the right time etc.)					
Technology	The risks originated from technology (e.g., development environment, new or complex technologies, performance requirements, tools etc.)					
Suppliers	The risks are associated with a supplier (e.g., delays in supplies, capability of suppliers etc.)					
Customer	The risks derived from the customer (e.g., unclear requirements, requirement volatility, change in project scope, delays in response etc.)					
Security	The risks are associated with information security, security of personnel, security of assets, and security of intellectual property					
Legal	The risks are associated with legal issues that may impact the project					
Project Management	The risks are associated with project management processes, organizational maturity etc.					

Table 1.4 Risk Identification (sources of risk)

The table above shows some examples of source of risks, from those associated with human resources to those related to project management.

Risk Parameters:

Parameters for evaluating, categorizing, and prioritizing risks include the following:

• Risk likelihood (i.e., probability of risk occurrence)

- Risk consequence (i.e., impact and severity of risk occurrence)
- Thresholds to trigger management activities

Risk parameters are used to provide common and consistent criteria for comparing the various risks to be managed and to prioritize the necessary actions required for risk mitigation planning.

2. Risk Assessment

The *risk assessment* consists of evaluating the range of possible impacts should the risk occur.

Follow these steps when assessing risks:

- Define the various impacts of each risk
- Rate each impact based on a logical severity level
- Sort and evaluate risks by severity level
- Determine if any controls already exist
- Define potential mitigation actions

We must prioritize and act on risks in order of priority. In reality, we may be able to fix simple project risks immediately, so we must also use our common sense when prioritizing. Prioritizing is good practice when we need to allocate resources and keep records of the process. The project core team must review *all* risks to ensure the full impact of risk on a project has been identified and estimated.

3. Risk Mitigation Planning

The risk owners are responsible for planning and implementing mitigation actions with support from the project team.

All team members, inclusive of partners and suppliers, may be requested to identify and develop mitigation measures for identified risks. The project core team members are responsible for identifying an appropriate action owner for each identified risk.

After mitigation actions are defined, the project core team will review the actions. The risk owner must track all mitigation actions and expected completion dates. The risk owner and the project core team members must hold all action owners accountable for the risk mitigation planning.

Documenting the process allows us to systematically address the risk that exists in the project. Once the process is documented, action plans can be formulated and responsibilities allocated for controls that need to be implemented to eliminate or reduce the risk of those hazards.

An action plan for each identified risk can include multiple actions involving the team members. In case of risk occurrence, or risk triggers, the risk owner reports to the project manager and implements the response plans.

4. Risk Mitigation Action Implementation

The *action implementation* is the responsibility of the risk owner. The action owners are responsible for the execution of the tasks or activities necessary to complete the mitigation action and eliminate or minimize the risk. The risk owner or the project manager will monitor completion dates of the mitigation action implementation.

Risk Occurrence and Contingency Plans

Whenever any risk occurs, the project team should implement *contingency plans* to ensure that project deliverables can be met. The details of each occurrence should be recorded in the risk register or other tracking tool.

The *risk register* or *risk management plan* will be maintained by the project manager and reviewed regularly. The risk register is a document that contains the results of various risk management processes. It is often displayed in a table or spreadsheet format.

5. Risk Tracking and Reporting

Risk tracking and reporting provides critical visibility to all risks. Risk owners must report on the status of their mitigation actions.

The most efficient method to track and report risk is to do it as an integral part of project team meetings through all stages in the project lifecycle. Depending on the risk severity, project managers need to report the risk status of each category of risk to senior management in the form of dashboards or through weekly status reports.

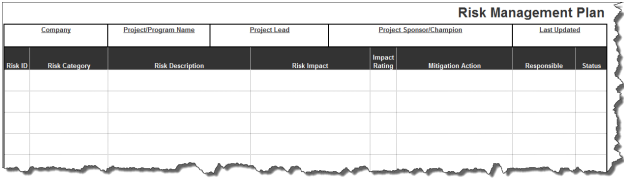


Fig. 1.35 Risk Management Plan Example

6. Risk Closure

The risk owners are responsible for recommending the risk closure to the project manager.

A risk is *closed* only when the item is not considered a risk to the project anymore. When a risk is closed, the project manager needs to update the risk status in the lessons learned document.

Risk Analysis Features

The risk analysis should be:

• Systematic

• Comprehensive

- Data driven
- Adherent to evidence
- Logically sound

Project Risk Analysis Advantages

The risk analysis:

- Aids strategic & business planning
- Meets customer requirements
- Reduces schedule delays
- Minimizes cost overruns
- Promotes effective use of resources

- Practically acceptable
- Open to critique
- Easy to understand
- Promotes continuous improvement
- Helps to achieve project goals
- Prevents surprises
- Uncovers new opportunities
- Enhances communication

1.3.5 PROJECT PLANNING

What is Project Management?

Project management is the process of defining, planning, organizing, managing, leading, securing, and optimizing the resources to achieve a set of planned goals and objectives. It is the application of knowledge, skills, tools, and techniques to project activities in order to meet project requirements.

Project management involves understanding project needs, planning those needs, and properly allocating resources to achieve the desired results.

The basic steps of project management as we will cover in greater detail later are: initiating the project, planning the project, executing the project, and closing the project.

What is a Project Plan?

A *project plan* is a crucial step in project management for achieving a project's goals. A project plan is a formal approved document used to guide and execute project tasks. It provides an overall framework for managing project tasks, schedules, and costs. Project plans are coordinating tools and communication devices that help teams, contractors, customers, and organizations define the crucial aspects of a project or program. Project planning involves defining clear, distinct tasks and deliverables, and the work needed to complete each task and phase of the project.

Project Planning Stages

The stages of project planning are:

• Determine project scope and objectives: Explore opportunities, identify and prioritize needs, consider project solutions

- Plan the project: Identify input and resources requirements such as human resources, materials, software, hardware, and budgets.
- Prepare the project proposal: Based on stakeholder feedback, plan the necessary resources, timeline, budget etc.
- Implement the project: Implement the project by engaging responsible resources and parties. Ensure execution and compliance of the defined plans.
- Evaluate the project: Regularly review progress and results. Measure the project's effectiveness against *quantifiable* requirements

The five key stages of project planning are:

- 1. Determining project scope and objectives
- 2. Project planning
- 3. Project proposal and approval
- 4. Project implementation
- 5. Project evaluation

Planning and Scheduling Objectives

The goals of planning and scheduling are:

- To optimize the use of resources (both human and other resources)
- To increase productivity
- To achieve desired schedules and deliverables
- To establish an approach to minimize long-term maintenance costs
- To minimize the chaos and productivity losses resulting from planned production schedules, priority changes, and non-availability of resources
- To assess current needs and future challenges

Proper project planning is necessary to coordinate all resources and tasks.

Project Planning Activities

Project planning activities involve the creation of:

- Statement of work
- Work breakdown structure
- Resource estimation plan

- Budget or financial plan
- Communication plan
- Risk management plan

• Project schedule

Project Planning Activities: Statement of Work

Statement of Work (SOW) is a formal document often accompanying a contract that outlines specific expectations, limitations, resources and work guidelines intended to define the work or project.

SOWs:

- Define the scope of the project
- Establish expectations and parameters of the project
- Identify technical requirements for the project
- Provide guidance on materials to be used
- Establish timeline expectations

SOWs or statements of work are important "contracts" that outline many critical elements of a project or body of work. SOWs clarify expectations, deliverables, timelines, budgets, and responsibilities.

Project Planning Activities: Work Breakdown Structure

Work Breakdown Structure (WBS) is a decomposition of project components into small and logical bodies of work or tasks.

WBSs:

- Identify all required components of a project
- Cascades components into sub-components and tasks

WBSs are not by themselves project plans or schedules but they are a necessary step to help establish project plans and timelines. WBSs also enable logical reporting and summarizations of project progress.

Level 1	Level 2	Level 3						
1.0 Bicycle	1.1 Frame	1.1.1 Set Body Frame						
		1.1.2 Assemble Handlebars						
		1.1.3 Install Seat & Seat Post						
		1.1.4 Assemble Wheel Bearings & Axle						
		1.1.5 Install Wheels						
		1.1.6 Attach Brake System to Frame Set						
	1.2 Wheels	1.2.1 Stage Rims						
		1.2.2 Install Spokes						
		1.2.3 Insert Tubes in Tires						
		1.2.4 Assemble Tires on Rims						
		1.2.5 Install Reflectors						
	1.3 Brake System	1.3.1 Connect Brake Cables To Hand Levers						
		1.3.2 Connect Brake Cables to Brake Harness						
		1.3.3 Attach Brake Pads to Brake Harness						
		1.3.4 Adjust/Calibrate Brake System						

Fig. 1.36 Work Breakdown Structure Example

A WBS is an important parameter in project planning because projects must inevitably be divided into smaller, more manageable tasks and subtasks. These breakdowns must be aligned in a logical manner to meet customer defined target dates. The WBS is required to prepare a project schedule and/or Gantt chart as well as to identify project milestones.

Project Planning Activities: Resource Planning

Resource Estimation Plan

- Estimate Resource Requirements (Use Your WBS)
- Parts, Hardware, Software, Human Resources etc.
- Plan resources
- Establish who is responsible for what and when
- Determine quantity requirements and delivery dates etc.

Level 3	Parts Required	Quantity	Cost	Part Inventory	Responsible
1.1.1 Set Body Frame	Body Frame	1	\$9.00	4	John
1.1.2 Assemble Handlebars	Handlebars	1	\$4.00	4	John
1.1.3 Install Seat & Seat Post	Seat & Post	1 Each	\$3.00	4 Each	John
1.1.4 Assemble Wheel Bearings & Axle	Bearing & Axle	2 Each	\$1.00	8 Each	John
1.1.5 Install Wheels	Wheels	2	\$0.50	8	John
1.1.6 Attach Brake System to Frame Set					John
1.2.1 Stage Rims	Wheel Rim	2	\$0.30	12	Cathy
1.2.2 Install Spokes	Spokes	48	\$0.03	72	Cathy
1.2.3 Insert Tubes in Tires	Tubes & Tires	2 Each	\$0.33	0 Tubes 8 Tires	Cathy
1.2.4 Assemble Tires on Rims					Cathy
1.2.5 Install Reflectors	Reflectors	4	\$0.10	18	Cathy
1.3.1 Connect Brake Cables To Hand Levers	Cables & Levers	2 Each	\$0.90	12 Each	Lisa
1.3.2 Connect Brake Cables to Brake Harness	Brake Harness	2	\$0.40	8	Lisa
1.3.3 Attach Brake Pads to Brake Harness	Brake Pads	4	\$0.15	12	Lisa
1.3.4 Adjust/Calibrate Brake System					Lisa

Fig. 1.37 Resource Estimation Plan Example

The estimation of resource requirements (human, software, hardware, materials etc.) is necessary for each task and is useful for the creation of a project plan. Use your work breakdown structure to boil it down to tasks so that you can determine specific resource requirements, dependencies etc.

Project Planning Activities: Project Scheduling Project Scheduling

- Assign beginning times to each activity in the WBS (days are used for start and durations times in example below)
- Assign duration times to each activity in the WBS
- Identify People Responsible and set completion dates
- Represent schedules as Gantt charts or network diagrams
- Identify critical dependencies between tasks

Level 2	Level 3	Task Start	Task Duration	Responsible
1.1 Frame	1.1.1 Set Body Frame	0	2	John
	1.1.2 Assemble Handlebars	2	2	John
	1.1.3 Install Seat & Seat Post	4	2	John
	1.1.4 Assemble Wheel Bearings & Axle	6	4	John
	1.1.5 Install Wheels	14	4	John
	1.1.6 Attach Brake System to Frame Set	18	3	John
1.2 Wheels	1.2.1 Stage Rims	0	1	Cathy
	1.2.2 Install Spokes	1	7	Cathy
	1.2.3 Insert Tubes in Tires	10	2	Cathy
	1.2.4 Assemble Tires on Rims	12	2	Cathy
	1.2.5 Install Reflectors	8	2	Cathy
1.3 Brake System	1.3.1 Connect Brake Cables To Hand Levers	0	8	Lisa
	1.3.2 Connect Brake Cables to Brake Harness	8	8	Lisa
	1.3.3 Attach Brake Pads to Brake Harness	16	2	Lisa
	1.3.4 Adjust/Calibrate Brake System	21	2	Lisa

Fig. 1.38 Project Scheduling Example

Scheduling is critical because without it you will miss deadlines and poorly organize dependent tasks and activities. The result of the lack of scheduling, or of poor scheduling, can be catastrophic to a project.

Project Planning Activities: Project Scheduling Project Schedule – Gantt Chart

The advantage of a Gantt chart is its ability to display the status of each task/activity at a glance. Because it is a graphic representation, it is easy to demonstrate the schedule of tasks and timelines to all the stakeholders.

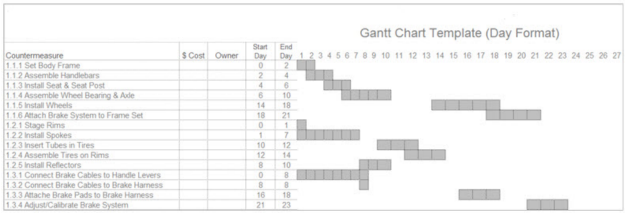


Fig. 1.39 Gantt Chart Example

A Gantt chart offers visibility to a project plan. It displays the tasks of the project plan in a simple and well-organized manner.

There are many tools available to create Gantt charts. Some of the most common and compatible for businesses are Microsoft Project, Visio, and Excel. In Microsoft Excel, there are no native Gantt chart templates so we have established a simple one to use with our bicycle example (shown in fig. 1.39). This template is available for free on our website under the "Tools" link.

Project Scheduling: Critical Path Method Critical Path Method (CPM)

Critical path method (CPM) is a project modeling technique used to identify the set of activities that are most influential to a project's completion timeline. Critical path tasks are those that others are dependent upon.

Project timelines cannot be shortened without shortening the tasks or activities that are identified as critical path items.

Steps to Using the Critical Path Method

- 1. List all activities necessary to complete the project.
- 2. Determine the time or duration of each activity.
- 3. Identify the dependencies between the activities.

CPM is fundamental to project management. You must understand all required project tasks, their durations and dependencies. The critical path method can help you organize and understand how each task behaves relative to others and how the collection of all tasks fit in the grand scheme of a project.

Critical Path Method Example: Let's continue with our bicycle example referencing fig. 1.39. Note that the tasks in the WBS are already set to start based on their dependencies. For example, installing the wheels in step 1.1.5 requires the whole set of steps in 1.2 to be completed so 1.1.5 starts on day 14 when the wheels are ready.

By using your WBS, you can efficiently evaluate task durations and dependencies to find the critical tasks that will affect the final timeline of your project.

Continuing with this example; the last task, "1.3.4 Adjusting the Brake System" completes on day 23 and cannot begin until day 21 because the brake system must be attached to the bike frame before being calibrated. The reason for such a delay can be found in the earlier example. You should have noticed in the resource plan that tire tubes stock was depleted.

Let us assume the tubes take 21 days to receive from the supplier. Below is the adjustment to our Gantt chart. Notice that the project timeline is pushed out by ten days, finishing on day 33 instead of day 23.

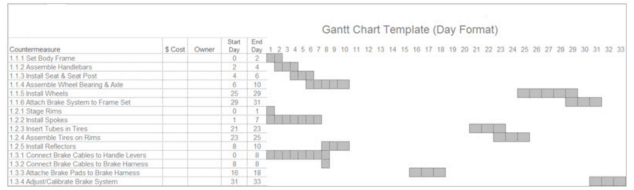


Fig. 1.40 Gantt Chart Revision (21 days for Tubes)

By coordinating the use of all project planning tools and methods covered this far (WBS, Resource Planning, Gantt Chart, CPM) you can not only depict the project tasks and timelines but you can also better manage resources as a result. The Tubes example demonstrates that there is nearly a full FTE resource wasted in waiting time. By knowing this you might be able to re-organize your projects tasks and resources to save on the cost of labor.

Project Scheduling: PERT

Program Evaluation and Review Technique (PERT)

The program evaluation and review technique (PERT) is a method of evaluation that can be applied to time or cost. PERT provides a weighted assessment of time or cost.

PERT uses three parameters for estimation:

- 1. Optimistic or Best Case Scenario represented by "O"
- 2. Pessimistic or Worst Case Scenario represented by "P"
- 3. Most Likely Scenario represented by "ML"

The PERT equation is:

$$E = \frac{O + 4ML + P}{6}$$

Where: E = estimate (of time or cost).

The PERT equation provides for heavier weighting of the most likely scenario but also considers the best and worst cases. In the event that a best or worst case scenario is an extreme situation, the PERT will account for it in a weighted manner.

Using the PERT formula, it is the sum of the estimates of the best case, the worst case, and four times the most likely case, all divided by six.

Let us apply the PERT formula to our estimates of receiving our needed tire tubes from the supplier and then change our schedule based on the result. Your experience with this supplier tells you that they typically overestimate the time required to deliver. Therefore, their 21 day lead time on the tubes should be considered a "worst case scenario."

Procurement has indicated that tubes have arrived in as few as six days from this supplier. So, your best-case scenario is six days. The most likely scenario you decide will be the median of this supplier's delivery time, which is 10 days.

Therefore:

$$E = \frac{0 + 4ML + P}{6}$$
$$E = \frac{6 + 4 \times 10 + 21}{6}$$
$$E = \frac{6 + 40 + 21}{6}$$
$$E = \frac{67}{6}$$
$$E = 11$$

Using the PERT formula in this example: add the best-case scenario (6) to "four times the most likely scenario" (4*10) plus the worst case scenario (21). The result is sixty-seven. We then divide 67 by 6, which yields 11 (rounded). Eleven days is your adjusted estimate of delivery for the tire tubes. Using 11 days in your project plan instead of 21 gives you a more confident estimate of time and removes tire tubes as the primary detracting critical path item.

Project Planning Activities: Budgeting or Financial Planning

Create a preliminary project budget that outlines the planned expenses and revenues pertaining to the project. Preliminary budgets are commonly used for project justification and future business forecasts.

Most organizations expect to see some financial analysis regarding the payback of a project. IRR (Internal Rate of Return) and NPV (Net Present Value) are two very common and acceptable financial measures that can be used to justify a project.

Project Planning Activities: Communication Plan

Communication plans are used to establish communication procedures among management, team members, and relevant stakeholders. It is appropriate and necessary to determine the communication schedule and define the acceptable modes of communication with your project team, stakeholders and steering committee.

Communication Plan Template											
Process/Funct	ion Name	Project/Prog	Project	t Lead	Project Sponsor/Champion						
Communication Purpose:											
Target Audience	Key Message	Message Dependencies	Delivery Date	Location	Medium	Follow up Medium	Messenger	Escalation Path	Contact Information		
Concernant of				and the second second			and the second second		Chan Chan		

Fig. 1.41 Communication Plan Example

Effective communication and frequency of communication are vital to any project. Keeping participants, stakeholders, and customers updated with progress and forecasts helps ensure good communication.

Project Planning Activities: Risk Management Plan

Risk management plans help to identify the sources of project risks and estimate the effects of those risks. Risks may arise from new technology, availability of resources, lack of inputs from customers, business risks or other unexpected sources.

It is important to assess the impact of risk to customers and project stakeholders. Calculate the probability of risk occurrence based on previous or similar projects and industry benchmarks. Then put a plan together to manage the risks.

Most projects that fail are due to the lack of, or improper, risk management. Identify sources of risk as early as possible in the project. Determine the severity and frequency of possible risk occurrences and establish controls or mitigation plans. Prioritize your energy and resources based on risk severity.

Project Planning Tools Advantages

Project planning tools are very useful to organize and communicate project plans, status, and projections. The project planning tools we just reviewed (WBS, Schedule, Gantt, PERT etc.) are some of the most popular and readily available tools and methods used by professionals today.

Project planning tool advantages:

- Help link tasks and sub-tasks or other work elements to get a whole view of what needs to be accomplished
- Allow a more objective comparison of alternative solutions and provide consistent coverage of responsibilities
- Allow for effective scope control and change management
- Facilitate effective communication with all project participants and stakeholders
- Help define management reviews

- Act as an effective monitoring mechanism for the project
- Establish project baselines for progress reviews and control points

1.4 LEAN FUNDAMENTALS

1.4.1 LEAN AND SIX SIGMA

What is Lean?

A Lean enterprise is one which intends to eliminate waste and allow only value to be pulled through its system. A Lean enterprise can be achieved by identifying and effectively eliminating all waste (which will result in a flowing, cost-effective system).

A Lean manufacturing system drives value, flows smoothly, maximizes production, and minimizes waste. Lean manufacturing is characterized by:

- Identifying and driving value
- Establishing flow and pull systems
- Creating production availability and flexibility
- Zero waste
- Waste elimination
- Waste identification and elimination is critical to any successful Lean enterprise
- Elimination of waste enables flow, drives value, cuts cost, and provides flexible and available production

The Five Lean Principles

The following five principles of Lean are taken from the book *Lean Thinking* (1996) by James P. Womack and Daniel T. Jones.

- 1. Specify value desired by customers.
- 2. Identify the value stream.
- 3. Make the product flow continuous.
- 4. Introduce pull systems where continuous flow is possible.
- 5. Manage toward perfection so that the number of steps and the amount of time and information needed to serve the customer continually falls.

Principle 1: Specify Value Defined by Customers

Only a small fraction of the total time and effort spent in an organization actually adds value for the end customer. With a clear definition of value (from the customer's perspective), it is much easier to identify where the waste is.

Principle 2: Identify the Value Stream

The value stream is the entire set of activities across all parts of the organization involved in jointly delivering the product or service. It is the end-to-end process that delivers the value to the customer. As stated above, once you understand what determines value to the customer, it is easier to determine where the waste is.

Principle 3: Create Flow

Typically, you will find that only 5% of activities add value, but the percentage can be as high as 45% in a service environment. Eliminating most or all of this waste ensures that your product or service flows better. Thus, bringing timely, desired value to the customer without interruption or waiting.

Principle 4: Create Pull

When possible, understand the customer demand on your product or service. Then, create a process to respond to it. In other words, only produce what the customer wants when they want it.

Principle 5: Pursue perfection

At first, reorganizing process steps can improve flow, and as this is done, layers of waste become exposed and exploited. Continuing the process of eliminating, reorganizing, and exploiting waste will continue to drive down the number of steps and time needed to serve the customer. This continuous cycle should be pursued to perfection.

Lean and Six Sigma

Lean and Six Sigma both have the objectives of producing high value (quality) at lower costs (efficiency). They approach these objectives in somewhat different manners but in the end, both Lean and Six Sigma drive out waste, reduce defects, improve processes, and stabilize the production environment.

Lean and Six Sigma are a perfect combination of tools for improving quality and efficiency. The two methodologies complement each other to improve quality, efficiency, and ultimately profitability and customer satisfaction.

1.4.2 HISTORY OF LEAN

History of Lean

Lean thinking has been traced as far back as the 1400s but the concepts and principles of Lean were more rigorously and methodically applied starting in the early 1900s. The first person to truly apply the principles in a production process was Henry Ford. He established the first mass production system in 1913 by combining standard parts, conveyors, and work flow. In 1913, Ford brought together the concepts of consistently interchangeable parts, standard work, and moving conveyance to create what he called *flow production*.

Decades later, between 1948 and 1975, Kiichiro Toyoda and Taiichi Ohno at Toyota improved and implemented various new concepts and tools (e.g., value stream, takt time, Kanban etc.) many based on Ford's efforts.

Toyota developed what is known today as the Toyota Production System (TPS) based in Lean principles. TPS advanced the concepts of Ford's system through a series of innovations that provided the continuity of flow *and* a variety of product offerings.

The term Lean, was later coined in the 1990's by John Krafcik who was a graduate student at MIT working on a research project for the book <u>The Machine That Changed the World</u> by Jim Womack. Womack, the founder of the Lean Enterprise Institute helped to refine and advance collectively, the principles known today as Lean.

1.4.3 SEVEN DEADLY MUDA

The Seven Deadly Muda

Muda is a Japanese word meaning "futility, uselessness, idleness, superfluity, waste, wastage, wastefulness." The seven commonly recognized forms of waste, also called the "Seven Deadly Muda," as defined by Taiicho Ohno (the Toyota Production System), are:

- 1. Defects
- 2. **Overproduction**—Producing more than what your customers are demanding
- 3. **Over-Processing**—Unnecessary time spent (e.g., relying on inspections instead of designing a process to eliminate problems)
- 4. **Inventory**—Things awaiting further processing or consumption
- 5. Motion—Extra, unnecessary movement of employees
- 6. Transportation—Unnecessary movement of goods
- 7. **Waiting**—For an upstream process to delivery, for a machine to finish processing, or for an interrupted worker to get back to work

The Seven Deadly Muda: Defects

Defects or defectives are an obvious waste for any working environment or production system. Defects require rework during production or afterwards when the product is returned from an unhappy customer.

Some defects are difficult to solve and often create workarounds and hidden factories. Nobody likes defects. They are an obvious form of waste—wasted time, wasted materials, and wasted money. Defects take time and resources to be fixed, and can be especially costly if they get into the hands of a customer. Eliminating defects is a sure way to improve product quality, customer satisfaction, and production costs.

The Seven Deadly Muda: Overproduction

Overproduction is wasteful because your system expends energy and resources to produce more materials than the customer or next function requires. Overproduction is one of the most detrimental of the seven deadly muda because it leads to other wastes such as overproduction, inventory, transportation and it also exacerbates existing wastes already inherent in the process such as defects, waiting or needless motion.

The Seven Deadly Muda: Over-Processing

Over-processing occurs any time more work is done than is required by the next process step, operation, or consumer. Over-processing also includes being over capacity (scheduling more workers than required or having more machines than necessary).

Another form of over-processing can be buying tools or software that are more precise, complex, or more expensive than required. You might think of this as over-engineering a process or just doing more than is required at a specific step of the process (having more labor available than necessary, having more machines than are needed, or having tools that are not adding incremental value because they do more than the product requires).

The Seven Deadly Muda: Inventory

Inventory is an often-overlooked waste and one that needs to be managed meticulously in order to optimize turns and resources. Take this book for instance. Our distributors require inventory so that they can pack and ship this book when the customer orders it. Unfortunately, inventory is costly up front while revenues lag. If we print too many books and never sell them, then we have excess waste in the form of inventory.

If on the other hand, we create too little inventory and our distributors run out of stock leaving orders unfulfilled then we have an opportunity cost in the form lost or delayed revenues due to poor inventory management.

In either case, inventory has a significant impact on cash flow which is a vital resource for businesses to survive and prosper. Proper inventory management is important to managing waste and opportunity costs.

The 7 Deadly Muda: Motion

Motion is another form of waste often occurring as a result of poor setup, configuration, or operating procedures. Wasted motion can be experienced by machines or humans and is exaggerated by repetition or recurring tasks.

Wasted motion is very common with workers who are unaware of the impact of small unnecessary movements in repetitive tasks. This can result from an improperly planned setup, work area configuration, and operating procedure. Small movements of an employee probably do not seem very wasteful by themselves. But when you consider repetitive motion that occurs over a long period, your perspective may change.

The Seven Deadly Muda: Transportation

Transportation is the unnecessary movement of goods, raw materials or finished products. Transportation is considered wasteful because it does *nothing* to add value or transform the product.

Imagine for a moment driving to and from work twice before getting out of your car to go into work . . . That is waste in the form of transportation. Driving less is better! In a similar way, the less transportation a product must endure, the better. There would be fewer opportunities for delay, destruction, loss, damage etc.

Transportation is wasteful because it takes energy to move something. And, it may take costly tools to move it. Transportation also introduces the risk of something getting damaged, lost, misplaced, delayed etc. Unless something is being transported into the hands of the customer, it probably means it is going somewhere to be stored (which is also wasteful).

The Seven Deadly Muda: Waiting

Waiting is an obvious form of waste and is typically a symptom of an upstream problem. Waiting is usually caused by inefficiency, bottlenecks, or poorly-designed work flows within the value stream, but it can also be caused by inefficient administration.

Reduction in waiting time will require thoughtful applications of Lean and process improvement. Waiting is when a downstream process is being starved because there is something wrong upstream in the process (like a machine having to finish processing or a worker that has stepped away from his or her station). In other words, there is a "bottleneck" that inhibits the flow of the process.

1.4.4 FIVE-S (5S)

What is 5S?

5S is systematic method used to organize, order, clean, and standardize a workplace . . . and, to keep it that way! *5S* is a methodology originally developed in Japan and is used for improving the Lean work environment. *5S* is summarized in five Japanese words all starting with the letter *S*:

- 1. Seiri (sorting)
- 2. Seiton (straightening)
- 3. Seiso (shining)
- 4. Seiketsu (standardizing)
- 5. Shisuke (sustaining)

To separate the needed from the unnecessary and prioritize what is needed. Sorting in a 5s activity is accomplished by many practitioners using the red tag method whereby red tags are placed on items in a work environment that are no longer useful or need to be removed. A common mantra in the sorting stage is "when in doubt" "throw it out".

Seiton (Set in order or straighten)

To set in order is to organize the work environment. This means to place all tools and equipment (those remaining from the sort stage) into the most useful, logical and accessible locations. Organization should be such that it promotes productivity and avails itself to the concepts of the visual factory.

Another common mantra used in 5S is "A place for everything and everything in its place". This saying keeps the focus on orderliness and not just during the 5S activity but for many years thereafter.

One of the most memorable examples of proper "set in order" results is that of the tool outlines on a peg board. When a tool is removed from the board it becomes very evident that the tool is no longer in its storage location. If it's not in use, then it should be where it can be found by other workers.

Seiso (Shine)

Shine means to clean the work area. This is not a casual sweep and "de-clutter" activity either. Shine in 5S calls for "getting dirty" while cleaning. Examples are machinery to be maintained, equipment cleaned and treated or painted, machinery vacuumed, equipment greased, lubed and brought to optimal performance levels.

This is the stage where a measure of pride becomes evident, when other departments and other work areas begin to express interest in conducting 5S on their own work environments.

Seiketsu (Standardize)

Standardizing is the stage where work practices are staged to be performed consistently. Clear roles and responsibilities are outlined and everyone is expected to know their role. Standardize also means to make workstations and tool layouts common among like functions and normalize equipment and parts across all workstations so that workers can be moved around to any workstation at any time and perform the same task.

Shisuke (Sustain)

Sustaining in 5S is also called self-discipline. It takes discipline to maintain 5S. Obviously, it takes time for new habits to form, and there needs to be a culture within the team to follow this methodology. It is critical not to fall back into old ways. Therefore, it is critical that the sustain stage of 5s:

- Creates the culture in the team to follow the first four S's consistently.
- Avoids falling back to the old ways of doing things with cluttered and unorganized work environments.
- Establishes and maintains the momentum of optimizing the workplace.
- Promotes innovations of workplace improvement.
- Sustains the first four stages using:
 - o 5S Maps

- o 5S Schedules
- 5S Job cycle charts
- Integration of regular work duties
- 5S Blitz schedules
- \circ $\,$ Daily workplace scans to maintain and review standards $\,$

Goals and Benefits of 5S

The goals of 5S are to reduced waste, cut unnecessary expense, optimize efficiency and establish a work environment that is:

- Self-explaining
- Self-ordering

- Self-regulating
- Self-improving

Thus, 5S should create a work environment where this is no more wandering and searching for tools or parts. Workers should experience fewer delays or wait time due to equipment down time and the overall workplace will be safer, cleaner and more efficient. Some of the more common benefits reported by 5S participants are:

- Reduced changeovers
- Reduced defects
- Reduced waste
- Reduced delays
- Reduced injuries
- Reduced breakdowns

- Reduced complaints
- Reduced red ink
- Higher quality
- Lower costs
- Safer work environment
- Greater capacity

By following the disciplined 5S method and meeting the goals established by the 5S team, the following quantifiable benefits have been realized:

- Cut in floor space: 60%
- Cut in flow distance: 80%
- Cut in accidents: 70%
- Cut in rack storage: 68%
- Cut in number of forklifts: 45%
- Cut in changeover time: 62%

- Cut in physical inventory: 50%
- Cut in training requirements: 55%
- Cut in nonconformance: 96%
- Increase in test yields: 50%
- Late deliveries: 0%
- Increase in throughput: 15%

5S Disclaimer

As a method, 5S is one of many Lean methods that creates immediate and observable improvements. It is tempting and often attempted by organizations to implement 5S alone without considering the entire value stream. However, it is advisable to consider a well-planned Lean manufacturing approach to the entire production system.

2.0 MEASURE PHASE

2.1 PROCESS DEFINITION

2.1.1 CAUSE AND EFFECT DIAGRAM

What is a Cause and Effect Diagram?

A *cause and effect diagram* is also called a *fishbone diagram* or *Ishikawa diagram*. It was created by Kaoru Ishikawa and is used to identify, organize, and display the potential causes of a specific effect or event in a graphical way similar to a fishbone.

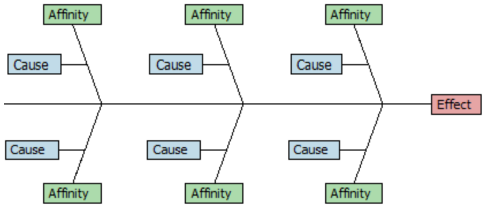


Fig. 2.1 Cause & Effect Diagram

A powerful brainstorming tool, the cause and effect diagram illustrates the relationship between one specified event (output) and its categorized potential causes (inputs) in a visual and systematic way.

Major Categories of Potential Causes

Typically, the possible causes fall into six major categories (or branches in the diagram). The acronym P4ME is a helpful way to remember the following standard categories which can be used when no other categories are known or obvious for your project:

- **P**eople: People who are involved in the process
- Methods: How the process is completed (e.g., procedures, policies, regulations, laws)
- Machines: Equipment or tools needed to perform the process
- Materials: Raw materials or information needed to do the job
- Measurements: Data collected from the process for inspection or evaluation
- Environment: Surroundings of the process (e.g., location, time, culture).

How to Plot a Cause and Effect Diagram

Step 1: Identify and define the effect being analyzed.

• Clearly state the operational definition of the effect or event of interest.

- The effect can be the positive outcome desired or negative problem targeted to solve.
- Enter the effect in the end box of the fishbone diagram and draw a spine pointed to it.
- Plotting a cause and effect diagram is a simple four-step process.

Step 2: Brainstorm the potential causes or factors of the effect occurring.

- Identify any factors with a potential impact on the effect and include them in this step.
- Put all the identified potential causes aside for use later.

Step 3: Identify the main categories of causes and group the potential causes accordingly.

- Besides P4ME (i.e., people, methods, machines, materials, measurements, and environment), you can group potential causes into other customized categories.
- Below each major category, you can define sub-categories and then classify them to help you visualize the potential causes.
- Enter each cause category in a box and connect the box to the spine. Link each potential cause to its corresponding cause category.

You can use P4ME, or you can use customized main categories to group the potential causes. Under each main category, you can define sub-categories if additional levels of detail are needed.

Step 4: Analyze the cause and effect diagram.

- A cause and effect diagram includes all the possible factors of the effect being analyzed.
- You can use a Pareto chart to filter the causes the project team needs to focus on.
- Identify causes with high impact that the team can act upon.
- Determine how to measure causes and effects quantitatively.
- Prepare for further statistical analysis.

Benefits to Using Cause and Effect Diagram

The benefits to using a cause and effect diagram to analyze an effect or event are:

- Helps to quickly identify and sort the potential causes of an effect.
- Provides a systematic way to brainstorm potential causes.
- Identifies areas requiring data collection for further quantitative analysis.
- Locates "low-hanging fruit."

Limitation of Cause and Effect Diagrams

• A cause and effect diagram only provides qualitative analysis of correlation between each cause and the effect. More analysis is needed to quantify relationships between causes and effects.

- One cause and effect diagram can only focus on *one* effect or event at a time. Effects can have different causes.
- Further statistical analysis is required to quantify the relationship between various factors and the effect and identify the root causes.

Cause and Effect Diagram Example

Let us follow an example through the process of using a cause and effect diagram. A real estate company is interested to find the root causes of high energy costs of its properties. The cause and effect diagram is used to identify, organize, and analyze the potential root causes.

Step 1: Identify and define the effect being analyzed: In this example, the effect we are concerned with is high energy costs of buildings.

Step 2: Brainstorm the potential causes or factors of the high-energy costs: A team is assembled to brainstorm possible reasons for high energy costs of buildings. They provided the following list of possible causes:

- Poor Maintenance
- Bad Habits
- Lighting Schedule
- HVAC Schedule
- Temperature Set Point
- Old HVAC Units
- Open Dampers

- Light Bulbs
- Inefficient Building Materials
- Meter Float
- Meter Accuracy
- Increasing Fuel Cost
- Building Air Leakage
- Humidity

Step 3: Identify the main categories of causes and group the potential causes accordingly. The team used the standard P4ME categories for this exercise and bucketed the causes above based on the most appropriate categories.

Step 4: Analyze the cause and effect (C&E) diagram.

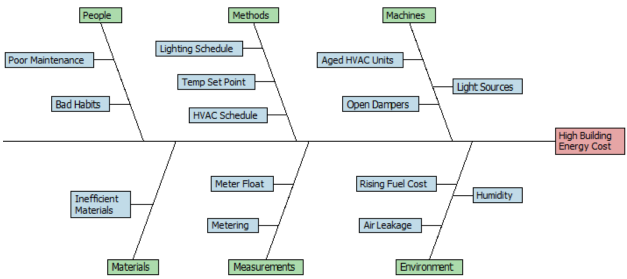


Fig. 2.2 Populated Cause & Effect Diagram with P4ME Categories

After completing the C&E diagram, the real estate company conducts further research on each potential root cause. It is discovered that:

- The utility metering is accurate
- The building materials are acceptable and there is no significant air leakage
- The fuel prices increased recently but were negligible
- Most lights are off during the non-business hours except that some lights have to be on for security purposes
- The temperature set points in the summer and winter are both adequate and reasonable
- The high-energy costs are probably caused by the poor HVAC maintenance on aged units and the wasteful energy consuming habits

Next, the real estate company needs to collect and analyze the data to check whether root causes identified in the C&E diagram have any validity. Data should be used to support this conclusion. Therefore, the team need to consider how to collect data to prove or disprove their assertion.

2.1.2 CAUSE AND EFFECTS MATRIX

What is a Cause and Effect Matrix?

The *cause and effect matrix* (XY Matrix) is a tool to help subjectively quantify the relationship of several X's to several Y's. Among the Y's under consideration, two important ones should be the *primary* and *secondary metrics* of your Six Sigma project. The X's should be derived from your cause and effect diagram. Let us take a peek as to what it looks like on the next page.

A cause and effect matrix is a powerful yet easy tool to transition from brainstorming (which you did using the cause and effect diagram) to estimating, ranking, and quantifying the

relationship between those things you found during brainstorming and the Y's that are important to the project.

Cause and Effects Matrix

Project: XY Matrix Owner:											
Ar Matrix Owner.											
Output Measures (Y's)	Y ₁	Y ₂	Y ₃	Y ₄	Y ₅	Y ₆	Y ₇	Y ₈	Y ₉	Y ₁₀	
Weighting (1-10):											
Input Variables (X's)#		For	each X, so	ore its imp	act on eac	h Y listed a	bove (use a	1 0,3,5,7 sc	:ale)		Score
X ₁											0
X ₂											0
X ₃											0
X ₄											0
X ₂₉											0
X ₃₀		0		0	0						0

Fig. 2.3 Cause & Effects Matrix

Figure 2.3 represents a clean cause and effect matrix where you can translate all of the potential X's brainstormed during the creation of your cause and effect diagram session.

Once you have that information entered into the matrix you can add one or more Y's to the top of the matrix. The Y's should be weighted relative to each other based on their importance to the project.

Once this is complete, you can begin rating each X against each Y on a scale of 0, 3, 5, 7 (0 means no impact; 3 means weak impact; 5 is moderate impact; and 7 is strong impact). Try to use the whole spectrum of your scale during this process because you will need the resolution to help differentiate the X's from one another.

How to Use a Cause and Effect Matrix

- 1. Across the top enter your output measures. These are the Y's that are important to your project.
- 2. Next, give each Y a weight. Use a 1–10 scale, 1 being least important and 10 most important.
- 3. Below, in the leftmost column, enter all the variables you identified with your cause and effect diagram.

- 4. Within the matrix, rate the strength of the relationship between the X in the row and the corresponding Y in that column. Use a scale of 0, 3, 5, and 7.
- 5. Lastly, sort the "Score" column to order the most important X's first.

Cause and Effect Matrix Notes

When complete, sort the X's based on their scores and graph them (use a Pareto chart). What the results will tell you should not be "taken to the bank" as this entire process from brainstorming X's to rating them against Y's is a subjective exercise.

However, if you have chosen your team wisely, it is very common that someone or several SMEs on your project team know the real solutions and/or the best way to narrow down to find them. Therefore, it is also important not to dismiss the results of your cause and effect matrix. Look at the image with comments embedded to summarize what you have just read.

After You Have Completed the C&E Matrix

After you have completed your cause and effects matrix, build a strategy for validating and/or eliminating the x's as significant variables to the Y=f(x) equation.

- Build a data collection plan
- Prepare and execute planned studies
- Perform analytics
- Review results with SMEs, etc.

The combination of the C&E diagram and the C&E matrix is powerful. This is where some of the best ideas and theories about the Y=f(x) equation begin.

Now, it is time to start eliminating and/or validating these variables. Put a plan together to do so. Use the resources and tools you have already learned as well as those that you are about to learn throughout this course.

2.1.3 FAILURE MODES AND EFFECTS ANALYSIS (FMEA)

What is FMEA?

							-	CLAR .	are meduco	a Linout	- many on	~			-
Product or Process Step	Potential Failure Mode	Potential Failure Effects	s	Potential Causes	0	Current Controls	D	R P N	Recommended Actions	Responsible	Actions Taken	s	0	D	R P N
						1		0			6				0
					1.1			0							0
								0					1		0
								0							0
					-			0					-		0
					. 8			0							0
								0					1		0

Failure Modes & Effects Analysis - FMEA

Fig. 2.4 Failure Modes & Effects Analysis Template

Figure 2.4 is a template of an FMEA. The *FMEA (Failure Modes and Effects Analysis)* is a tool and an analysis technique used to identify, evaluate, and prioritize potential deficiencies in a process so that the project team can design action plans to reduce the probability of those failures from occurring.

FMEA activity often follows process mapping. A process map is commonly used during the FMEA to determine where potential process breakdowns can occur. At each step, the team must ask, "What can go wrong?" A Pareto analysis may follow to quantitatively validate or disprove the results of the FMEA.

FMEA is best completed in cross-functional brainstorming sessions where attendees have a good understanding of the entire process or of a segment of it.

Important FMEA Terms

- *Process Functions* Process steps depicted in the process map. FMEA is based on a process map and one step/function is analyzed at a time.
- *Failure Modes* Potential and actual failure in the process function/step. It usually describes the way in which failure occurs. There might be more than one failure mode for one process function.
- *Failure Effects* Impact of failure modes on the process or product. One failure mode might trigger multiple failure effects.
- *Failure Causes* Potential defect of the design that might result in the failure modes occurring. One failure mode might have multiple potential failure causes.
- *Severity Score* The seriousness of the consequences of a failure mode occurring. Ranges from 3 to 9, with 9 indicating the most severe consequence.
- *Occurrence Score* The frequency of the failure mode occurring. Ranges from 3 to 9, with 9 indicating the highest frequency.
- *Detection Score* How easily failure modes can be detected. It is a rating for how effective any controls are in the process. Ranges from 3 to 9, with 9 indicating the most difficult detection.
- *RPN (Risk Prioritization Number)* The product of the severity, occurrence, and detection scores. It is a calculated value that ultimately is used to prioritize the FMEA. Ranges from 1 to 1000. The higher RPN is, the more focus the particular step/function needs.
- *Recommended Actions* The action plan recommended to reduce the probability of failure modes occurring or at least enable better controls to detect it when failure occurs.
- *Current Controls* Procedures currently conducted to prevent failure modes from happening or to detect the failure mode occurring.

How to Conduct an FMEA

Conducting an FMEA is not a difficult process, but can be time consuming when done correctly, particularly the first time it is performed on a process. To demonstrate how an FMEA is used we will be sharing a super-simple example.

Joe is trying to identify, analyze, and eliminate the failure modes he experienced in the past when preparing his work bag before heading to the office every morning. He decides to run an FMEA for his process of work bag preparation. There are only two steps involved in the process: putting the work files in the bag and putting a water bottle in the bag.

											~				
Product or Process Step	Potential Failure Mode	Potential Failure Effects	S	Potential Causes	0	Current Controls	D	R P N	Recommended Actions	Responsible	Actions Taken	S	0]	D	R P N
								0							0
						······		0							0
								0						1	0
						······		0						1	0
								0						1	0
						······		0						1	0
		÷	ę		÷				•		÷	ę			

Failure Modes & Effects Analysis - FMEA

Figure 2.5 shows a clearer picture of the header of the FMEA template. For the example we are about to share, we will be starting in the left column and for the most part, we will work through the template from left to right.

FMEA Step 1: List the critical functions of the process based on the process map created. Here you see that the process steps are listed under the "Process Function" heading. In this case, the process is extremely simple so that we can focus on how to use FMEA as a tool. The two process steps populated are Joe putting work files in the bag and Joe putting his water bottle in the bag.

Product or Process Step	Potential Failure Mode	Potential Failure Effects
Place files in bag		
Put water bottle in bag		

Fig. 2.6 FMEA List Process Steps

FMEA Step 2: List all the potential failure modes that might occur in each function. In figure 2.7 you can see that we have chosen to list only one failure for each process step. In your case, be sure to be thorough and not overlook any possible failure modes. We could add other failure modes such as placed files in wrong pocket, placed bent/crumpled file in bag etc.

Fig. 2.5 FMEA Header

Product or Process Step	Potential Failure Mode	Potential Failure Effects
Place files in bag	Incorrect files put in the bag	
Put water bottle in bag	Water leaks	

Fig. 2.7 FMEA Potential Failure Modes

FMEA Step 3: List all the potential failure effects that might affect the process. In this situation, if Joe places the wrong files in his bag, his work will be delayed when he arrives at his destination because he will have the wrong files. Or, his files will be destroyed because his water bottle leaked.

Product or Process Step	Potential Failure Mode	Potential Failure Effects
Place files in bag	Incorrect files put in the bag	Work is delyed
Put water bottle in bag	Water leaks	Files in bag damaged

Fig. 2.8 FMEA Failure Effects

FMEA Step 4: List all the possible causes that may lead to the failure mode happening. Joe may have had a loose cap on his water bottle or disorganized files which caused his failures. There can be more than one cause to the same failure. List each one separately.

Potential Failure Effects	S	Potential Causes	0
Work is delyed		Files are not organized well	
Files in bag damaged		Cap on water bottle not tight	

Fig. 2.9 FMEA Failure Causes

FMEA Step 5: List the current control procedures for each failure cause. In many cases, you'll find that there are no controls or weak controls relative to your failure causes. The FMEA will help you and your team identify these weaknesses.

Potential Causes	0	Current Controls	D
Files are not organized well		Check if files are needed	
Cap on water bottle not tight		Check bottle cap before inserting	

Fig. 2.10 FMEA Current Controls

FMEA Step 6: Determine the severity rating for each potential failure mode. At this point it is necessary to attempt to quantify the failure effects in terms of their severity. As you progress, will see that there will be 3 ratings to be assigned in an FMEA (Severity, Occurrence and Detection). These quantifications will result in a total "Risk Priority Number" (RPN) which will help you determine the failure modes that are most detrimental to your process.

For severity rankings. We recommend using a scale range of 3,5,7,9 with 9 representing the most severe and 3 being the least severe. If a failure mode has more than one cause, severity rankings should be the same for each cause because you're ranking the severity of the failure not of the cause.

Potential Failure Effects	S	Potential Causes	0	Current Controls
Work is delyed	9	Files are not organized well		Check if files are needed
Files in bag damaged	7	Cap on water bottle not tight		Check bottle cap before inserting

Fig. 2.11 FMEA Failure Effects Severity Rating

FMEA Step 7: Determine the occurrence rating for each potential failure cause. Using the same 3,5,7,9 scale. Rate each failure causes based on how frequently you believe they occur. A rating of 9 will represent a high occurrence rating and 3 will represent an infrequent or low occurrence rating. If there is more than one cause to a single failure mode, rate each cause based on how frequently the failure occurs due to that particular cause.

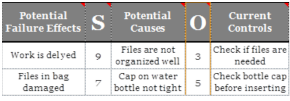


Fig. 2.12 FMEA Failure Occurrence Rating

FMEA Step 8: Determine the detection rating for each current control procedure. Detection ratings are intuitively different and inverse of the others two ratings. High detection ratings mean that control procedure has a poor ability to detect the failure when it occurs. Conversely, a low detection rating of 3 indicates that the control procedure is very capable of detection. This ensures that the product of all 3 ratings will yield an RPN that when ranked, the highest RPN will indicate which failure mode is in need of the most attention.

S	Potential Causes	0	Current Controls	D	R P N
9	Files are not organized well	3	Check if files are needed	5	135
7	Cap on water bottle not tight	5	Check bottle cap before inserting	5	175

Fig. 2.13 FMEA Failure Detection Rating

FMEA Step 9: Calculate the Risk Prioritization Number (RPN). The RPN is calculated by multiplying all 3 ratings together $S \ge 0 \ge D = RPN$.

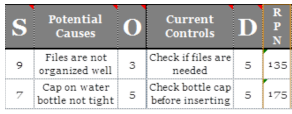


Fig. 2.14 FMEA Risk Priority Number (RPN)

FMEA Step 10: Rank the failures using the RPN and determine the precedence of problems or critical inputs of process. Ranking should be from highest to lowest.

S	Potential Causes	0	Current Controls	D	R P N
7	Cap on water bottle not tight	5	Check bottle cap before inserting	5	175
9	Files are not organized well	3	Check if files are needed	5	135

Fig. 2.15 FMEA Sort by RPN

FMEA Step 11: Brainstorm and create recommended action plans for each failure mode. There can be multiple actions per failure mode, or cause or control procedure. The purpose is to thoroughly address root causes as well as your ability to detect failures. The goal would be to prevent failures.

Current Controls	D	R P N	Recommended Actions
Check bottle cap before inserting	5	175	Obtain new water bottle
Check if files are needed	5	135	Organize & Categorize Files

Fig. 2.16 FMEA Recommended Actions

FMEA Step 12: Determine and assign task owners and projected completion dates for each action item. This is an important accountability step and will be further supported by management routines that review progress and the effectiveness of the action items.

D	R P N	Recommended Actions	Responsible
5	175	Obtain new water bottle	Joe
5	135	Organize & Categorize Files	Joe

Fig. 2.17 FMEA Action Responsibility

FMEA Step 13: Determine new severity ratings after actions have been taken. FMEAs are often considered living and breathing documents because they should continuously address failure modes and require continued updating to actions, rankings and RPNs.

Recommended Actions	Responsible	Actions Taken	S	O D	R P N
Obtain new water bottle	Joe				0
Organize & Categorize Files	Joe				0

Fig. 2.18 FMEA After Action Failure Severity

FMEA Step 14: Determine the new occurrence ratings after actions are taken.

Recommended Actions	Responsible	Actions Taken	S	0	D	R P N
Obtain new water bottle	Joe					0
Organize & Categorize Files	Joe					0

Fig. 2.19 FMEA After Action Failure Occurrence Rating

FMEA Step 15: Determine new detection ratings after actions are taken.

Recommended Actions	Responsible	Actions Taken	S	0	D	R P N
Obtain new water bottle	Joe					0
Organize & Categorize Files	Joe					0

Fig. 2.20 FMEA After Action Failure Detection Rating

FMEA Step 16: Update the RPN based on new severity, occurrence, and detection ratings.

Recommended Actions	Responsible	Actions Taken	S	O D	R P N
Obtain new water bottle	Joe				0
Organize & Categorize Files	Joe				0

Fig. 2.21 FMEA List Process Steps

Make sure the corrective action plan is robust. Make sure it is clear what actions are to be taken, who owns the task, and what the completion date is for the task. Re-assess the severity, occurrence, and detection ratings if the corrective action plan is carried out, and update the RPN. Continue this cycle as needed.

2.1.4 Theory of Constraints

What is Theory of Constraints (TOC)?

Processes, systems, and organizations are all vulnerable to their weakest part. Any manageable system is limited by constraints in its ability to produce more (and there is always at least one constraint).

A common analogy for the theory of constraints is, "a chain is no stronger than its weakest link." This concept was introduced by Eliyahu M. Goldratt in his 1984 book titled *The Goal*.

Theory of Constraints (TOC) is an important tool for improving process flows. For most organizations, the goal is to make money. For some organizations (e.g., non-profits) making money is a necessary condition for pursuing the goal.

Constraints can come in many forms. Some examples are production capacity, material, logistics, the market (demand), employee behavior, or even management policy. The underlying premise of TOC is that organizations can be measured and controlled through variation of three measures—throughput, operational expense, and inventory.

TOC Performance Measures

Making sound financial decisions based on these three measures is a critical requirement.

- Throughput—Rate at which a system generates money through sales
- Operational Expense—Money spent by the system to turn inventory into throughput
- Inventory—Money the system has invested in purchasing things it intends to sell

Why is throughput defined by sales? Goods are not considered an asset until sold. This contradicts the common accounting practice of listing inventory as an asset even if it may never be sold. Units that are produced but not sold are inventory.

The objective of a firm is to increase throughput and/or decrease inventory and operating expense in such a way as to increase profit, return on investment, and cash flow.

TOC Five Focusing Steps

The objective is to ensure ongoing improvement efforts are focused on the constraints of a system.

- 1. Identify the system's constraints.
- 2. Decide how to exploit the constraints.
- 3. Subordinate everything else to the decision in step 2.
- 4. Elevate the constraints.
- 5. If in previous steps a constraint has been broken, return to step 1, but do not allow inertia to cause a system's constraint.

TOC focuses on the output of an entire system versus a discrete unit of the components. The five focusing steps help to identify the constraint that towers above all others within the

system. It is an iterative process. Once the largest constraint is addressed (strengthened), then the next weakest link in the chain should be addressed, and so on. It is an ongoing system of process improvement.

Logical Thinking Processes

	Focusing Step	Thinking Process	Tools
1	Identify the system's constraint(s)	 Identify the problems Find the root causes	Cause and effect diagram
2	Decide how to exploit the constraint(s)	Develop a solution	Future reality tree
3	Subordinate everything else to the decision in step 2	 Identify the conflict preventing the solution Remove the conflict	Evaporating cloud
4	Elevate the constraint	• Construct and execute an implementation plan	Prerequisite tree Transition tree
5	If in previous steps a cons to cause a system's cons	straint has been broken, return to traint	step 1, but do not allow inertia

Fig. 2.22 TOC Logical Thinking Processes

Used in conjunction with the five focusing steps. There are many logic tools to help a firm analyze and verbalize cause and effect:

- E-C-E (Effect-Cause-Effect) Diagramming
- Current reality tree, also known as cause and effect diagram -Root cause identification
- Future reality tree—Assess solutions
- Evaporating cloud—Conflict resolution tool
- Prerequisite tree—Implementation tool
- Transition tree—Implementation plan

Simulation Exercise

Resources needed:

- Three "production line" participants
- One timer per each production line participant
- Five small boxes of 15 widgets each (paperclips, pens/pencils, candy, etc.)

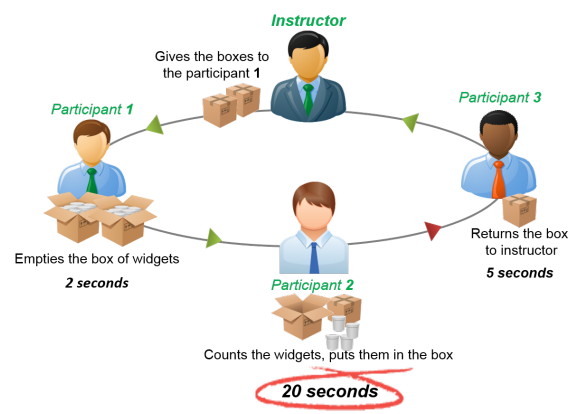


Fig. 2.23 TOC Widget Value Chain Simulation

Perform the process slowly the first time. The instructor gives the boxes, one at a time, to the first participant in the production line who empties the contents and hands the box and widgets to the next participant. The second participant counts the widgets, puts them in the box, and hands it to the third person who returns the box to the instructor. The instructor varies the rates at which the five boxes of widgets are handed to the first person in the "value chain" and times are recorded for each member's part in the process.

Considerations or Observations:

- As the boxes are distributed slowly, it is clear that all participants have plenty of time to do their activities, but they are all being starved of work and the "bottleneck" is external to the process (the instructor).
- As the instructor speeds up the process and hands over the boxes at a much faster rate, all three participants start to work faster, with the person in the middle always busy as the other two participants are waiting for him or her. In essence, the bottleneck has now changed to Person B, and a process constraint or bottleneck has been identified.
- Have the timers provide an average time for each of the three steps. Calculate throughput" for each step in the value chain (follow example below).

Participant 1: Time = 2 seconds (Throughput = 30 boxes/minute) Participant 2: Time = 20 seconds (Throughput = 3 boxes/minute) ← Bottleneck Participant 3: Time = 5 seconds (Throughput = 12 boxes/minute)

The efficiency of the process has slowed down to the slowest resource. In the example above, the output of the whole process is three boxes a minute; and Participant 1 and Participant 3 are not contributing to the overall efficiency. If they were to slow down, they would not have an adverse impact on the process, since they will always have idle time. Such resources with extra capacity are non-bottleneck resources.

Key question: What would you do to improve the efficiency of this system? Use the five focusing steps to lead to a conclusion and an action plan.

- 1. Identify the system's constraints.
- 2. Decide how to exploit the constraints.
- 3. Subordinate everything else to the decision in step 2.
- 4. Elevate the constraints.
- 5. If in previous steps a constraint has been broken, return to step 1, but do not allow inertia to cause a system's constraint.

Focusing Step 1: Identifying the constraint. In this exercise, "Count widgets and fill box" is the bottleneck; however, it is not always that easy to identify the bottleneck, so brainstorming tools like cause and effect diagrams are helpful.

Focusing Step 2: Determine how to exploit the constraint? This is where you would brainstorm potential solutions.

- More resources to perform step 2 (adds cost to the process).
- Sharing the work with Participant 1 or 3 (no added cost).
- Other ideas?

Focusing Step 3: Determine what conflicts prevent the solution?

- Perhaps there is resistance to spending more on additional resources?
- Maybe resources performing other steps require additional training/certification to perform another function?

Focusing Step 4: Determine the plan to implement?

- Hiring new resources? Training?
- What other considerations for implementing a solution?

Focusing Step 5: Implement the plan and then consider where the new bottleneck is.

2.2 SIX SIGMA STATISTICS

2.2.1 BASIC STATISTICS

What is Statistics?

Statistics is the science of collection, analysis, interpretation, and presentation of data. In Six Sigma, we apply statistical methods and principles to quantitatively measure and analyze process performance to reach statistical conclusions and help solve business problems.

sta-tis-tics (as defined on dictionary.com)

1. The science that deals with the collection, classification, analysis, and interpretation of numerical facts or data, and that, by use of mathematical theories of probability, imposes order and regularity on aggregates of more or less disparate elements.

2. The numerical facts or data themselves.

Types of Statistics

Descriptive statistics simply describe what is happening, while *inferential statistics* help you make comparisons, draw conclusions, or make judgments about data.

Descriptive Statistics

Descriptive statistics is applied to describe the main characteristics of a collection of data. It summarizes the features of the data quantitatively only and it does not make any generalizations beyond the data at hand. The data used for descriptive statistics are for the purpose of representing or reporting.

Inferential Statistics

Inferential statistics is applied to "infer" the characteristics or relationships of the populations from which the data are collected. Inferential statistics draws statistical conclusions about the population by analyzing a sample of data subject to random variation. A complete data analysis includes both descriptive statistics and inferential statistics.

Statistics vs. Parameters

The word *statistic* refers to a numeric measurement calculated using a sample data set, for example, sample mean or sample standard deviation. Its plural is *statistics* refers to the scientific discipline of statistical analysis.

Parameter refers to a numeric metric describing a population, for example, population mean and population standard deviation. Unless you have the full data set of the population, you will not be able to know population parameters. Almost always, you will be working with statistics, but they will allow you to make inferences about the population.

Continuous Variable vs. Discrete Variable

Data can take different forms, which is important to recognize when it is time to analyze the data. Continuous variables are measured and can be divided infinitesimally (they can take any value within a range) and there is an infinite number of values possible. Examples are temperature, height, weight, money, time etc.

Discrete variables are sometimes referred to as count data which can literally be counted. Discrete data is finite with limited numbers of values available. Examples are count of people, count of countries, count of defects or count of defectives.

Types of Data

Nominal Data: Nominal Data are categorical but have no natural rank order. Examples of nominal data are colors, states, days of the week etc. Nominal comes from the Latin word "nomen" meaning "name" and nominal data are items that are differentiated by a simple naming system.

Be careful of nominal items that have numbers assigned to them and may appear ordinal but are not. These are used to simplify capture and referencing. Nominal items are usually categorical and belong to a definable category.

Ordinal Data: Ordinal Data are rank order data. Examples of ordinal data are the first, second and third place in a race or scores on an exam. Items on an ordinal scale are set into some kind of *order* by their position on the scale. The order of items is often defined by assigning numbers to them to show their relative position. Nominal and Ordinal data are categorical and thus referred to as discrete data.

Interval: Interval data are data measured on a quantifiable scale with equidistant values throughout the scale. Examples of interval data are temperature as measured on the Fahrenheit or Celsius scale.

Ratio Data: Ratio data are the ratio between the magnitude of a continuous value and the unit value of the same category. Examples of ratio data are weight, length, time etc.

In a ratio scale, numbers can be compared as multiples of one another. Thus, one person can be twice as tall as another person. Also important, the number zero has meaning. Thus, the difference between a person of 35 and a person 38 is the same as the difference between people who are 12 and 15. A person can also have an age of zero.

Interval and ratio data measure quantities and hence are continuous. Because they can be measured on a scale, they may also be referred to as scale data.

2.2.2 Descriptive Statistics

Basics of Descriptive Statistics

Descriptive statistics

Descriptive statistics provide a quantitative summary for the data collected. It summarizes the main features of the collection of data (shape, location and spread). Descriptive statistics is a presentation of data collected and it does *not* provide any inferences about a more general condition. Instead, descriptive statistics describe what the data "looks like."

Shape of the Data

When we describe the "shape" of the data, we typically refer to the *distribution* of the data. Distribution, also called frequency distribution, summarizes the frequency of an individual value or a range of values for a variable (either continuous or discrete). It is sometimes referred to as a "frequency distribution" because it depicts how often a particular value occurs within a data set. Distribution is depicted as a table or graph.

Simple Example of Distribution

We are tossing a fair die. The possible value we obtain from each toss is a value between 1 and 6. Each value between 1 and 6 has a 1/6 chance to be hit for each toss. The distribution of this game describes the relationship between every possible value and the percentage of times the value is hit (or count of times the value is hit).

In this example, there is an equal probability for each value on the die occurring. If you were to roll the die a high number of times, what might the distribution look like? If you rolled it 600 times, you would likely get each value nearly 100 times.

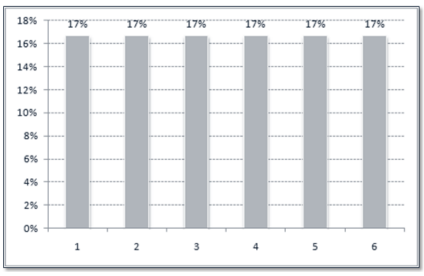


Fig. 2.24 Fair Die Distribution Probability

Common Types of Continuous Distributions

- Normal Distribution
- T distribution
- Chi-square distribution

• F distribution

Common Types of Discrete Distributions

- Binomial distribution
- Poisson distribution

There are different types of distributions that are commonly found in statistics. It is important to recognize that there are different types of data distributions because the statistics can tell you different things about the data and different inferential statistics have to be used for different situations.

Location of the Data

The location (i.e., central tendency) of the data describes the value where the data tend to cluster around. There are multiple measurements to capture the location of the data, the three most common and meaningful are Mean, Median and Mode.

Mean

The mean is the arithmetic average of a data set. It is easily calculated by dividing the sum of the data points by n (n is the number of values in the data set).

$$\bar{\mathbf{x}} = \frac{1}{n} \sum_{i=1}^{n} \mathbf{x}_i$$

For example, we have a set of data: 2, 3, 5, 8, 5, and 9. The arithmetic mean of the data set is

$$\frac{2+3+5+8+5+9}{6} = 5.33$$

In the example above, there are six data points. The sum of the six data points is 32, and when dividing by the number of data points, 6, the arithmetic mean is 5.33.

Median

The median is the middle value of the data set in numeric order. It separates a finite set of data into two even parts, one with values higher that the median and the other with values lower than the median. For example, we have a set of data: 45, 32, 67, 12, 37, 54, and 28. The median is 37 since it's the middle value of the sorted list of those values (i.e., 12, 28, 32, 37, 45, 54, and 67). There are three values lower and three values higher than the median.

Mode

The mode is the value that occurs most often in the data set. If no number is repeated, there is no mode for the data set. For example, we have a data set: 55, 23, 45, 45, 68, 34, 45, 55. The mode is 45 since it occurs most frequently. The numbers 23, 34, and 68 occur once, 55 occurs twice, and 45 occurs three times.

Spread of the Data

The spread (i.e., variation) of the data describes the degree of data dispersing around the center value. There are multiple measurements to capture the spread of the data, the most common and meaningful are Range, Variance and Standard Deviation.

Range

The range is the numeric difference between the greatest and smallest values in a data set. Only two data values (i.e., the greatest and the smallest values) are accounted for calculating the range. For example, we have a set of data: 34, 45, 23, 12, 32, 78, and 23. The largest value is 78, and the smallest is 12. Therefore, the range is 78 - 12 = 66.

Variance

The variance measures how far on average the data points spread out from the mean. More specifically, it is the average squared deviation of each value from its mean. All data points are accounted for by calculating the variance:

$$s^2 = \frac{1}{n} \sum_{i=1}^n (x_i \text{-} \overline{x})^2$$

Where: *n* is the number of values in the data set.

$$\bar{\mathbf{x}} = \frac{1}{n} \sum_{i=1}^{n} \mathbf{x}_{i}$$

You will notice that the variance and standard deviation are directly related to each other. Why is the term squared? First, because squared terms always give a positive value and we are only trying to measure the distance from the mean (positives and negatives will negate each other). Second, squaring emphasizes the larger terms (or differences) between the mean and each individual data point.

Standard Deviation

Standard deviation describes how far the data points spread away from the mean. It is simply the square root of the variance. All data points are considered by calculating the standard deviation:

$$s = \sqrt{\frac{1}{n} \sum_{i=1}^{n} (x_1 \cdot \overline{x})^2}$$

Where: *n* is the number of values in the data set.

$$\bar{\mathbf{x}} = \frac{1}{n} \sum_{i=1}^{n} \mathbf{x}_{i}$$

2.2.3 NORMAL DISTRIBUTION AND NORMALITY

What is Normal Distribution?

Normal distribution, also called Gaussian distribution, is the probability distribution of a continuous random variable whose values spread symmetrically around the mean. A normal distribution can be completely described by using its mean (μ) and variance (σ^2).

When a variable x is normally distributed, we note $x \sim N(\mu, \sigma^2)$.

Z Distribution

The Z distribution, also known as standard normal distribution, is the simplest normal distribution with the mean equal to zero and the variance equal to one. Any normal distribution can be transferred to a Z distribution by applying the following equation:

$$Z = \frac{x - \mu}{\sigma}$$

Where: $x \sim N(\mu, \sigma^2)$ and $\sigma \neq 0$.

The standard normal distribution is important because the probabilities and quantiles of any normal distribution can be computed from the standard normal distribution if the mean and standard deviation are known.

Shape of Normal Distribution

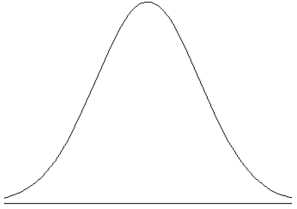


Fig. 2.25 Shape of a Normal Distribution

You might remember teachers from your academic days referring to "the curve" when talking about how they will grade performance. The probability density function curve of normal distribution is bell-shaped (see figure 2.25) and is expressed by the equation:

$$f(x) = \frac{1}{\sqrt{2\pi\sigma^2}} e^{\frac{(x-\mu)^2}{2\sigma^2}}$$

The equation just means that within the population that is described by this distribution, the highest probability is for values closest to the mean, then the probability drops as you move further from the mean (in either direction).

Location of Normal Distribution

If a variable is normally distributed, the mean, the median, and the mode have approximately the same value. The probability density curve of normal distribution is symmetrical around a center value which is the mean, median, and mode at the same time.

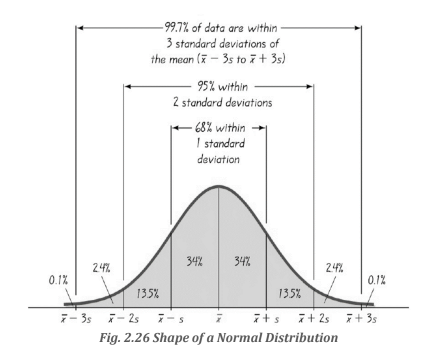
Spread of Normal Distribution

The spread or variation of the normally-distributed data can be described using the variance or the standard deviation (which is the square root of the variance). The smaller the variance or the standard deviation, the less variability in the data set.

The Empirical Rule (68–95–99.7 Rule)

The *68–95–99.7 rule* or the *empirical rule in statistics* states that for a normal distribution:

- About 68% of the data fall within one standard deviation of the mean
- About 95% of the data fall within two standard deviations of the mean
- About 99.7% of the data fall within three standard deviations of the mean



One way to think about this rule is to consider the area under the normal distribution curve. The 68–95–99.7 rule describes what percent of data fall within one, two, and three standard deviations from the mean, respectively. Most of the data values are grouped around the center.

Normality

Not all the distributions with a bell shape are normal distributions. It is important to understand whether or not data being analyzed fit the normal distribution because there are implications for how the data are analyzed and what inferences are made from the data.

To check whether a group of data points are normally distributed, we need to run a normality test. There are different normality tests available, such as Anderson–Darling test, Sharpiro–Wilk test, or Jarque–Bera test. More details of normality test will be introduced in the Analyze module.

Normality Testing

To check whether the population of our interest is normally distributed, we need to run normality test.

- Null Hypothesis (H₀): The data points *are* normally distributed.
- Alternative Hypothesis (H_a): The data points are *not* normally distributed.

Use Minitab to Run a Normality Test

Normality Test Case study: Using a data set provided, we are interested to know whether the height of basketball players is normally distributed. This example demonstrates how to perform this analysis using Minitab.

Introde .

Data File: "One Sample T-Test" tab in "Sample Data.xlsx"

- Null Hypothesis (H₀): The height of basketball players is normally distributed.
- Alternative Hypothesis (H_a): The height of basketball players is not normally distributed.

Steps to run a normality test in Minitab:

- 1. Click Stat \rightarrow Basic Statistics \rightarrow Normality Test.
- 2. A new window named "Normality Test" pops up.

3. Select "HtBk" as the "Variable."

Normality Test	
	Variable: HtBk Percentile Lines (None At Y values: At data values: Tests for Normality (Anderson-Darling Ryan-Joiner (Similar to Shapiro-Wilk) Kolmogorov-Smirnov
Select	Title:
Help	OK Cancel

Fig. 2.27 Normality Variable Selection

- 4. Click "OK."
- 5. The normality test results appear in the new window.

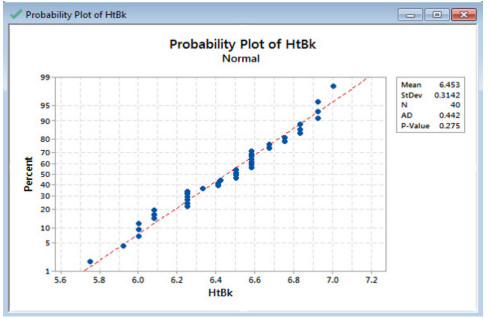


Fig. 2.28 Normality Test Output

Figure 2.28 shows the output from Minitab. We need to know the p-value to determine if we should accept or reject the null hypothesis. The p-value is 0.275 and is greater than alpha (or significance level). Since p > 0.05, we fail to reject the null hypothesis, which was that the data is normal. Therefore, *the data are normally distributed*.

2.2.4 GRAPHICAL ANALYSIS

What is Graphical Analysis?

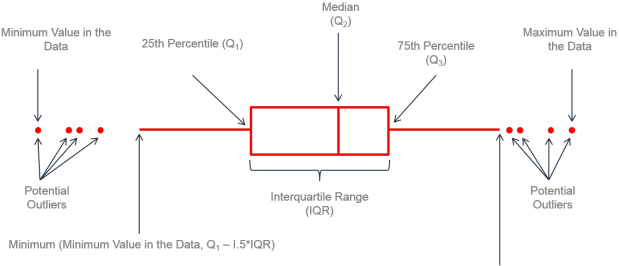
In statistics, *graphical analysis* is a method to visualize the quantitative data. Graphical analysis is used to discover structure and patterns in the data. The presence of which may explain or suggest reasons for additional analysis or consideration. A complete statistical analysis includes both quantitative analysis and graphical analysis.

When we think of statistics, we often think of numbers. However, graphical analysis is complementary to purely quantitative statistics because it allows us to visualize structure and patterns in the data. Often, the graphical presentation of data can tell a story on its own, but the total statistical analysis is important to complete the picture. Sometimes our eyes can deceive us. There are various graphical analysis tools available. The most commonly used and hence the ones we shall cover here are Box Plot, Histogram, Scatter Plot and Run Chart.

Box Plot

A *box plot* is a graphical method to summarize a data set by visualizing the minimum value, 25th percentile, median, 75th percentile, the maximum value, and potential outliers. A percentile is the value below which a certain percentage of data fall. For example, if 75% of the observations have values lower than 685 in a data set, then 685 is the 75th percentile of the data. At the 50th percentile, or median, 50% of the values are lower and 50% are higher than that value.

Box Plot Anatomy



Maximum (Maximum Value in the Data, $Q_3 + I.5*IQR$)

Fig. 2.29 Interpreting a Box Plot

Figure 2.29 describes how to read a box plot. Here are a few explanations that may help. The middle part of the plot, or the "interquartile range," represents the middle quartiles (or the 75th minus the 25th percentile). The line near the middle of the box represents the median (or middle value of the data set). The whiskers on either side of the IQR represent the lowest and

highest quartiles of the data. The ends of the whiskers represent the maximum and minimum of the data, and the individual dots beyond the whiskers represent outliers in the data set.

How to Use Minitab to Generate a Box Plot

Data File: "Box Plot" tab in "Sample Data.xlsx"

Steps to render a box plot in Minitab:

- 1. Click Graph \rightarrow Boxplot.
- 2. A new window named "Boxplots" pops up.

Boxplots				×
One Y Simple	With Groups			
Multiple Y's Simple	With Groups $\begin{array}{ c c c c c } \hline U & U & U & U \\ \hline U & U & U & U \\ \hline U & U & U & U \\ \hline U & U & U & U \\ \hline A & \frac{1}{12} & \frac{1}{12} & \frac{1}{12} \\ \hline A & \frac{1}{12} & \frac{1}{12} & \frac{1}{12} \end{array}$			
Help		<u>O</u> K	Cano	el

Fig. 2.30 Boxplot Dialog Screen

- 3. Click "OK" in the window "Boxplots."
- 4. Another new window named "Boxplot One Y, Simple" pops up.

Boxplot: One	Y, Simple			— X
C1 HtBk		<u>G</u> raph variables: HtBk		* *
		<u>S</u> cale <u>M</u> ultiple Graphs	Labels D <u>a</u> ta Options	Data View
S	elect			
Help			<u>O</u> K	Cancel

Fig. 2.31 Boxplot – One Y, Simple

- 5. Select "HtBk" as the "Graph Variables."
- 6. Click the "Data View" button and a new window named "Boxplot Data View" pops up.
- 7. Check the boxes "Median symbol" and "Mean symbol."

8. Click "OK" in the window "Boxplot – Data View."

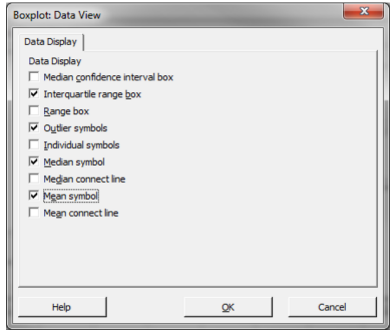


Fig. 2.32 Boxplot - Data View

- 9. Click "OK" in the window "Boxplot One Y, Simple."
- 10. The box plot appears automatically in the new window.

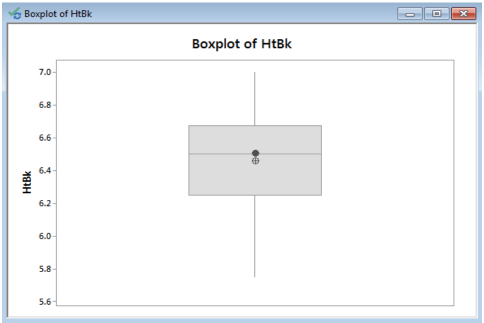


Fig. 2.33 Boxplot Output

Figure 2.33 demonstrates the result of the boxplot after navigating through the Minitab menus to yield this output. Notice the interquartile range between the 25th and 75th quartiles, the median line, the mean, and the whiskers.

Histogram

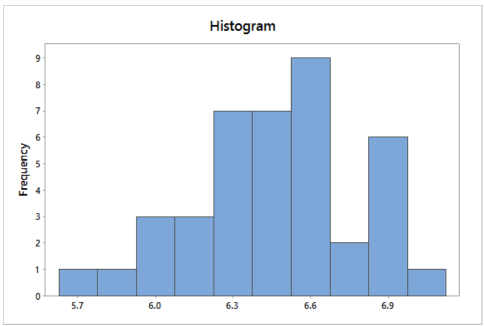


Fig. 2.34 Simple Histogram Example

A *histogram* is a graphical tool to present the distribution of the data. The X axis of a histogram represents the possible values of the variable and the Y axis represents the frequency of the value occurring. A histogram consists of adjacent rectangles erected over intervals with heights equal to the frequency density of the interval. The total area of all the rectangles in a histogram is the number of data values.

A histogram can also be normalized. In the case of normalization, the X axis still represents the possible values of the variable, but the Y axis represents the percentage of observations that fall into each interval on the X axis. The total area of all the rectangles in a normalized histogram is 1. Using histograms, we have a better understanding of the shape, location, and spread of the data.

How to Use Minitab to Generate a Histogram

Data File: "Histogram" tab in "Sample Data.xlsx"

Steps to render a histogram in Minitab:

- 1. Click Stat \rightarrow Basic Statistics \rightarrow Graphical Summary.
- 2. A new window named "Graphical Summary" pops up.

3. Select "HtBk" as the "Variables."

Graphical Summary	X
C1 HtBk	Variables:
	~
	By variables (optional):
	Confidence level: 95.0
Select	
Help	OK Cancel

Fig. 2.35 Graphical Summary Dialog Box Variable

- 4. Click "OK."
- 5. The histogram appears in the new window.

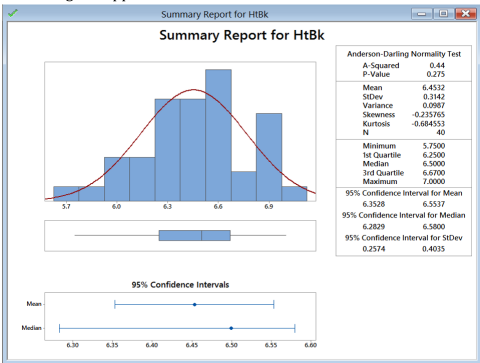


Fig. 2.36 Graphical Summary Output - Histogram

The output from the previous steps has generated a graphical summary report of the data set HtBk. Among the information provided is a histogram. The image shows the frequency of the data for the numerical categories ranging from 5.7 to approximately 6.9. You can see the shape of the data roughly follows the bell curve.

Scatter Plot

A *scatter plot* is a diagram to present the relationship between two variables of a data set. A scatter plot consists of a set of data points. On the scatter plot, a single observation is presented by a data point with its horizontal position equal to the value of one variable and its vertical position equal to the value of the other variable. A scatter plot helps us to understand:

- Whether the two variables are related to each other or not
- What the strength of their relationship
- The shape of their relationship
- The direction of their relationship
- Whether outliers are present

How to Use Minitab to Generate a Scatter Plot

Data File: "Scatter Plot" tab in "Sample Data.xlsx"

Steps to render a Scatterplot in Minitab:

- 1. Click Graph \rightarrow Scatterplot.
- 2. A new window named "Scatterplots" pops up.

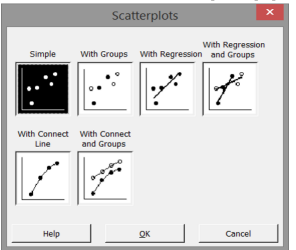


Fig. 2.37 Scatterplots Dialog Box

- Leaving "Simple" selected, click "OK" A new window "Scatterplot – Simple" pops up.
- 4. Select "MPG" as the "Y variables."

5. Select "weight" as the "X variables."

Scatterplot: Simple			
C1 MPG C2 weight	Y variables 1 MPG 2 - 3 - 4 - 5 - 6 - 7 -	X variables weight	Data View
		0 K	Cancel
Help		<u>O</u> K	

Fig. 2.38 Scatterplots - Simple X & Y Variables

- 6. Click "OK."
- 7. The scatter plot appears in the new window.

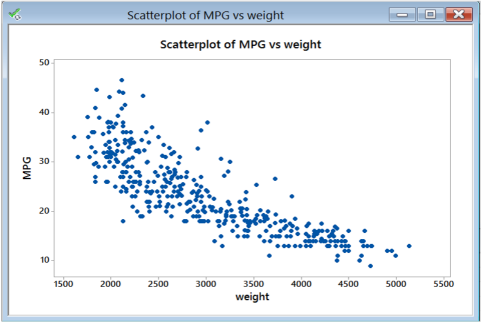


Fig. 2.39 Scatterplot

Figure 2.39 is Minitab's output of the scatterplot data. You can immediately see the value of graphical displays of data. This information obtainable by viewing this output shows a relationship between weight and MPG. This scatterplot shows that the heavier the weight the lower the MPG value and vice versa.

Run Chart

A *run chart* is a chart used to present data in time order. Run charts capture process performance over time. The X axis of a run chart indicates time and the Y axis shows the observed values. A run chart is similar to a scatter plot in that it shows the relationship

between X and Y. Run charts differ however because they show how the Y variable changes with an X variable of time.

Run charts look similar to control charts except that run charts do not have control limits and they are much easier to produce thank a control chart. A run chart is often used to identify anomalies in the data and discover pattern over time. They help to identify trends, cycles, seasonality and other anomalies.

How to Plot a Run Chart in Minitab

Steps to plot a run chart in Minitab:

Data File: "Run Chart" tab in "Sample Data.xlsx"

- 1. Click Stat \rightarrow Quality Tools \rightarrow Run Chart.
- 2. A new window named "Run Chart" pops up.
- 3. Select "Measurement" as the "Single Column."
- 4. Enter "1" as the "Subgroup Size."

Run Chart		
C1 Measuremen C2 Cycle C3 Trend	t Data are arranged as Single column: Measurement Subgroup size: 1 (use a constant or an ID column) Subgroups across rows of: For data in subgroups Plot subgroup means Plot subgroup medians	Options
Select Help		OK Cancel

Fig. 2.40 Run Chart Variable Selection

5. Click "OK."

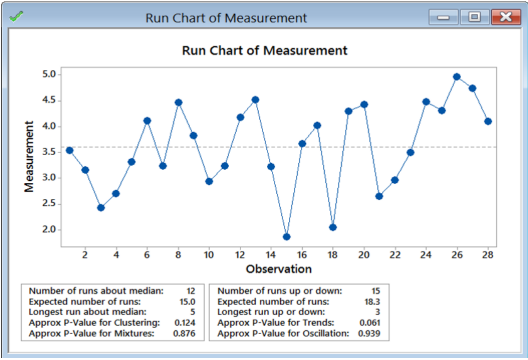


Fig. 2.41 Minitab Run Chart Output

Figure 2.41 is a run chart created with Minitab. The time series displayed by this chart appears stable. There are no extreme outliers, no visible trending or seasonal patterns. The data points seem to vary randomly over time.

Now, let us look at another example which may give us a different perspective. We will create another run chart using the data listed in the column labeled "Cycle". This column is in the same file used to generate Figure 2.41. Follow the steps used for the first run chart and instead of using "Measurement" use "Cycle" in the Run Chart dialog box pictured in figure 2.40.

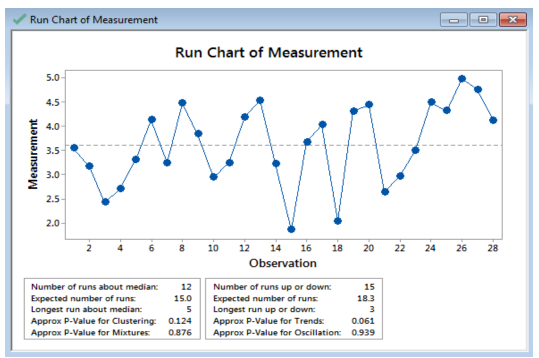


Fig. 2.42 Minitab Run Chart for the "Cycle" data

In this example (figure 2.42), the data points are clearly exhibiting a pattern. It could be seasonal or it could be something cyclical. Imagine that the data points are taken monthly and this is a process performing over a period of 2.5 years. Perhaps the data points represent the number of customers buying new homes. The home buying market tends to peak in the summer months and dies down in the winter.

Using the same data tab lets create a final run chart. This time use the "Trend" data. Again, follow the steps outlined previously to generate a run chart.

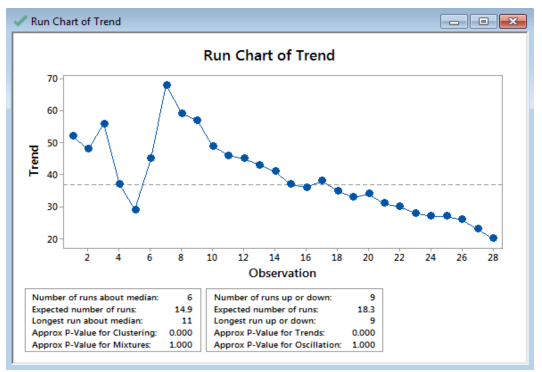


Fig. 2.43 Run Chart Using Trend Data

In this example, the process starts out randomly, but after the seventh data point almost every data point has a lower value than the one before it. This clearly illustrates a downward trend. What might this represent? Perhaps a process winding down? Product sales at the end of a product's life cycle? Defects decreasing after introducing a process improvement?

It should be clearly evident through our review of Histograms, Scatterplots and Run Charts, that there is great value in "visualizing" the data. Graphical displays of data can be very telling and offer excellent information.

2.3 MEASUREMENT SYSTEM ANALYSIS

2.3.1 PRECISION AND ACCURACY

What is Measurement System Analysis?

Measurement System Analysis (MSA) is a systematic method to identify and analyze the variation components of a measurement system. It is a mandatory step in any Six Sigma project to ensure the data are reliable before making any data-based decisions. A MSA is the check point of data quality before we start any further analysis and draw any conclusions from the data. Some good examples of data-based analysis where MSA should be a prerequisite:

- Correlation analysis
- Regression analysis
- Hypothesis testing

- Analysis of variance
- Design of experiments
- Statistical process control

You will see where and how the analysis techniques listed above are used throughout this training program. It is critical at this stage however to know that any variation, anomalies, or trends found in your analysis are due to the data and not due to the inaccuracies or inadequacies of a measurement system. Therefore, the need for a MSA is vital.

Measurement System

A measurement system is a process used to obtain data and quantify a part, product or process. Data obtained with a measurement device or measurement system are the *observed* values. Observed values are comprised of two elements

- 1. True Value = Actual value of the measured part
- 2. Measurement Error = Error introduced by the measurement system.

The *true value* is what we are ultimately trying to determine through the measurement system. It reflects the true measurement of the part or performance of the process.

Measurement error is the variation introduced by the measurement system. It is the bias or inaccuracy of the measurement device or measurement process.

The *observed value* is what the measurement system is telling us. It is the measured value obtained by the measurement system. Observed values are represented in various types of measures which can categorized into two primary types discrete and continuous. Continuous measurements are represented by measures of weight, height, money and other types of measures such as ratio measures. Discrete measures on the other hand are categorical such as Red/Yellow/Green, Yes/No or Ratings of 1–10 for example.

A *variable MSA* is designed assess continuous measurement systems while *attribute MSAs* have been designed to assess discrete measurement systems.

Sources of measurement errors

There are numerous ways to introduce error into a measurement system. The following are just a few common ones:

- Human Error imagine reading a ruler or a scale, or think about systems where a person must make a qualitative judgment
- Environment Sometimes the temperature of the air or humidity can affect how a measurement system reads or the equipment itself may need to be calibrated from time to time
- Equipment
- Sample Sampling can introduce variation based on how or where or when we choose to sample a process
- Process
- Materials
- Methods

Fishbone diagrams can help to brainstorm potential factors affecting the accuracy or validity of measurement systems. The more errors a measurement system allows, the less reliable the observed values are. A valid measurement system brings in minimum amount of measurement error. Ultimately, the goal of a measurement system analysis is to understand how much error exists in the system by analyzing the characteristics of the system. Then, decisions can be made as to whether the system needs to be changed, replaced, modified or accepted.

Characteristics of a Measurement System

Any measurement systems can be characterized by two aspects:

- Accuracy (location related)
- Precision (variation related)

A valid measurement system is *both* accurate and precise. Being accurate does not guarantee the measurement system is precise. Being precise does not guarantee the measurement system is accurate.

Accuracy vs. Precision

Accuracy is the level of closeness between the average observed value and the true value. Accuracy tells us how well the observed value reflects the true value.

Precision is the spread of measurement values. It is how consistent the repeated measurements deliver the same values under the same circumstances.

To simplify, *accuracy* describes how close the average observed value is to the true value for what is being measured. *Precision* describes how close or repeatable the measurement is. Let us take a look at a few representations of accuracy and precision.

Accurate and precise (high accuracy and high precision)

Figure 2.44 represents a target with shots landing in the center and all shots close to one another. This is a depiction of both accuracy and precision. For the purposes of an MSA, think of the center of the target as the "true value" and the black dots as "observed values." All six measurements are located at the center of the target and are grouped tightly together. This means that the measurements are both accurate and precise.



Fig. 2.44 Accurate and Precise

Accurate and not precise (high accuracy and low precision) Figure 2.45 is an example with all the observations located around the center of the target, but they are spread further apart from one another, more so than in figure 2.44. This measurement system would be described as accurate, but not precise.



Fig. 2.45 Accurate but Not Precise

Precise and not accurate (high precision and low accuracy) Figure 2.46 shows all observations grouped closely together, but they are not located near the center of the target. This system is precise, but not accurate.



Fig. 2.46 Precise but Not Accurate

Not accurate and not precise (low accuracy and low precision) Lastly, figure 2.47 shows a system that is neither accurate nor precise. The observations are not located around the center and they are spread far apart from each other.



Fig. 2.47 Not Accurate and Not Precise

MSA Conclusions

If the measurement system is considered *both* accurate and precise, we can start the databased analysis or decision making. If the measurement system is either not accurate or not precise, we need to identify the factors affecting it and calibrate the measurement system until it is both accurate and precise.

Stratifications of Accuracy and Precision

- Accuracy
 - o Bias
 - o Linearity
 - \circ Stability
- Precision
 - Repeatability
 - Reproducibility

2.3.2 BIAS, LINEARITY, AND STABILITY

Bias

Bias is the difference between the observed value and the true value of a parameter or metric being measured. Think of it as a measurement system's absolute correctness versus a standard (or true value). Bias is calculated by subtracting the reference value from the average value of the measurements.

Bias = Grand Mean-Reference Value

where the reference value is a standard agreed upon value.

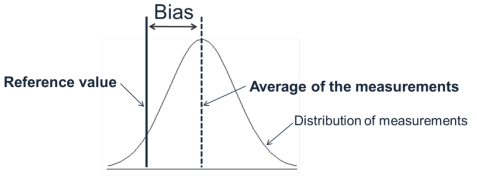


Fig. 2.48 Measurement Bias

In figure 2.48 you can see that the group of observations is consistently to the right of the reference value. The dotted line represents the grand mean described on the last page. The goal is to reduce the distance between the grand mean and the reference value, thereby reducing bias. The closer the average of all measurements is to the reference level, the smaller the bias. The reference level is the average of measurements of the same items using the master or standard instrument.

To determine whether the difference between the average of observed measurement and the reference value is statistically significant (we will explain more details about statistical significance in the Analyze module), we can either conduct a *hypothesis test* or *compare* the reference value against the confidence intervals of the average measurements. If the reference

value falls into the confidence intervals, the bias is not statistically significant and can be ignored. Otherwise, the bias is statistically significant and must be fixed.

Potential causes of bias are:

- Errors in measuring the reference value
- Lack of proper training for appraisers
- Damaged equipment or instrument
- Measurement instrument not calibrated precisely
- Appraisers read the data incorrectly

In the first case, for example, there could be something wrong with the standard (or reference value). For example, think about weight standards that are used to calibrate scales. If the weight is damaged in some way such that it no longer represents the standard, then obviously, we will not be able to measure the correct value.

Linearity

Linearity is the degree of the consistency of bias over the entire expected measurement range. It quantifies how the bias changes over the range of measurement. For example, a scale is off by 0.01 pounds when measuring an object of 10 pounds. However, it is off by 10 pounds when measuring an object of 100 pounds. The scale's bias is not linear.

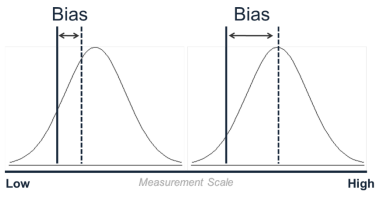


Fig. 2.49 Measurement Linearity

Figure 2.49 represents the bias of a measurement system across its measurement scale from low measures to high measures. The image suggests that as the measurement scale increases so does the bias. The linearity for this measurement system would be positive and undesirable.

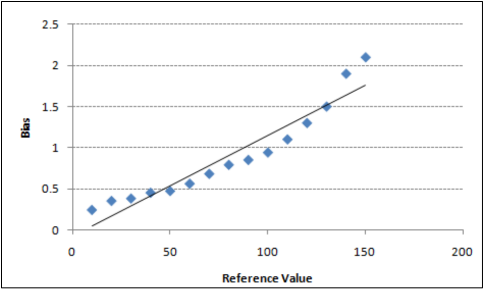


Fig. 2.50 Bias Scatterplot for Linearity

Figure 2.50 uses a scatterplot to demonstrate the linearity of a measurement system. The X axis represents references values for the measurement scale and the Y axis represents bias. The best fit line shows the slope of the bias. The ideal linearity of a measurement system would have a slope of zero which would imply no change in bias over the scale of the measurement system. The slope in figure 2.50 is positive suggesting that as the measurement scale increases so does the bias.

The formula of the linearity of a measurement system is:

Where:

Slope =
$$\frac{\sum_{i=1}^{n} x_i y_i - \frac{1}{n} \sum_{i=1}^{n} x_i \sum_{i=1}^{n} y_i}{\sum_{i=1}^{n} (x_i^2) - \frac{1}{n} (\sum_{i=1}^{n} x_i)^2}$$

and where:

- *x_i* is the reference value
- *y_i* is the bias at each reference level
- *n* is the sample size.

As demonstrated in the previous figures, we know that slope is an important computation in determining the linearity of a measurement system. This is based on the fact that the equation of a line defines the relationship between the bias and the reference values of the parts or samples.

Potential causes of linearity

The causes of linearity include:

- Errors in measuring the lower end or higher end of the reference value
- Lack of proper training for appraisers
- Damaged equipment or instrument
- Measurement instrument not calibrated correctly at the lower or higher end of the measurement scale
- Innate nature of the instrument

As you notice, some of the things we discussed as causes for bias are also potential causes for linearity issues: training, damaged or improperly-calibrated equipment. How these issues manifest themselves are the difference, because bias can become an issue at the low and high ends of a measurement range.

Stability

Stability is the consistency level needed to obtain the same values when measuring the same objects over an extended period. A measurement system that has low bias and linearity close to zero but cannot consistently perform well over time would not deliver reliable data.

Much like we discussed linearity being the degree of bias over the range of a measurement scale, stability is the degree of bias over time. Figure 2.50 demonstrates increasing bias as time passes.

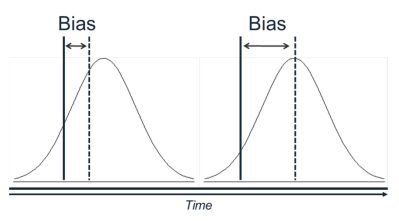


Fig. 2.51 Measurement Stability: Bias over Time

Control charts are the most common tools used to evaluate the stability of a measurement system. By plotting bias calculations of a measurement system over time and using a control chart to visualize scale and variation, it allows us to determine if bias is stable or out of control. As you will learn in the Control phase, if no measures on a control chart are "out of control" then we can consider bias to be stable.

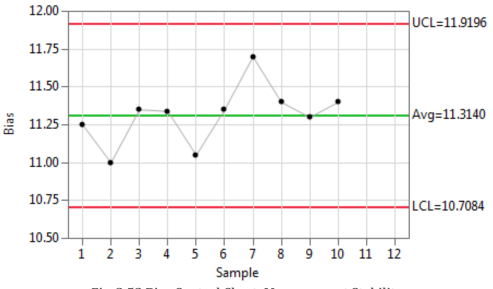


Fig. 2.52 Bias Control Chart: Measurement Stability

Potential causes of instability

The causes of instability include:

- Inconsistent training for appraisers
- Damaged equipment or instrument
- Worn equipment or instrument
- Measurement instrument not calibrated
- Appraisers do not follow the procedure consistently

Consider two appraisers who were trained at two different times, and they would measure differently, or consider two appraisers who do not follow the procedures in the same way. Measurement equipment can wear out over time, lose calibration, or even break, obviously affecting its ability to measure accurately over time.

2.3.3 GAGE REPEATABILITY AND REPRODUCIBILITY

Repeatability

Repeatability evaluates whether the same appraiser can obtain the same value multiple times when measuring the same object using the same equipment under the same environment. It refers to the level of agreement between the repeated measurements of the same appraiser under the same condition. Repeatability measures the inherent variation of the measurement instrument. To simplify repeatability, think of it as your own personal ability to measure something 3 times and get the same answer all three times. If you can't then you're unable to successfully "repeat" a measurement.

Reproducibility

Reproducibility evaluates whether different appraisers can obtain the same value when measuring the same object independently. It refers to the level of agreement between different appraisers. While repeatability is inherent to the measurement system, reproducibility is not caused by the inherent variation of the measurement instrument. It reflects the variability caused by different appraisers, locations, gauges, environments etc. The simple explanation of reproducibility is best described by asking, can you and someone else measure something and get the same answer. If not, the reproducibility of the measurement system is inadequate.

Gage Repeatability & Reproducibility

Gage R&R (Gage Repeatability & Reproducibility) is a method used to analyze the variability of a measurement system by partitioning the variation of the measurements using ANOVA (Analysis of Variance). Gage R&R primarily addresses the precision aspect of a measurement system. It is a tool used to understand if a measurement system can repeat and reproduce and if not, help us determine what aspect of the measurement system is broken so that we can fix it.

Gage R&R requires a deliberate study with parts, appraisers and measurements. Measurement data must be collected and analyzed to determine if the measurement system is acceptable. Typically Gage R&Rs are conducted by 3 appraisers measuring 10 samples 3 times each. Then, the results can be compared to determine where the variability is concentrated. The optimal result is for the measurement variability to be due to the parts.

Data collection of a gage R&R study:

Let k appraisers' measure n random samples independently and repeat the process p times. Different appraisers perform the measurement independently. The order of measurement (e.g., sequence of samples and sequence of appraisers) is randomized, as evidenced by the representative letters k, n, and p.

Potential sources of variance in the measurement: Appraisers: $\sigma^2_{appraisers}$

Parts: σ_{parts}^2

Appraisers × Parts: $\sigma^2_{appraisers \times parts}$

Repeatability: $\sigma^2_{repeatability}$

Variance Components: $\sigma_{total}^2 = \sigma_{appraisers}^2 + \sigma_{parts}^2 + \sigma_{appraisers \times parts}^2 + \sigma_{repeatability}^2$

As can be seen in this equation for total measurement variance, there are several components: the appraisers, the parts being measured, the interaction between the appraisers and parts, and the repeatability. A valid measurement system has low variability in both repeatability and

reproducibility so that the total variability observed can reflect the true variability in the parts being measured.

$$\sigma_{total}^{2} = \sigma_{reproducibility}^{2} + \sigma_{repeatability}^{2} + \sigma_{parts}^{2}$$

Where:

$$\sigma^2_{reproducibility} = \sigma^2_{appraisers} + \sigma^2_{appraisers \times parts}$$

Gage R&R variance reflects the precision level of the measurement system.

$$\sigma^2_{R\&R} = \sigma^2_{repeatability} + \sigma^2_{reproducibility}$$

As you can see, we can also show the total variance of the measurement system is a function of the reproducibility, the repeatability, and the variance in parts. It ties back to the equation on the previous page when you substitute variance in the appraisers and the variance in interaction between appraisers and parts with reproducibility. The goal is to minimize reproducibility variance and repeatability variance so you are only left with variance in parts—the variance in the parts is the "true" variance.

The *precision* of the measurement system is represented by the sum of the repeatability variance and reproducibility variance.

Variation Components

The variation of each component can be represented by the standard deviation (or sigma) for each component multiplied by Z₀, which is a sigma multiplier that assumes a specific confidence level for the spread of the data. This is important because we will use the variation to calculate each component's contribution to the total measurement variation of the system.

Variation_{total} = $Z_0 \times \sigma_{total}$

 $Variation_{repeatability} = Z_0 \times \sigma_{repeatability}$

 $Variation_{reproducibility} = Z_0 \times \sigma_{reproducibility}$

 $Variation_{parts} = Z_0 \times \sigma_{parts}$

Where:

$$\sigma_{total}^2 = \sigma_{reproducibility}^2 + \sigma_{repeatability}^2 + \sigma_{parts}^2$$

The percentage of variation R&R contributes to the total variation in the measurement:

Contribution
$$\%_{R\&R} = \frac{Variation_{R\&R}}{Variation_{total}} \times 100\%$$

Where:

$$Variation_{R\&R} = Z_0 \times \sqrt{\sigma_{repeatability}^2 + \sigma_{reproducibility}^2}$$

The goal for the Contribution $\mathcal{W}_{R\&R}$ calculation is to determine how much variation is contributed by repeatability and reproducibility. As you can imagine, the lower the contribution from these components the better, but there are some guidelines for how to assess the quality of the measurement system. Table 2.1 represents the standard interpretations of varying degrees of contribution percentages.

Contribution %R&R	Interpretation
Less than 1%	Satisfactory – Measurement System is Acceptable
Between 1% and 9%	Grey Area – Requires Decisions and Trade-offs
Greater than 10%	Unsatisfactory – Measurement System is Unacceptable

Table 2.1 Gage R&R Contribution% Interpretation Guidelines

2.3.4 VARIABLE AND ATTRIBUTE MSA

Variable Gage R&R

Whenever something is measured repeatedly or by different people or processes, the results of the measurements will vary. Variation comes from two primary sources:

- Differences between the parts being measured
- The measurement system

We can use a gage R&R to conduct a measurement system analysis to determine what portion of the variability comes from the parts and what portion comes from the measurement system. There are key study results that help us determine the components of variation within our measurement system.

Key Measures of a Variable Gage R&R

- %Contribution The percent of contribution for a source is 100 times the variance component for that source divided by the total variance.
- %Study Var (6*SD) The percent of study variation for a source is 100 times the study variation for that source divided by the total variation
- %Tolerance (SV/Tolerance) The percent of spec range taken up by the total width of the distribution of the data based on variation from that source
- Distinct Categories The number of distinct categories of parts that the measurement system is able to distinguish. If a measurement system is not capable of distinguishing at least five types of parts, it is probably not adequate.

Variable Gage R&R Guidelines (AIAG)

The guidelines for acceptable or unacceptable measurement systems can vary depending on an organizations tolerance or appetite for risk. The common guidelines used for interpretation are published by the Automotive Industry Action Group (AIAG). These guidelines are considered standard for interpreting the results of a measurement system analysis using Variable Gage R&R. Table 2.2 summarizes the AIAG standards.

Measurement System	% Study Var	% Contribution	Distinct Categories
Acceptable	10% or less	1% or Less	5 or Greater
Marginal	10% - 30%	1% - 9%	
Unacceptable	30% or Greater	9% or Greater	Less than 5

Table 2.2 Variable Gage R&R Guidelines

Use Minitab to Implement a Variable MSA

Data File: "Variable MSA" tab in "Sample Data.xlsx"

Let's take a look at an example of a Variable MSA using the data in the Variable MSA tab in your "Sample Data.xlsx" file. In this exercise, we will first walk through how to set up your study using Minitab and then we will perform a Variable MSA using 3 operators who all measured 10 parts three times each. The part numbers and operators and measurement trials are all generic so that you can apply the concept to your given industry. First, we need to set up the study:

- 1. Click on Stat \rightarrow Quality Tools \rightarrow Gage R&R \rightarrow Create Gage R&R Study Worksheet.
- 2. A new window named "Create Gage R&R Study Worksheet" pops up.

Create Gage R&R Study Worksheet						
Number	of parts: 10 •	Number of operators:	3 • Options			
Part	Part Name	Operator	Operator Name			
1	1	1	Operator A			
2	2	2	Operator B			
3	3	3	Operator C			
4	4		,			
5	5					
6	6					
7	7					
8	8					
9	9					
10	10					
Number of replicates:						

Fig. 2.53 "Create Gage R&R Study Worksheet" dialog box

3. Select 10 as the "Number of Parts." Select 3 as the "Number of Operators." Select 3 as the "Number of Replicates." Enter the part name (e.g., Part 01, Part 02, and Part 03). Enter the operator name (e.g., Operator A, Operator B, Operator C). Click on the "Options" button, another window named "Create Gage R&R Study Worksheet – Options" pops up.

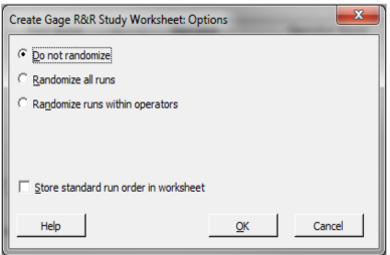


Fig. 2.54 complete "Create Gage R&R Study Worksheet" dialog box

- 4. Select the radio button "Do not randomize."
- 5. Click "OK" in the window "Create Gage R&R Study Worksheet Options."
- 6. Click "OK' in the window "Create Gage R&R Study Worksheet."
- 7. A new data table is generated.

Uorksheet 2 ***					
÷	C1	C2-T	C3-T	C4	
	RunOrder	Parts	Operators	Measurement	
4	4	2	Operator A		
5	5	2	Operator B		
6	6	2	Operator C		
7	7	3	Operator A		
8	8	3	Operator B		
9	9	3	Operator C		
10	10	4	Operator A		
11	11	4	Operator B		
12	12	4	Operator C		
13	13	5	Operator A		
14	14	5	Operator B		

Fig. 2.55 New Minitab Worksheet Ready to Populate with Variable MSA Study Results

Step 2: Data collection

In the newly-generated data table, Minitab has provided the data layout for your data collection for your variable MSA study. We have added the header "Measurement" for this example. You would have to do something similar.

When you conduct your variable MSA in your work environment it would be necessary to set up your study just as we have in the previous steps and then you could collect your measurement data properly. However, for our purposes today, we have provided you with an MSA that is setup with data already collected. We will use our "Variable MSA" tab in "Sample Data.xlsx," for the next steps.

÷	C1	C2-T	С3-Т	C4	^
	Run Order	Part	Operator	Measurement	
1	1	Part 01	Operator A	0.29	
2	2	Part 01	Operator A	0.41	
3	3	Part 01	Operator A	0.64	
4	4	Part 02	Operator A	-0.56	
5	5	Part 02	Operator A	-0.68	
6	6	Part 02	Operator A	-0.58	
7	7	Part 03	Operator A	1.34	
8	8	Part 03	Operator A	1.17	
9	9	Part 03	Operator A	1.27	
10	10	Part 04	Operator A	0.47	
11	11	Part 04	Operator A	0.50	
12	12	Part 04	Operator A	0.64	
13	13	Part 05	Operator A	-0.80	
14	14	Part 05	Operator A	-0.92	
15	15	Part 05	Operator A	-0.84	
16	16	Part 06	Operator A	0.02	
17	17	Part 06	Operator A	-0.11	
18	18	Part 06	Operator A	-0.21	~

Fig. 2.56 Pre-Populated Variable MSA data from the Sample Data.xlsx file

Step 3: Activate the Minitab worksheet with our Variable MSA data prepopulated. Step 4: Implement Gage R&R

- 1. Click Stat \rightarrow Quality Tools \rightarrow Gage Study \rightarrow Gage R&R Study (Crossed).
- 2. A new window named "Gage R&R Study (Crossed)" appears.
- 3. Select "Part" as "Part numbers."
- 4. Select "Operator" as "Operators."
- 5. Select "Measurement" as "Measurement data."

Gage R&R Study (Crossed)				
	Part numbers:	Part	Gage Info	
	Operators:	Operator	Options	
	Measurement data:	Measurement	Conf Int	
			Storage	
	 Method of Analysis ANOVA 			
Select				
			OK	
Help			Cancel	

Fig. 2.57 Gage R&R Study (Crossed) dialog box with variables

- 6. Click on the "Options" button and another new window named "Gage R&R Study (Crossed) ANOVA Options" pops up.
- 7. Enter 5.15 as the "Study variation (number of standard deviations)".

The value 5.15 is the recommended standard deviation multiplier by the Automotive Industry Action Group (AIAG). It corresponds to 99% of data in the normal distribution. If we use 6 as the standard deviation multiplier, it corresponds to 99.73% of the data in the normal distribution.

Gage R&R Study (Crossed): ANOVA Options	J
Study variation: 5,15 (number of standard deviations)	
Process tolerance	1
Enter at least one specification limit	
Lower spec:	1
Upper spec:	l
C Upper spec - Lower spec:	
Historical standard deviation:	
Use historical standard deviation to estimate process variation	
Alpha to remove interaction term: 0.05	
I Display probabilities of misclassification	
Do not display percent contribution	
Do not display percent study variation	1
Draw graphs on separate graphs, one graph per page	
Title:	
Help OK Cancel	

Fig. 2.58 Gage R&R Study (Crossed) ANOVA Options dialog box

- 8. Click "OK" in the window "Gage R&R Study (Crossed) ANOVA Options."
- 9. Click "OK" in the window "Gage R&R Study (Crossed)."
- 10. The MSA analysis results appear in the new window and the session window.

Variance Components									
	0	6 Contribution							
Source	VarComp	(of VarComp)	_						
Total Gage R&R	0.09143	7.76							
Repeatability	0.03997	3.39							
Reproducibility	0.05146	4.37							
Operator	0.05146	4.37							
Part-To-Part 1.08645 92.24									
Part-To-Part	1.00045	JZ.2-							
Total Variation	1.17788	100.00							
	1.17788	100.00	%Study Va						
Total Variation	1.17788	100.00 Study Var	-						
Total Variation	1.17788	100.00 Study Var	(%SV						
Total Variation Gage Evaluatio Source	1.17788 DN StdDev (SD)	100.00 Study Var (5.15 × SD)	(%SV 27.8						
Total Variation Gage Evaluatio Source Total Gage R&R	1.17788 Dn StdDev (SD) 0.30237	100.00 Study Var (5.15 × SD) 1.55721 1.02966	(%SV 27.8 18.4						
Total Variation Gage Evaluation Source Total Gage R&R Repeatability	1.17788 Dn StdDev (SD) 0.30237 0.19993	5tudy Var (5.15 × SD) 1.55721 1.02966 1.16821	(%SV 27.8 18.4 20.9						
Total Variation Gage Evaluation Source Total Gage R&R Repeatability Reproducibility	1.17788 Dn StdDev (SD) 0.30237 0.19993 0.22684	5tudy Var (5.15 × SD) 1.55721 1.02966 1.16821 1.16821	%Study Va (%SV 27.8(18.4) 20.9(20.9(96.0-						

Fig. 2.59 Gage R&R Study (Crossed) output

Step 5: Interpret the MSA results

The result of this Gage R&R study leaves room for consideration on one key measure. As noted in previous pages, the targeted percent contribution R&R should be less than 9% and study variation less than 30%. With % contribution at 7.76% it is below our 9% unacceptable threshold and similarly, Study variation at 26.86% is also below the threshold of 30% but this result is at best marginal and should be heavily scrutinized by the business before concluding that the measurement system does not warrant further improvement.

Visual evaluation of this measurement system is another effective method of evaluation but can at times be misleading without the statistics to support it. Diagnosing the mean plots above should help in the consideration of measurement system acceptability, you may benefit from taking a closer look at operator C.

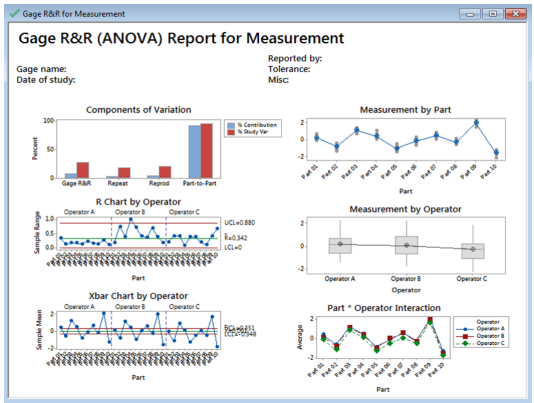


Fig. 2.60 Gage R&R Study (Crossed) additional output

Visual evaluation of this measurement system alone might mislead you to a conclusion of a passing gage study (most of the variation seems to be in part to part, which is what we hope to see). However, an experienced practitioner will note such things such as the range chart being out of control. This may help provide clues regarding what to look for when trying to further diagnose the validity of this measurement system. An out of control Range chart in a variable MSA suggests that an operator or all operators are too inconsistent in their repeated measures causing wide ranges and out of control conditions.

Whenever a range chart is out of control, the accuracy of the X chart is automatically called into question. You will learn in future lessons that the control limits of an X chart are calculated

using the mean of the R chart. If the R chart shows out-of-control conditions, then the mean is likely misrepresented and any calculation using it should be questioned.

Use Minitab to Implement an Attribute MSA

Data File: "Attribute MSA" tab in "Sample Data.xlsx" (an example in the AIAG MSA Reference Manual, 3rd Edition).

Steps in Minitab to run an attribute MSA:

Step 1: Reorganize the original data into four new columns (i.e., Appraiser, Assessed Result, Part, and Reference).

- 1. Click Data \rightarrow Stack \rightarrow Blocks of Columns.
- 2. A new window named "Stack Blocks of Columns" pops up.
- 3. Select "Appraiser A," "Part," and "Reference" as block one.
- 4. Select "Appraiser B," "Part," and "Reference" as block two.
- 5. Select "Appraiser C," "Part," and "Reference" as block three.
- 6. Select the radio button of "New worksheet" and name the sheet "Data."
- 7. Check the box "Use variable names in subscript column."
- 8. Click "OK."

Stack	Blocks of Colum	ins 💌
C1	Part	Stack two or more blocks of columns on top of each other:
C2	Reference	'Appraiser A' Part Reference
C3 C4 C5	Appraiser A Appraiser B Appraiser C	'Appraiser B' Part Reference
	Appraiser C	'Appraiser C' Part Reference
		1
		Store stacked data in:
		New worksheet
,		Name: (Optional)
	- L - L	C Columns of current worksheet:
_	Select	
		Store subscripts in: (Optional)
		✓ Use variable names in subscript column
	Help	OK Cancel

Fig. 2.61 Stack Blocks of Columns dialog box with variables

9. The stacked columns are created in the new worksheet named "Data."

	🛄 Data ***							
Γ	Ŧ	C1-T	C2	C3	C4			
		Subscripts						
L	1	Appraiser A	1	1	1			
	2	Appraiser A	1	1	1			
Γ	3	Appraiser A	1	1	1			
Г	4	Appraiser A	1	2	1			
Г	5	Appraiser A	1	2	1			
Γ	6	Appraiser A	1	2	1			
Γ	7	Appraiser A	0	3	0			
Γ	8	Appraiser A	0	3	0			
	9	Appraiser A	0	3	0			

Fig. 2.62 Stacked Columns output

10. Name the four columns from left to right in worksheet "Data": Appraiser, Assessed Result, Part, and Reference.

🖽 Da	🔲 Data ***									
÷	C1-T	C2	C3	C4						
	Appraiser	Assessed Result	Part	Reference						
1	Appraiser A	1	1	1						
2	Appraiser A	1	1	1						
3	Appraiser A	1	1	1						
4	Appraiser A	1	2	1						
5	Appraiser A	1	2	1						
6	Appraiser A	1	2	1						
7	Appraiser A	0	3	0						
8	Appraiser A	0	3	0						
9	Appraiser A	0	3	0						

Fig. 2.63 Stacked Columns output with named columns

Step 2: Run a MSA using Minitab

- 1. Click Stat \rightarrow Quality Tools \rightarrow Attribute Agreement Analysis.
- 2. A new window named "Attribute Agreement Analysis" pops up.
- 3. Click in the blank box next to "Attribute column" and the variables appear in the list box on the left.
- 4. Select "Assessed Result" as "Attribute column."
- 5. Select "Part" as "Sample."
- 6. Select "Appraiser" as "Appraisers."

7. Select "Reference" as "Known standard/attribute."

Attribute Agreement A	nalysis		X
	Data are arranged as Attribute column: Samples: Appraisers: Multiple columns:	Assessed Result Part Appraiser	Information Options Graphs Results
	(Enter trials for each Number of appraise Number of trials: Appraiser names (or		-
Select	Known standard/attribu	ite: Reference ttribute data are ordere	(Optional)
Help			Cancel

Fig. 2.64 Attribute Agreement Analysis dialog box with variables

- 8. Click the "Options" button and another window named "Attribute Agreement Analysis Options" pops up.
- 9. Check the boxes of both "Calculate Cohen's kappa if appropriate" and "Display disagreement table."

Attribute Agreement Analysis: Options						
Calculate Cohen's kappa if appropriate						
✓ Display disagreement table						
Confidence level: 95.0						
Help OK Cancel						

Fig. 2.65 Attribute Agreement Analysis: Options dialog box with selections

- 10. Click "OK" in the window "Attribute Agreement Analysis Options."
- 11. Click "OK" in the window "Attribute Agreement Analysis."

12. The MSA results appear in the newly-generated window and the session window.

Attribute Agreement Analysis for Assessed Result									
Within Appraisers									
Assessmer	nt Agreem	ent							
Appraiser	# Inspected	# Match	ned	Percent	95%	CI			
Appraiser A	50		42	84.00	(70.89, 9	92.83)			
Appraiser B	50		45	90.00	(78.19, 9	96.67)			
Appraiser C	50		40	80.00	(66.28, 8	89.97)			
# Matched: Ap	praiser agrees w pa Statisti		elf a	cross trials.					
Appraiser	Response	Карра	S	Е Карра	Z	P(vs > 0)		
Appraiser A	0	0.760000	0.0	816497	9.3081	0.000	C		
	1	0.760000	0.0	816497	9.3081	0.000	C		
Appraiser B	0	0.845073	0.0	816497	10.3500	0.000	C		
	1	0.845073	0.0	816497	10.3500	0.000	C		
Appraiser C	0	0.702911	0.0	816497	8.6089	0.000	C		
	1	0.702911	0.0	816497	8.6089	0.000	0		

Fig. 2.66 Attribute Agreement Analysis output (Within Appraisers)

The rater scores represent how the raters agree with themselves. Appraiser A, for instance, agreed with himself on 84% of the measurements made.

Between Appraisers								
Assessment Agreement								
# Inspecte	d # Match	ed Percent	t 95%	CI				
5	0	39 78.00) (64.04, 8	38.47)				
# Matched: All appraisers' assessments agree with each other. Fleiss' Kappa Statistics								
Response	Карра	SE Kappa	Z	P(vs > 0)				
0	0 0.793606 0.0235702 33.6698 0.0000							
0 0.793606 0.0235702 33.6698 0.0000 1 0.793606 0.0235702 33.6698 0.0000								

Fig. 2.67 Attribute Agreement Analysis output (Between Appraisers)

All Appraisers vs Standard									
Assessment Agreement									
# Inspected	d # Match	ed	Percent	95%	CI				
50)	39	78.00	(64.04, 8	38.47)				
	# Matched: All appraisers' assessments agree with the known standard. Fleiss' Kappa Statistics								
Response	Карра	SI	Е Карра	Z	P(vs > 0)				
0	0.859184	0.0	471405	18.2260	0.0000				
1	0.859184	0.0	471405	18.2260	0.0000				
Cohen's I	Cohen's Kappa Statistics								
Response	Карра	SI	Е Карра	Z	P(vs > 0)				
0	0.859313	0.0	470879	18.2491	0.0000				
1	0.859313	0.0	470879	18.2491	0.0000				

Fig. 2.68 Attribute Agreement Analysis output (All Appraisers vs Standard)

The important numbers are called out here. Of the 50 total measurements performed, for 78% of those (#39) the appraisers agreed with both themselves and the other appraisers.

Kappa statistic is a coefficient indicating the agreement percentage above the expected agreement by chance. Kappa ranges from -1 (perfect disagreement) to 1 (perfect agreement). When the observed agreement is less than the chance agreement, Kappa is negative. When the observed agreement is greater than the chance agreement, Kappa is positive. Rule of thumb: If Kappa is greater than 0.7, the measurement system is acceptable. If Kappa is greater than 0.9, the measurement system is excellent.

Attribute Agreement Analysis for Assessed Result

Within Appraisers

Assessment Agreement

Appraiser	# Inspected	# Matched	Percent	95% CI
Appraiser A	50	42	84.00	(70.89, 92.83)
Appraiser B	50	45	90.00	(78.19, 96.67)
Appraiser C	50	40	80.00	(66.28, 89.97)

Matched: Appraiser agrees with him/herself across trials.

Fleiss' Kappa Statistics

Appraiser	Response	Карра	SE Kappa	Z	P(vs > 0)
Appraiser A	0	0.760000	0.0816497	9.3081	0.0000
	1	0.760000	0.0816497	9.3081	0.0000
Appraiser B	0	0.845073	0.0816497	10.3500	0.0000
	1	0.845073	0.0816497	10.3500	0.0000
Appraiser C	0	0.702911	0.0816497	8.6089	0.0000
	1	0.702911	0.0816497	8.6089	0.0000

Fig. 2.69 Kappa Statistic within Appraisers

Each Appraiser vs Standard					
Appraiser	Response	Карра	SE Kappa	Z	P(vs > 0)
Appraiser A	0	0.880236	0.0816497	10.7806	0.0000
	1	0.880236	0.0816497	10.7806	0.0000
Appraiser B	0	0.922612	0.0816497	11.2996	0.0000
	1	0.922612	0.0816497	11.2996	0.0000
Appraiser C	0	0.774703	0.0816497	9.4881	0.0000
	1	0.774703	0.0816497	9.4881	0.0000
Cohen's Kappa Statistics					
Appraiser	Response	Карра	SE Kappa	Z	P(vs > 0)
Appraiser A	0	0.880395	0.0815419	10.7968	0.0000
	1	0.880395	0.0815419	10.7968	0.0000
Appraiser B	0	0.922634	0.0816199	11.3040	0.0000
	1	0.922634	0.0816199	11.3040	0.0000
Appraiser C	0	0.774910	0.0815140	9.5065	0.0000
	1	0.774910	0.0815140	9.5065	0.0000

Fig. 2.70 Kappa Statistic of Each Appraiser vs. Standard

In all cases the Kappa indicates that the measurement system is acceptable.

2.4 PROCESS CAPABILITY

2.4.1 CAPABILITY ANALYSIS

What is Process Capability?

Process capability measures how well the process performs to meet given specified outcome. It indicates the conformance of a process to meet given requirements or specifications. *Capability analysis* helps to better understand the performance of the process with respect to meeting customer's specifications and identify the process improvement opportunities.

Process Capability Analysis Steps

Step 1: Determine the metric or parameter to measure and analyze.

Step 2: Collect the historical data for the parameter of interest.

Step 3: Prove the process is statistically stable (i.e., in control).

Step 4: Calculate the process capability indices.

Step 5: Monitor the process and ensure it remains in control over time. Update the process capability indices if needed.

Process Capability Indices

Process capability can be presented using various indices depending on the nature of the process and the goal of the analysis. Popular process capability indices are:

P_{pk} C_{pm}

- C_p
- Pp
- Cpk

Capability of the Process (Cp)

 C_p can be calculated with the formula:

$$C_{\rm p} = \frac{\rm USL-LSL}{6 \times \sigma_{\rm within}}$$

Where:

$$\begin{split} \sigma_{within} &= \frac{s_p}{c_4(d+1)} \\ s_p &= \sqrt{\frac{\sum_i \sum_j (x_{ij} \cdot \overline{x}_i)}{\sum_i (n_i \cdot 1)}} \end{split}$$

$$d = \sum_{i} (n_{i}-1)$$
$$c_{4} = \frac{4(n-1)}{(4n-3)}$$

The C_p index is process capability. It assumes the process mean is centered between the specification limits and essentially is the ratio of the distance between the specification limits to six process standard deviations. Obviously, the higher this value the better, because it means you can fit the process variation between the spec limits more easily. C_p measures the process' potential capability to meet the two-sided specifications. It does not take the process average into consideration.

High C_p indicates the small spread of the process with respect to the spread of the customer specifications. C_p is recommended when the process is centered between the specification limits. C_p works when there are both upper and lower specification limits. The higher C_p the better, meaning the spread of the process is smaller relative to the spread of the specifications.

Performance of the Process (P_p)

 P_p can be calculated with the formula:

$$P_{\rm p} = \frac{\rm USL-LSL}{6 \times \sigma_{\rm overall}}$$

Where:

$$\sigma_{overall} = \frac{s}{c_4(n)}$$

$$s = \sqrt{\sum_{i} \sum_{j} \frac{\left(x_{ij} - \bar{x}\right)^2}{n-1}}$$
$$c_4 = \frac{4(n-1)}{(4n-3)}$$

Like C_p , P_p assumes the process mean is centered between the spec limits and essentially is the ratio of the distance between the spec limits to six process standard deviations. Obviously, the higher this value the better, because it means you can fit the process variation between the spec limits more easily. Similar to C_p , P_p measures the capability of the process to meet the two-sided specifications. It only focuses on the spread and does not take the process centralization into consideration. It is recommended when the process is centered between the specification limits.

 C_p considers the within-subgroup standard deviation and P_p considers the total standard deviation from the sample data. P_p works when there are both upper and lower specification limits. P_p is a more conservative method for determining capability because it considers the total variation of a process instead of a subgroup of data.

Capability of the Process with a k Factor Adjustment (Cpk)

 $C_{\mbox{\scriptsize pk}}$ can be calculated with the formula:

$$C_{pk} = (1-k) \times C_p$$

Where:

$$k = \frac{|m-\mu|}{\frac{USL-LSL}{2}}$$
$$m = \frac{USL + LSL}{2}$$

The *m* is essentially the center point between the spec limits (it is calculated like an average). When calculating *k*, the numerator is the absolute value of the difference between the center point between the spec limits (*m*) and the process mean (μ); the denominator is half the distance between the specs. As you can guess from the calculation of *k*, it takes into consideration the location of the mean between the spec limits. The formulas to calculate C_{pk} can also be expressed as follows:

$$C_{pk} = \min(\frac{USL-\mu}{3\times\sigma_{within}}, \frac{\mu-LSL}{3\times\sigma_{within}})$$

Where:

$$\sigma_{\text{within}} = \frac{s_p}{c_4(d+1)}$$
$$s_p = \sqrt{\frac{\sum_i \sum_j (x_{ij} \cdot \bar{x}_i)}{\sum_i (n_i \cdot 1)}}$$
$$d = \sum_i (n_i \cdot 1)$$
$$c_4 = \frac{4(n \cdot 1)}{(4n \cdot 3)}$$

The expression for C_{pk} is indicating that the limiting factor is the specification limit that the mean is closest to. C_{pk} measures the process' actual capability by taking both the variation and average of the process into consideration. The process does not need to be centered between the specification limits to make the index meaningful. C_{pk} is recommended when the process is

not in the center between the specification limits. When there is only a one-sided limit, C_{pk} is calculated using C_{pu} or C_{pl} . C_{pk} for upper specification limit:

$$C_{pu} = \frac{USL-\mu}{3 \times \sigma_{within}}$$

 $C_{\mbox{\scriptsize pk}}$ for lower specification limit:

$$C_{\rm pl} = \frac{\mu - LSL}{3 \times \sigma_{\rm within}}$$

Where:

- *USL* and *LSL* are the upper and lower specification limits.
- μ is the process mean.

Performance of the Process with a k Factor Adjustment (P_{pk})

Ppk can be calculated with the formula:

$$P_{pk} = (1-k) \times P_p$$

Where:

$$k = \frac{|m-\mu|}{\frac{USL-LSL}{2}}$$
$$m = \frac{USL + LSL}{2}$$

Where:

- USL and LSL are the upper and lower specification limits
- μ is the process mean.

P_{pk} is to P_p as C_{pk} is to C_p. The *k* factor is the same used earlier to calculate C_{pk}.

The formulas to calculate P_{pk} can also be expressed as follows:

$$P_{pk} = \min(\frac{USL-\mu}{3\times\sigma_{overall}}, \frac{\mu-LSL}{3\times\sigma_{overall}}, \frac{\sigma_{overall}}{\sigma_{overall}}, \frac{\sigma_{overal$$

$$c_4 = \frac{4(n-1)}{(4n-3)}$$

Similar to C_{pk} , P_{pk} measures the process capability by taking both the variation and the average of the process into consideration. P_{pk} solves the decentralization problem P_p cannot overcome. C_{pk} considers the within-subgroup standard deviation, while P_{pk} considers the total standard deviation from the sample data. When there is only a one-sided specification limit, P_{pk} is calculated using P_{pu} or P_{pl} . P_{pk} for upper specification limit:

$$P_{pu} = \frac{USL-\mu}{3 \times \sigma_{overall}}$$

P_{pk} for lower specification limit:

$$P_{\rm pl} = \frac{\mu - \rm LSL}{3 \times \sigma_{\rm overall}}$$

Where:

- USL and LSL are the upper and lower specification limits
- μ is the process mean.

Taguchi's Capability Index (Cpm)

 C_p , P_p , C_{pk} , and P_{pk} all consider the variation of the process. C_{pk} and P_{pk} take both the variation and the average of the process into consideration when measuring the process capability. It is possible that the process average fails to meet the target customers require while the process still remains between the specification limits. C_{pm} (Taguchi's capability index) helps to capture the variation from the specified target. Rather than only telling us if a measurement is good (between the spec limits), C_{pm} helps us understand *how good* the measurement is. C_{pm} can be calculated with the formula:

$$C_{pm} = \frac{\min(T-LSL, USL-T)}{3 \times \sqrt{s^2 + (\mu - T)^2}}$$

Where:

- USL and LSL are the upper and lower specification limits
- *T* is the specified target
- μ is the process mean.

Note: C_{pm} can work only if there is a target value specified.

Capability Interpretation

Interpreting the results of a capability analysis is very dependent upon the nature of the business, product and customer. The Automotive Industry Action Group (AIAG) suggests that P_p and P_{pk} should be > 1.67 while C_p and C_{pk} should be > 1.33. The reality is that anything

greater than 1.0 is a fairly capable process but your business needs to assess the costs vs. benefits of achieving capability greater than 1.67 or even higher. If you're dealing with customer safety or life and death influences then obviously the product necessitates a capability greater than 2.0. Below is a simple table for quick interpretation with sigma level references.

Index	Value	Interpretation	Sigma Level
Cpk	<1.0	Not Very Capable	<3
Cpk	1.0-1.99	Capable	3-6
Cpk	>2.0	Very Capable	>6

Use Minitab to Run a Process Capability Analysis

J Data File: "Capability Analysis" tab in "Sample Data.xlsx"

Steps in Minitab to run a process capability analysis:

- 1. Click Stat \rightarrow Basic Statistics \rightarrow Normality Test.
- 2. A new window named "Normality Test" pops up.
- 3. Select "HtBk" as the variable.

Normality Test	X
	Variable: HtBk Percentile Lines
	None O At data values:
	Tests for Normality • Anderson-Darling
	C Ryan-Joiner (Similar to Shapiro-Wilk) C Kolmogorov-Smirnov
Select	Title:
Help	OK Cancel

Fig. 2.71 Normality Test dialog box with variable

4. Click "OK."

5. The histogram and the normality test results appear in the new window.

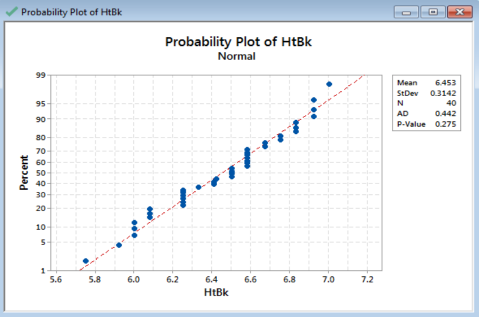


Fig. 2.72 Normality Test output

In this example, the p-value is 0.275, greater than the alpha level (0.05). We fail to reject the hypothesis and conclude that the data are normally distributed.

- 6. Click Stat \rightarrow Quality Tools \rightarrow Capability Analysis \rightarrow Normal.
- 7. A new window named "Capability Analysis (Normal Distribution)" pops up.
- 8. Select "HtBk" as the single column and enter "1" as the subgroup size.
- 9. Enter "6" as the "Lower spec" and "7" as the "Upper spec"

Capability Analysis (No	ormal Distribution)		×
	Data are arranged as G Single column: HtBk Subgroup size: 1 (use a constant or an ID col C Subgroups across rows of:	lumn)	Transform Estimate Options Storage
	Lower spec:	6	Boundary
	Upper spec:	7	Boundary
	Historical mean:		(optional)
Select	Historical standard deviation:		(optional)
Help			OK Cancel

Fig. 2.73 Capability Analysis (Normal Distribution) dialog box with variable

10. Click "Options" button and another new window named "Capability Analysis (Normal Distribution) – Options" pops up.

11. Enter "6.5" as the target and click "OK."

Capability Analysis (Normal Distribution)): Options
Target (adds Cpm to table): 6.5	
Use tolerance of $K\times\sigma$ for capability statist	ics K = 6
 ✓ Within subgroup analysis ✓ Overall analysis 	Display Parts per million Percents
	 Capability stats (Cp, Pp) C Benchmark Z's (σ level)
	TInclude confidence intervals
	Confidence level: 95.0
	Confidence intervals: Two-sided 🗨
Title:	
Help	OK Cancel

Fig. 2.74 Capability Analysis (Normal Distribution): Options dialog box with variable

- 12. Click "OK" in the "Capability Analysis (Normal Distribution)" window.
- 13. The capability analysis results app ear in the new window.

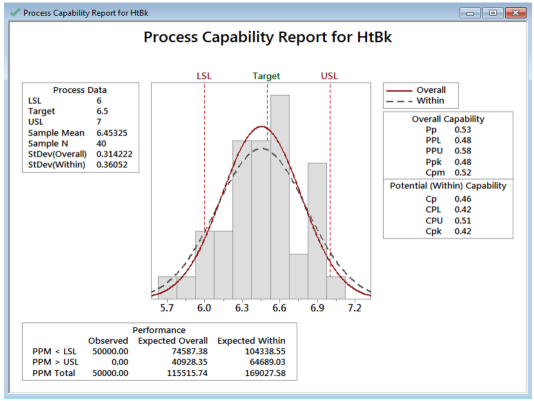


Fig. 2.75 Capability Analysis Output

If the p-value of the previous normality test is smaller than the alpha level (0.05), we would reject the null hypothesis and conclude that the data are not normally distributed. Thus, we would perform a Non-Normal Capability analysis as follows:

- 1. Click Stat \rightarrow Quality Tools \rightarrow Capability Analysis \rightarrow Non-Normal
- 2. A new window named "Capability Analysis (Non-Normal Distribution)" pops up.

- 3. Select "HtBk" as the single column.
- 4. Enter "6" as the "Lower spec" and "7" as the "Upper spec."

Capability Analysis (Nonnormal Distribution)				
	Data are arrange Single column Subgroups ac Fit distribution:	Estimate Options Storage		
Select	Lower spec: Upper spec:		oundary	
Help			OK Cance	4

Fig. 2.76 Capability Analysis (Non-Normal Distribution) dialog box with variable

- 5. Click "Options" button and another new window named "Capability Analysis (Non-Normal Distribution) – Options" pops up.
- 6. Enter "6.5" as the target and click "OK."

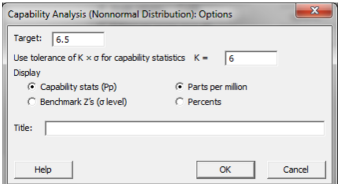


Fig. 2.77 Capability Analysis (Non-Normal Distribution): Options

7. Click "OK" in the "Capability Analysis (Non-Normal Distribution)" window.

8. The capability analysis results appear in the new window.

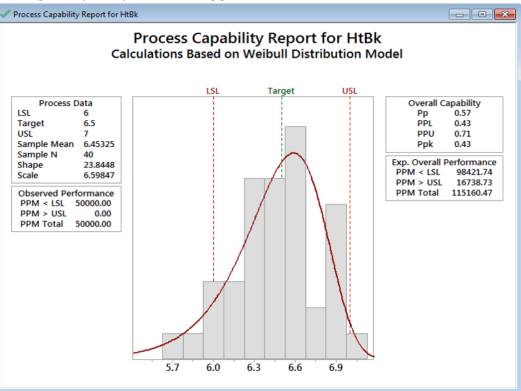


Fig. 2.78 Capability Analysis (Non-Normal Distribution) output

With P_{pk} of less than 1.0 we can conclude that the capability of this process is not very good. Anything less than 1.0 should be considered not capable and we should strive for P_{pk} to reach levels of greater than 1 and preferably over 1.67.

2.4.2 CONCEPT OF STABILITY

What is Process Stability?

A process is said to be stable when:

- The process is in control
- The future behavior of the process is predictable at least between some limits
- There is only random variation involved in the process.
- The causes of variation in the process are only due to chance or common causes
- There are not any trends, patterns, or outliers in the control chart of the process

Stability means that the process is in control; there are no special causes (only random variation) influencing the process outcome. Future process outcomes are predictable within a range of values, and there are no trends, patterns, or outliers in a control chart of the process.

Root Causes of Variation in the Process

To discuss stability, it is important that we define and differentiate common cause and special cause variation. Common causes are unable to be eliminated from the process. Typical common causes are:

- Chance
- Random and anticipated
- Natural noise
- Inherent in the process

Special causes are:

- Assignable cause
- Unanticipated
- Unnatural pattern
- The signal of changes in the process

Special causes are able to be eliminated from the process. Common cause variation is left to chance. It is random, it is expected, and inherent to the process; therefore, it is unable to be eliminated. Special cause, on the other hand, can be attributed to an "assignable cause." It is not expected (or anticipated) and is external to the process. These causes signal changes in the process and can be eliminated.

Control Charts

Control charts are the graphical tools to analyze the stability of a process. A control chart is used to identify the presence of potential special causes in the process and to determine whether the process is statistically in control. If the samples or calculations of samples are all in control, the process is stable and the data from the process can be used to predict the future performance of the process.

Popular Control Charts

- I-MR Chart—Individuals and moving range
- Xbar-R Chart—Xbar is the average and R is range
- Xbar-S Chart—Xbar is the average and S is standard deviation
- C Chart—Uses count data to chart non-conformances or defects)
- U Chart—U stands for units. It is the same as a C chart, but a C chart assumes subgroup side is constant, while a U chart assumes uneven subgroup sizes.
- P Chart—Proportions chart, which shows the proportion defective
- NP Chart—Shows number defective (with equal subgroup size)
- EWMA Chart—Exponentially weighted moving average
- CUSUM Chart—Cumulative sum

Process Stability vs. Process Capability

Process stability indicates how stable a process performed in the past. When the process is stable, we can use the data from the process to predict its future behavior. Process capability indicates how well a process performs with respect to meeting the customer's specifications. The process capability analysis is valid only if the process is statistically stable (i.e., in control, predictable). Being stable does *not* guarantee that the process is also capable. However, being stable is the prerequisite to determine whether a process is capable.

2.4.3 ATTRIBUTE AND DISCRETE CAPABILITY

Process Capability Analysis for Binomial Data

If we are measuring the count of defectives in each sample set to assess the process performance of meeting the customer specifications, we use "%Defective" (percentage of items in the samples that are defective) as the process capability index. %Defective is given by a very simple calculation: determine the number of defective units divided by the number of units overall.

% Defective =
$$\frac{N_{defectives}}{N_{overall}}$$

Where:

- $\bullet \quad N_{defectives} \, is \, the \, total \, count \, of \, defectives \, in \, the \, samples$
- Noverall is the sum of all the sample sizes.

Process Capability Analysis for Poisson Data

If we are measuring the count of defects in each sample set to assess the process performance of meeting the customer specifications, we use Mean DPU (defects per unit of measurement) as the process capability index:

$$\text{DPU} = \frac{\text{N}_{\text{defects}}}{\text{N}_{\text{overall}}}$$

Where:

- N_{defects} is the total count of defects in the samples
- Noverall is the sum of all the units in the samples.

2.4.4 MONITORING TECHNIQUES

Capability and Monitoring

In the Measure phase of the project, process stability analysis and process capability analysis are used to baseline the performance of current process. In the Control phase of the project,

process stability analysis and process capability analysis are combined to monitor whether the improved process is maintained consistently as expected.

3.0 ANALYZE PHASE

3.1 INFERENTIAL STATISTICS

3.1.1 UNDERSTANDING INFERENCE

What is Statistical Inference?

Statistical inference is the process of making inferences regarding the characteristics of an unobservable population based on the characteristics of an observed sample. Statistical inference is widely used since it is difficult or sometimes impossible to collect the entire population data. It is rare that we ever know the characteristics of a population, so we need to take limited data and infer what the population looks like.

Outcome of Statistical Inference

The outcome or conclusion of statistical inference is a *statistical proposition* about the population, such as an estimate of the population mean or standard deviation.

Examples of statistical propositions:

- Estimating a population parameter
- Identifying an interval or a region where the true population parameter would fall with some certainty
- Deciding whether to reject a hypothesis made on characteristics of the population of interest
- Making predictions
- Clustering or partitioning data into different groups

Population and Sample

A *statistical population* is an entire set of objects or observations about which statistical inferences are to be drawn based on its sample. It is usually impractical or impossible to obtain the data for the entire population. For example, if we are interested in analyzing the population of all the trees, it is extremely difficult to collect the data for all the trees that existed in the past, exist now, and will exist in the future.

A *sample* is a subset of the population (like a piece of the pie above). It is necessary for samples to be *representative* of the population. The process of selecting a subset of observations within a population is referred to as *sampling*.

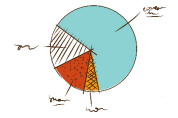


Fig. 3.18 Sampling a statistical population

When we talk about statistical inference, it is because we are trying to make inferences about a population based upon a sample size. Consider the presidential elections. The "population" includes every single person that casts a vote. However, on Election Day you will see the news outlets making projections (inferences) about how people have voted, based upon sampling being done throughout the day. Unless the population is very small, it is typically impractical or impossible to obtain measurements on an entire population.

Population and Sample

Population Parameters (Greek letters) Mean: μ Standard deviation: σ Variance: σ^2 Median: η Sample Statistics (Roman letters) Sample Mean: \overline{X} Standard deviation: sVariance: S^2 Median: \widetilde{X}

The *population parameter* is the numerical summary of a population. The *sample statistic* is the numerical measurement calculated based on a sample of that population. It is used to estimate the true population parameter. There is a different nomenclature and there are different symbols that we use when referring to a population or a sample.

Descriptive Statistics vs. Statistical Inference

Descriptive Statistics

Descriptive statistics summarize the characteristics of a collection of data. Descriptive statistics are descriptive only and they do not make any generalizations beyond the data at hand. Data used for descriptive statistics are for the purpose of representing or reporting.

Statistical Inference

Statistical inference makes generalizations from a sample at hand of a population. Data used for statistical inference are for the purpose of making inferences on the entire population of interest. A complete statistical analysis includes *both* descriptive statistics and statistical inference.

Error Sources of Statistical Inference

Statistical inference uses sample data to best approximate the true features of the population. A valid sample must be unbiased and representative of the population, meaning that the data should not be selected in a way that only represents a certain segment of the population. For example, what would happen if we wanted to understand income for the population of the US? In our sampling, if we only pulled data for professional athletes, or if we only pulled our sample in Beverly Hills, CA, what would happen to our data?

The two sources of error in statistical inference are:

- 1. Random Sampling Error
 - Is variation due to observations being selected randomly.
 - Is inherent to the sampling process and beyond one's control
- 2. Selection Bias
 - Non-random variation due to inadequate design of sampling
 - It can be improved by adjusting the sampling size and sampling strategy

Random sampling error is expected when observations are selected randomly. This error is expected when we are not able to look at an entire population.

Selection bias is when we do not consider the profile of the population in the design of our sampling strategy, meaning the design is inadequate to represent the population.

3.1.2 SAMPLING TECHNIQUES

What is Sampling?

Sampling is the process of selecting objects or observations from a population of interest about which we wish to make a statistical inference. It is extensively used to collect information about a population, which can represent physical or intangible objects.



Fig. 3.19 Sampling process

We know that we usually cannot get data for an entire population. When we want to understand the population, we need to make sure we are getting a good sample that will allow us to make a statistical inference.

Advantages of Sampling

It is usually impractical or impossible to collect the data of an entire population because it is costly and time consuming, but also because of the unavailability of historical records and the dynamic nature of the population.

Advantages of sampling a representative subset of the population:

- Lower cost
- Faster data collection
- Easier to manipulate

Uses of Samples

With valid samples collected, we can draw statistical conclusions about the population of our interest. There are two major uses of samples in making statistical inference:

- *Estimation*: Estimating the population parameters using the sample statistics
- *Hypothesis testing*: Testing a statement about the population characteristics using sample data

This module covers sample size calculation for estimation purposes only. Sample size calculations for hypothesis testing purposes will be covered in the Hypothesis Testing module.

Basic Sampling Steps

- 1. Determine the population of interest
- 2. Determine the sampling frame
- 3. Determine the sampling strategy
- 4. Calculate the sample size
- 5. Conduct sampling

Over the next few pages, we will go into some detail about these basic steps to follow when doing sampling.

- What population are we interested in knowing about?
- What are the "guardrails" we put around the sample?
- What is the sampling strategy?
- What is the size of the sample needed?
- Do the sampling.

Population

Population in statistics is the entire set of objects or observations about which we are interested to draw conclusions or generalize based on some representative sample data. The population covers all the items with characteristics we are interested to analyze.

A population can be either physical or intangible.

- Physical: trees, customers, monitors etc.
- Intangible: credit score, pass/fail decisions etc.

A population can be static or dynamic.

- Characteristics of individuals are relatively static over time.
- Items making up the population continue to change or be generated over time. For instance, the world population is constantly changing. On the other hand, the average stock price in 2011 is no longer changing.

Sampling Frame

In the ideal situation, the scope of a population could be defined. For example, Mary is interested to know how the soup she is cooking tastes. The population is simply the pot of soup. However, in some other situations, the population cannot be identified or defined precisely. For example, to collect the information for an opinion poll, we do not have a list of all the people in the world at hand.

A *sampling frame* is a list of items of the population (preferably the entire population; if not, approximate population). For example, the telephone directory would be a sampling frame for opinion poll data collection.

Basic Sampling Strategies

- Simple Random Sampling
- Matched Random Sampling
- Stratified Sampling

- Systematic Sampling
- Proportional to Size Sampling
- Cluster Sampling

Simple Random Sampling

Simple random samples are selected in such a way that each item in the population has an equal chance of being selected. There is no bias involved in the sample selection. Such selection minimizes the variation between the characteristics of samples and the population. It is the basic sampling strategy.

Matched Random Sampling

Matched random samples are samples randomly selected in pairs, each of which has the same attribute. For example, researchers are interested in understanding the weight of twins; researchers are interested in understanding the patients' blood pressure before and after taking some medicine.

Stratified Sampling

A population can be grouped or "stratified" into distinct and independent categories. An individual category can be considered as a sub-population. *Stratified samples* are randomly

selected in each category of the population. The categories can be gender, region, income level etc.

Stratified sampling requires advanced knowledge of the population characteristics, for example: A fruit store wants to measure the quality of all their oranges. They decide to use stratified sampling by region to collect sample data. Since about 40% of their oranges are from California, 40% of the sample is selected from the California oranges sub-population.

Systematic Sampling

Systematic samples are selected at regular intervals based on an ordered list where items in the population are arranged according to a certain criterion. Systematic sampling has a random start and then every *i*th item is selected going forward, for example, we are sampling the every 5th unit produced on the production line.

Cluster Sampling

Cluster sampling is a sampling method in which samples are only selected from certain clusters or groups of the population. It reduces the cost and time spent on the sampling but bears the risk that the selected clusters are biased, for example, selecting samples from the region where researchers are located so that the cost and time spent on travelling is reduced.

Sampling Strategy Decision Factors

When determining the sampling strategy, we need to consider the following factors:

- Cost and time constraints
- Nature of the population of interest
- Availability of advanced knowledge of the population
- Accuracy requirement

So how does one decide what sampling strategy to use? Consider how much time is available to collect the data and what cost restrictions are in place. Consider the nature of the population, and what you know about the population. Are there specific segments you are interested in? Are the samples streaming like in a manufacturing operation? Determine how accurate you need to be—this can compete with how much you must spend on sampling, but usually this will determine how big of a sample you should collect.

Sample Size

The *sample size* is a critical element that can influence the results of statistical inference. The smaller the sample size, the higher the risk that the sample statistic will not reflect the true population parameter. The greater the sample size, the more time and money we will spend on collecting the samples.

Sample Size Factors

There are a few factors to consider when determining the right sample size:

- Is the variable of interest continuous or discrete?
- How large is the population size?
- How much risk do you want to take regarding missing the true population parameters?
- What is the acceptable margin of error you want to detect?
- How much is the variation in the population? The larger the variation in the population, the larger the risk in sample variation.

Sample Size Calculation for Continuous Data

Sample size equation for continuous data

$$n_0 = \left(\frac{Z_{\frac{\alpha}{2}} \times s}{d}\right)^2$$

Where:

- *n* is the sample size we are trying to determine
- *Z* is a value that we look up based upon the confidence level. If we want 95% confidence, then alpha is 1 0.95 = 0.05 (or 5%). Therefore, we will look up the *Z* value for when alpha is 0.05, which is 1.96.
- *s* is the estimated standard deviation for the population
- *d* is the acceptable margin of error.

Sample Size Calculation for Continuous Data When the sample size calculated using the formula

$$n_0 = \left(\frac{Z_{\frac{\alpha}{2}} \times s}{d}\right)^2$$

exceeds 5% of the population size, we use this formula to calculate the sample size:

$$n = \frac{n_0}{\left(1 + \frac{n_0}{N}\right)}$$

Where:

 n_0 represents the sample size calculated using the formula on the previous page

N is the size of the population.

Sample Size Calculation for Discrete Data

Sample size equation for continuous data

$$n_0 = \left(\frac{Z_{\alpha/2}}{d}\right)^2 \times p \times (1-p)$$

Where:

- *n*⁰ is the number of samples
- $Z_{\alpha/2}$ is the *Z* score when risk level is $\alpha/2$. When α is 0.05, it is 1.96. When α is 0.10, it is 1.65.
- *s* is the estimation of standard deviation in the population
- *d* is the acceptable margin of error
- *p* is the proportion of one type of event occurring (e.g., proportion of passes)
- $p \times (1 p)$ is the estimate of variance.

Sample Size Calculation for Discrete Data

Sample size calculated using the formula:

$$n_0 = \left(\frac{Z_{\alpha/2}}{d}\right)^2 \times p \times (1-p)$$

When the sample size exceeds 5% of the population size, we use a correction formula to calculate the final sample size.

$$n = \frac{n_0}{\left(1 + \frac{n_0}{N}\right)}$$

Where:

- *n*₀ is the sample size calculated using the previous equation
- *N* is the population size.

Sampling Errors

Random Sampling Error

- Is random variation due to observations being selected randomly
- Is inherent in the sampling process and beyond one's control

Selection Bias

- Is non-random variation due to inadequate design of sampling
- Can be improved by adjusting the sampling size and sampling strategy

3.1.3 SAMPLE SIZE

Sample Size

In the last model, we talked about sample size considerations for when we are looking to estimate parameters for a population. Now we want to use sample size considerations for hypothesis tests. The sample size is a critical element that can influence the results of hypothesis testing. The smaller the sample size, the higher the risk that the statistical conclusions will not reflect the population relationship. The greater the sample size, the more time and money we will spend on collecting the samples. So, we must figure out where the right balance is.

Sample Size Calculation

General sample size formula for *continuous data*:

$$n = \left(\frac{\left(Z_{\alpha/2} + Z_{\beta}\right) \times s}{d}\right)^{2}$$

General sample size formula for *discrete data*:

$$n = \left(\frac{Z_{\frac{\alpha}{2}} + Z_{\beta}}{d}\right)^2 \times p \times (1-p)$$

Where:

- *n* is the number of observations in the sample
- *α* is the risk of committing a false positive error (or Type I error). A false positive error, commonly called a false alarm, is a result that indicates a given condition has been fulfilled, when it has not been fulfilled. In the case of "crying wolf," the condition tested for was "Is there a wolf near the herd?" The actual result was that there had not been a wolf near the herd. The shepherd wrongly indicated there was one, by calling "Wolf, wolf!"
- β is the risk of committing a false negative error (or Type II error). A false negative error is where a test result indicates that a condition failed, while it was successful. An example is a guilty prisoner freed from jail. The condition, "Is the prisoner guilty?" had a positive result (yes, he is guilty). But the test failed to realize this, and wrongly decided the prisoner was not guilty.

- *s* is the estimation of standard deviation in the population
- *d* is the size of effect you want to be able to detect
- *p* is the proportion of one type of event occurring (e.g., proportion of passes)

Use Minitab to Calculate the Sample Size

Case study: We are interested in comparing the average retail price of a product between two states. We will run a hypothesis test on the two sample means to determine whether there is a statistically significant difference between the retail prices in the two states. The average retail price of the product is \$23 based on our estimation and the standard deviation is 3. We want to detect at least a \$2 difference with a 90% chance when it is true and we can tolerate the alpha risk at 5%. What should the sample size be? To use the software to calculate the sample size, consider this example where we are trying to determine if there is a statistical difference between two sample means.

Use Minitab to Calculate the Sample Size

Steps to calculate the sample size in Minitab:

- 1. Click Stat \rightarrow Power and Sample Size \rightarrow 2-Sample t.
- 2. A window named "Power and Sample Size for 2-Sample t" appears.
- 3. Enter "2" as "Differences."
- 4. Enter "0.9" as "Power values."
- 5. Enter "3" as "Standard deviation."

Power and Sample Size for 2-Sample t		
Specify values for a Sample sizes:	ny two of the following:	_
Differences:	2	
Power values:	0.9	
Standard deviation:	3	
	Options Graph	
Help	OK Cancel	

Fig. 3.20 Power and Sample Size for 2-Sample t with variables

6. Click "OK."

7. The sample size calculation results appear in a new window.

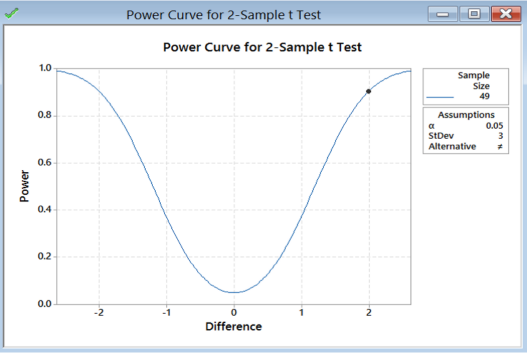


Fig. 3.21 Power and Sample Size for 2-Sample t output

The sample size for each group is 49 based on the sample size calculator. When the absolute value of the difference to detect decreases, the power decreases accordingly if the sample size is kept the same. 0.05 is the significance level by default. To customize the significance level, click on the "Options" button in the window "Power and Sample Size for 2-Sample t," specify the significance value in the box, and click "OK."

Power and Sample Size for 2-Sample t:	×
Alternative Hypothesis C Less than Not equal C Greater than	
Significance level: 0.05	
Help OK Cancel	

Fig. 3.22 Power and Sample Size for 2-Sample t Options

3.1.4 Central Limit Theorem

What is Central Limit Theorem?

The *Central Limit Theorem* is one of the fundamental theorems of probability theory. It states a condition under which the mean of a large number of independent and identically-distributed random variables, each of which has a finite mean and variance, would be approximately normally distributed. Let us assume $Y_1, Y_2...Y_n$ is a sequence of n i.i.d. random variables, each of which has finite mean μ and variance σ^2 , where $\sigma^2 > 0$. When n increases, the sample average of the n random variables is approximately normally distributed, with the mean equal to μ and variance equal to σ^2/n , regardless of the common distribution Y_i follows where i = 1, 2...n.

Independent and Identically Distributed

A sequence of random variables is *independent and identically distributed* (i.i.d.) if each random variable is independent of others and has the same probability distribution as others. It is one of the basic assumptions in Central Limit Theorem. Consider the law of large numbers (LLN)— It is a theorem that describes the result of performing the same experiment a large number of times. According to the LLN, the average of the results obtained from a large number of trials should be close to the expected value, and will tend to become closer as more trials are performed. The following example will explain this further.

Central Limit Theorem Example

Let us assume we have 10 fair die at hand. Each time we roll all 10 die together we record the average of the 10 die. We repeat rolling the die 50 times until we will have 50 data points. Upon doing so, we will discover that the probability distribution of the sample average approximates the normal distribution even though a single roll of a fair die follows a discrete uniform distribution. Knowing that each die has six possible values (1, 2, 3, 4, 5, 6), when we record the average of the 10 dice over time, we would expect the number to start approximating 3.5 (the average of all possible values). The more rolls we perform, the closer the distribution would be to a normal distribution centered on a mean of 3.5.

Central Limit Theorem Explained in Formulas

Let us assume $Y_1, Y_2 \dots Y_n$ are i.i.d. random variables with

$$E(Y_i) = \mu_Y \quad \text{where-} \infty < \mu_Y < \infty$$
$$var(Y_i) = \sigma_Y^2 \quad \text{where } 0 < \sigma_Y^2 < \infty$$

As $n \to \infty$, the distribution of \overline{Y} becomes approximately normally distributed with

$$E\left(\overline{Y}\right) = \mu_{Y}$$
$$\operatorname{var}(\overline{Y}) = \sigma_{\overline{Y}}^{2} = \frac{\sigma_{Y}^{2}}{n}$$

. .

Where:

- Y_{1} , Y_{2} , Y_{n} are independent and identically distributed random variables
- *E*(*Y_i*) = the expected value for Y is the mean for Y
- *Var*(*Y_i*) = the expected variance for Y.

As the *n* gets infinitely large, the distribution of Y means becomes approximately a normal distribution.

Central Limit Theorem Application

Use the sample mean to estimate the population mean if the assumptions of Central Limit Theorem are met:

$$E(Y_i) = E(\overline{y})$$

Where: *i* = 1, 2 . . . *n*.

Use standard error of the mean to measure the standard deviation of the sample mean estimate of a population mean.

$$SE_{\overline{Y}} = \frac{S}{\sqrt{n}}$$

Where: *s* is the standard deviation of the sample and *n* is the sample size.

Standard deviation of the population mean:

$$SD_{\overline{Y}} = \frac{\sigma}{\sqrt{n}}$$

Where:

- σ is the standard deviation of the population
- *n* is the sample size.

Use a larger sample size, if economically feasible, to decrease the variance of the sampling distribution. The larger the sample size, the more precise the estimation of the population parameter. Use a confidence interval to describe the region which the population parameter would fall in. The sample distribution approximates the normal distribution in which 95% of the data stays within two standard deviations from the center. Population mean would fall in the interval of two standard errors of the mean away from the sample mean, 95% of the time.

Confidence Interval

The *confidence interval* is an interval where the true population parameter would fall within a certain confidence level. A 95% confidence interval, the most commonly used confidence level, indicates that the population parameter would fall in that region 95% of the time or we are

95% confident that the population parameter would fall in that region. The confidence interval is used to describe the reliability of a statistical estimate of a population parameter.

The width of a confidence interval depends on the:

- Confidence level—The higher the confidence level, the wider the confidence interval
- Sample size—The smaller the sample size, the wider the confidence interval
- Variability in the data—The more variability, the wider the confidence interval

Confidence Interval of the Mean

Confidence interval of the population mean μ_Y of a continuous variable Y is:

$$\left[\overline{Y} - Z_{\alpha_{2}'} \times \left(\frac{\sigma}{\sqrt{n}}\right), \overline{Y} + Z_{\alpha_{2}'} \times \left(\frac{\sigma}{\sqrt{n}}\right)\right]$$

Where:

- \overline{Y} is the sample mean
- σ is the standard deviation of the population
- *n* is the sample size
- $Z_{\alpha/2}$ is the Z score when risk level is $\alpha/2$.

When α is 0.05, it is 1.96. When α is 0.10, it is 1.65.

α is (1 – confidence level). When confidence level is 95%, *α* is 5%. When the confidence level is 90%, *α* is 10%.

Minitab: Calculate the Confidence Interval of the Mean

Data File: "Central Limit Theorem" tab in "Sample Data.xlsx"

- 1. Click Stat \rightarrow Basic Statistics \rightarrow Graphical Summary.
- 2. A new window named "Graphical Summary" pops up.
- 3. Select "Cycle Time (Minutes)" as the variable.

4. The confidence level is 0.95 by default.

Graphical Summary	×
C1 Cycle Time (Minu	Variables:
	'Cycle Time (Minutes)'
	By variables (optional):
	Confidence level: 95.0
Select	
Help	OK Cancel

Fig. 3.23 Graphical Summary dialog box with variable

- 5. Click "OK."
- 6. A new window named "Summary for Cycle Time (Minutes)" pops up.

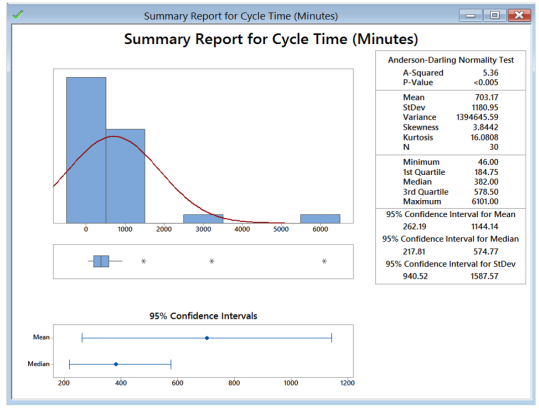


Fig. 3.24 Graphical Summary output

The 95% confidence interval of the mean is shown in the newly-generated "Summary for Cycle Time (Minutes)." The confidence level is 95% by default. In order to see the confidence interval

of "Cycle Time (Minutes)" at other confidence level, we need to enter the confidence level of our interest in the window "Graphical Summary" and click "OK."

Graphical Summary	×
	Variables: 'Cycle Time (Minutes)'
	By variables (optional):
	Confidence level: 90.0
Select	
Help	OK Cancel

Fig. 3.25 Graphical Summary dialog box

The following example shows how to generate 90% confidence interval of the mean.

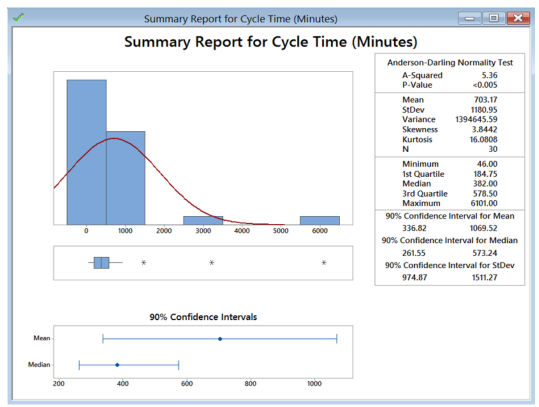


Fig. 3.26 Graphical Summary output with new confidence interval

Here the example shows what selections to make in Minitab to get the confidence intervals around the mean and standard deviation. While Minitab has the default confidence interval at 95%, you can see that other confidence levels can be selected as well.

3.2 Hypothesis Testing

3.2.1 GOALS OF HYPOTHESIS TESTING

What is Hypothesis Testing?

A *hypothesis test* is a statistical method in which a specific hypothesis is formulated about a population, and the decision of whether to reject the hypothesis is made based on sample data. Hypothesis tests help to determine whether a hypothesis about a population or multiple populations is true with certain confidence level based on sample data. Hypothesis testing is a critical tool in the Six Sigma tool belt. It helps us separate fact from fiction, and special cause from noise, when we are looking to make decisions based on data.

Hypothesis Testing Examples

Hypothesis testing tries to answer whether there is a difference between different groups or there is some change occurring.

- Are the average SAT scores of graduates from high school A and B the same?
- Is the error rate of one group of operators higher than that of another group?
- Are there any non-random causes influencing the height of kids in one state?

What is Statistical Hypothesis?

A *statistical hypothesis* is an assumption about one or multiple population. It is a statement about whether there is any difference between different groups. It can be a conjecture about the population parameters or the nature of the population distributions.

A statistical hypothesis is formulated in pairs:

- Null Hypothesis
- Alternative Hypothesis

Statements like the following reflect a null hypothesis:

- The data are normally distributed.
- The population mean is the same as X.
- Group A and Group B are the same.

And the alternative hypotheses are:

- The data are not normally distributed
- The population mean is not the same as X.
- Group A and Group B are not the same.

Null and Alternative Hypotheses

Null Hypothesis (H₀) states that:

- There is no difference in the measurement of different groups
- No changes occurred
- Sample observations result from random chance

Alternative Hypothesis (H₁ or H_a) states that:

- There is a difference in the measurement of different groups
- Some changes occurred
- Sample observations are affected by non-random causes

A statistical hypothesis can be expressed in mathematical language by using population parameters (Greek letters) and mathematical symbols.

- Population Parameters (Greek letters)
- Mean: μ
- Standard deviation: σ
- Variance: σ^2
- Median: η
- Mathematical Symbols
- Equal: =
- Not equal: ≠
- Greater than: >
- Smaller than: <

The following are examples of null and alternative hypotheses written in mathematical language.

$$\begin{cases} H_0; \mu_1 = \mu_2 \\ H_1; \mu_1 \neq \mu_2 \end{cases} \\ \begin{cases} H_0; \sigma_1 = 0 \\ H_1; \sigma_1 \neq 0 \end{cases} \\ \begin{cases} H_0; \eta_1 = 10 \\ H_1; \eta_1 > 10 \end{cases} \\ \begin{cases} H_0; \mu_1 = 10 \\ H_1; \mu_1 < 10 \end{cases}$$

The first pair reads: Null hypothesis where population mean 1 is equal to population mean 2. Alternative hypothesis is population mean 1 is NOT equal to population mean 2. Or in the third pair: Null hypothesis where population mean 1 = 10. Alternative hypothesis is population mean greater than 10.

Hypothesis Testing Conclusion

There are two possible conclusions of hypothesis testing:

- 1. Reject the null
- 2. Fail to reject the null

When there is enough evidence based on the sample information to prove the alternative hypothesis, we reject the null. When there is *not* enough evidence or the sample information is *not* sufficiently persuasive, we fail to reject the null.

Decision Rules in Hypothesis Testing

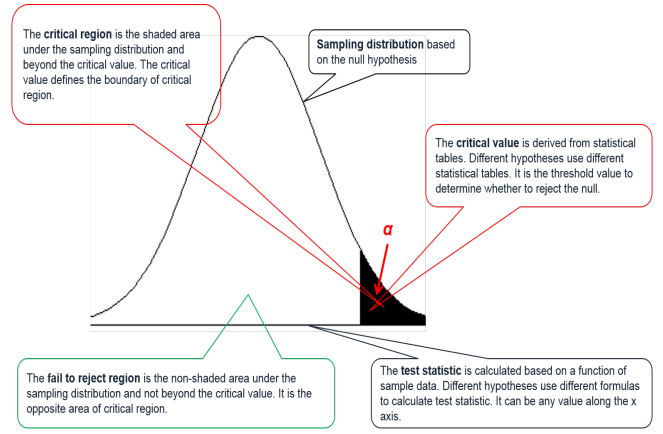


Fig. 3.27 Rules in Hypothesis Testing

As we get into the technicalities of hypothesis testing, it is important to understand some terms as they pertain to a distribution. Consider in this picture a normal distribution, and what we know about the distribution. Remember 68-95-99.7? The amount of data that falls within +/-1, 2, and 3 standard deviations of the mean.

The *test statistic* in hypothesis testing is a value calculated using a function of the sample. Test statistics are considered the sample data's numerical summary that can be used in hypothesis testing. Different hypothesis tests have different formulas to calculate the test statistic.

The *critical value* in hypothesis testing is a threshold value to which the test statistic is compared in order to determine whether the null hypothesis is rejected. The critical value is obtained from statistical tables. Different hypothesis tests need different statistical tables for critical values.

Hypothesis testing is made easy with our fancy software packages, but it is important that you understand how the theory and rules work behind the software.

- When the test statistic falls into the fail to reject region, we fail to reject the null and claim that there is no statistically significant difference between the groups.
- When the test statistic falls into the critical region, we reject the null and claim that there is a statistically significant difference between the groups.

The proportion of the area under the sampling distribution and beyond the critical value indicates α risk (also called α *level*). The most commonly selected α level is 5%. To illustrate, let us assume the test statistic is equal the critical value of 5%. In this case, there is a 5% risk that we will reject the null hypothesis when it is true. So, as the test statistic falls in the critical region, the risk of rejecting the null when it is actually true is less than 5%. Another way to state it is that we are 95% confident that we should reject the null hypothesis.

The proportion of the area under the sampling distribution and beyond the test statistic is the *p-value*. It is the probability of getting a test statistic at least as extreme as the observed one, given the null is true.

- When the p-value is smaller than the α level, we reject the null and claim that there is a statistically significant difference between different groups.
- When the p-value is higher than the α level, we fail to reject the null and claim that there is no statistically significant difference between different groups.

Steps in Hypothesis Testing

- Step 1: State the null and alternative hypothesis.
- Step 2: Determine α level.
- Step 3: Collect sample data.
- Step 4: Select a proper hypothesis test.
- Step 5: Run the hypothesis test.
- Step 6: Determine whether to reject the null.

3.2.2 Statistical Significance

Statistical Significance

In statistics, an observed difference is *statistically significant* if it is unlikely that the difference occurred by pure chance, given a predetermined probability threshold. Statistical significance indicates that there are some non-random factors causing the result to take place. The statistical significance level in hypothesis testing indicates the amount of evidence which is sufficiently persuasive to prove that a difference between groups exists not due to random chance alone.

When we talk about determining if something is statistically significant, we will use the p-value. The p-value is the probability that the difference between two groups is driven by chance. The lower the p-value, the more likely that there is some non-random factor that is causing the difference.

Practical Significance

An observed difference is *practically significant* when it is large enough to make a practical difference. A difference between groups that is *statistically significant* might not be large enough to be practically significant. Sometimes we determine with statistics that there are differences. However, while those differences might be statistically significant, they may not be large enough to make a practical difference from a business standpoint. When analysis determines that a difference is statistically significant, ask yourself the question, "Why does it matter?" or "Why should we care?"

Example: You started to use the premium gas recently, which was supposed to make your car run better. After running a controlled experiment to measure the performance of the car before and after using the premium gas, you performed a statistical hypothesis test and found that the difference before and after was statistically significant. Using premium gas did improve the performance. However, due to the high cost of the premium gas, you decided that the difference was not large enough to make you pay extra money for it. In other words, the difference is not practically significant.

In this example, the premium gas did make a difference in performance. However, you must ask what is important. If cost is important, does the increased cost of premium gas have long term savings benefit—better gas mileage, lower maintenance, and repair cost, longer engine life?

3.2.3 RISK; ALPHA AND BETA

Errors in Hypothesis Testing

In statistical hypothesis testing, there are two types of errors:

- *Type I Error*—A null hypothesis is rejected when it is true in fact
- *Type II Error*—A null hypothesis is not rejected when it is not true in fact

	Null hypothesis is true	Alternative hypothesis is true
Fail to reject null hypothesis	Correct	Incorrect (Type II Error)
Reject null hypothesis	Incorrect (Type I Error)	Correct

Fig. 3.28 Types of Errors in Hypothesis Testing

Type I error is when we react as though there is a change or a difference, when in fact it is only chance that is driving the difference.

Type II error is when we do not react when we should, meaning there is a difference but we act as though there is not.

Type I Error

Type I error is also called false positive, false alarm, or alpha (α) error. Type I error is associated with the risk of accepting false positives. It occurs when we think there is a difference between groups but in fact there is none, for example, telling a patient he is sick and in fact he is not. Type I error leads to "tampering," meaning we make changes or adjustments when we should not. We create a moving target and make it even more difficult to control the process.

Alpha (α)

Alpha (α) indicates the probability of making a type I error. It ranges from 0 to 1. α risk is the risk of making a type I error.

The most commonly used α *is 5%.* It corresponds to the fact that most commonly we use a 95% confidence level. The confidence level used to calculate the confidence intervals is ($100\% - \alpha$).

When making a decision on whether to reject the null, we compare the p-value against α :

- If p-value is smaller than *α*, we reject the null
- If p-value is greater than α , we fail to reject the null

To reduce the α risk, we decrease the α value to which the p-value is compared. When we make a decision about statistical significance, we compare the p-value against the alpha value. So, if our confidence level is 95%, our alpha is 5%; therefore, our p-value is 0.05. If p-value is less than alpha, then we reject the null hypothesis. To reduce risk, or increase confidence, we lower the alpha value that the p-value is compared to. To be 99% confident, our alpha would be (1 - 0.99) = 0.01.

Type II Error

Type II error is also called false negative, oversight, or beta (β) error. Type II error is associated with the risk of accepting false negatives. It occurs when we think there is not any difference between groups but in fact there is, for example, telling a patient he is not sick and in fact he is. Type II error describes when we are unresponsive to situations that we should respond to.

Beta (β)

 β indicates the probability of making a type II error. It ranges from 0 to 1. β risk is the risk of making a type II error. The most commonly used β is *10%*. It corresponds to the fact that most commonly we use a 90% power. The expression (*100%* – β) is called *power*, which denotes the probability of detecting a difference between groups when in fact the difference truly exists. To reduce the β risk, we increase the sample size. When holding other factors constant, β is inversely related to α .

3.2.4 Types of Hypothesis Tests

Two Types of Hypothesis Tests

When we talk about these two type of hypothesis tests, two-tailed and one-tailed, they are a direct reference to the tails of a data distribution.

Two-tailed hypothesis test

- Is also called two-sided hypothesis test
- Is a statistical hypothesis test in which the critical region is split into two equal areas, each of which stays on one side of the sampling distribution

One-tailed hypothesis test

- It is also called one-sided hypothesis test
- It is a statistical hypothesis test in which the critical region is only on one side of the sampling distribution

Two-Tailed Hypothesis Test

A two-tailed hypothesis test is used when we care about whether there is a difference between groups and we do not care about the direction of the difference.

Examples of two-tailed hypothesis tests:

$$\begin{cases} H_0: \mu_1 = 10 \\ H_a: \mu_1 \neq 10 \end{cases}$$

The null hypothesis (H₀) is rejected when:

- The test statistic falls into either half of the critical region
- The test statistic is either sufficiently small or sufficiently large
- The absolute value of the test statistic is greater than the absolute value of the critical value

A two-tailed hypothesis test is the most commonly used hypothesis test. In the next modules, we will cover more details about it. When we are comparing group A and group B, and we only

care to know if they are different, vs. one specific group being greater than another, we use a two-tailed test.

Two-Tailed Hypothesis Test

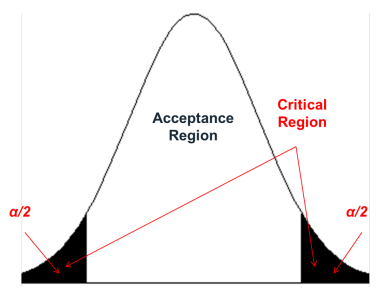


Fig. 3.29 Two-Tailed Hypothesis Test

As described on the previous page, the critical regions are on both (two) tails of the sample distribution.

One-Tailed Hypothesis Test

A one-tailed hypothesis test is used when we care about one direction of the difference between groups. Examples of one-tailed hypothesis tests:

$$\begin{cases} H_0 {:}\, \mu_1 = 10 \\ H_a {:}\, \mu_1 > 10 \end{cases}$$

The null hypothesis is rejected when the test statistic:

- Falls into the critical region which only exits on the right side of the distribution
- Is sufficiently large
- Is greater than the critical value

We use a one-tailed hypothesis test when we only care about one direction when comparing two groups, for instance, a null hypothesis where a population mean is equal to 10. An alternative hypothesis is where the population mean is not just unequal to 10, but greater than 10.

Of course, we look for the test statistic to fall into the critical region to reject the hypothesis. But in this situation, we are only looking for the test statistic to be sufficiently large—greater than the critical value.

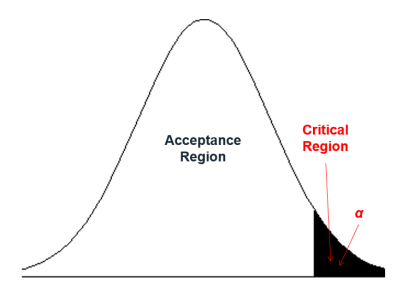


Fig. 3.30 One-Tailed Hypothesis Test

The critical region is only on one side of the distribution. The following are examples of onetailed hypothesis test.

The null hypothesis is rejected when the test statistic:

- Falls into the critical region which only exits on the left side of the distribution
- Is sufficiently small
- Is smaller than the critical value

This is the other side of the one-tailed hypothesis test, where the alternative hypothesis is that the population mean is less than 10.

We look for the test statistic to fall into the critical region to reject the hypothesis. But in this situation, we are only looking for the test statistic to be sufficiently small—less than the critical value.

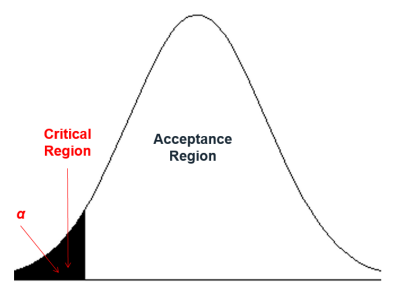


Fig. 3.31 One-Tailed Hypothesis Test

The critical region is on the other side of the distribution.

3.3 Hypothesis Tests: Normal Data

3.3.1 One and Two Sample T-Tests

What is a T-Test?

In statistics, a *t*-*test* is a hypothesis test in which the test statistic follows a *Student's t* distribution if the null hypothesis is true. We apply a t-test when the population variance (σ) is unknown and we use the sample standard deviation (*s*) instead.

The *t-statistic* was introduced in 1908 by William Sealy Gosset, a chemist working for the Guinness brewery in Dublin, Ireland ("Student" was his pen name). Gosset had been hired due to Claude Guinness's policy of recruiting the best graduates from Oxford and Cambridge to apply biochemistry and statistics to Guinness's industrial processes. Gosset devised the t-test as a cheap way to monitor the quality of stout. He published the test in Biometrika in 1908, but was forced to use a pen name by his employer, who regarded the fact that they were using statistics as a trade secret. In fact, Gosset's identity was known to fellow statisticians (http://en.wikipedia.org/wiki/Student's_t-test).

What is One Sample T-Test?

One sample t-test is a hypothesis test to study whether there is a statistically significant difference between a population mean and a specified value.

- Null Hypothesis (H₀): μ = μ₀
- Alternative Hypothesis (H_a): $\mu \neq \mu_0$

Where:

- μ is the mean of a population of our interest
- μ_0 is the specific value we want to compare against.

Assumptions of One Sample T-Test

- The sample data of the population of interest are unbiased and representative.
- The data of the population are continuous.
- The data of the population are normally distributed.
- The variance of the population of our interest is unknown.
- One sample t-test is more robust than the z-test when the sample size is small (< 30).

Normality Test

To check whether the population of our interest is normally distributed, we need to run normality test. While there are many normality tests available, such as Anderson–Darling, Sharpiro–Wilk, and Jarque–Bera, our examples will default to using the Anderson-Darling test for normality.

- Null Hypothesis (H₀): The data are normally distributed.
- Alternative Hypothesis (H_a): The data are not normally distributed.

Test Statistic and Critical Value of One Sample T-Test

To understand what is happening when you run a t-test with your software, the formulas here will walk you through the key calculations and how to determine if the null hypothesis should be accepted or rejected. To determine significance, you must calculate the t-statistic and compare it to the critical value, which is a reference value based on the alpha value and degrees of freedom (n - 1). The t-statistic is calculated based on the sample mean, the sample standard deviation, and the sample size.

Test statistic is calculated with the formula:

$$t_{calc} = \frac{\overline{Y}}{\frac{S}{\sqrt{n}}}$$

Where:

 \overline{Y} is the sample mean, *n* is the sample size, and *s* is the sample standard deviation

$$s = \sqrt{\frac{\sum_{i=1}^{n} (Y_i - \overline{Y})^2}{n-1}}$$

Critical value

- t_{crit} is the t-value in a Student's t distribution with the predetermined significance level α and degrees of freedom (n –1).
- t_{crit} values for a two-sided and a one-sided hypothesis test with the same significance level α and degrees of freedom (n 1) are different.

Decision Rules of One Sample T-Test

Based on the sample data, we calculated the test statistic t_{calc} , which is compared against t_{crit} to make a decision of whether to reject the null.

- Null Hypothesis (H₀): $\mu = \mu_0$
- Alternative Hypothesis (H_a): $\mu \neq \mu_0$
- If $|t_{calc}| > t_{crit}$, we reject the null and claim there is a statistically significant difference between the population mean μ and the specified value μ_0 .
- If $|t_{calc}| < t_{crit}$, we fail to reject the null and claim there is not any statistically significant difference between the population mean μ and the specified value μ_0 .

Use Minitab to Run a One-Sample T-Test

Case study: We want to compare the average height of basketball players against 7 feet.

Data File: "One Sample T-Test" tab in "Sample Data.xlsx"

- Null Hypothesis (H₀): *μ* = 7
- Alternative Hypothesis (H_a): $\mu \neq 7$

Step 1: Test whether the data are normally distributed

- 1. Click Stat \rightarrow Basic Statistics \rightarrow Normality Test.
- 2. A new window named "Normality Test" pops up.

3. Select "HtBk" as the variable.

Normality Test	
	Variable: HtBk
	Percentile Lines None At Y values:
	Tests for Normality Anderson-Darling Ryan-Joiner (Similar to Shapiro-Wilk) Kolmogorov-Smirnov
Select	Title:
Help	OK Cancel

Fig. 3.32 Normality Test dialog box with variable

- 4. Click "OK."
- 5. A new window named "Probability Plot of HtBk" appears, which covers the results of the normality test.

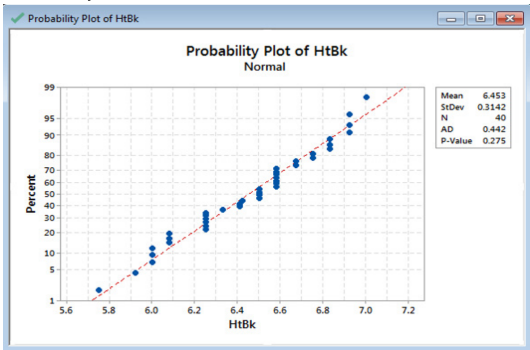


Fig. 3.33 Normality Test output

- Null Hypothesis (H₀): The data are normally distributed.
- Alternative Hypothesis (H_a): The data are not normally distributed.

Since the p-value of the normality is 0.275, which is greater than alpha level (0.05), we fail to reject the null and claim that the data are normally distributed. If the data are not normally distributed, you need to use hypothesis tests other than the one sample t-test. Now we can run the one-sample t-test, knowing the data are normally distributed.

Step 2: Run the one-sample t-test

- 1. Click Stat \rightarrow Basic Statistics \rightarrow 1 Sample t.
- 2. A new window named "One Sample t for the Mean" pops up.
- 3. Click the blank drop-down box and select "One or more samples, each in a column".
- 4. Select "HtBk" as the "Samples in columns."
- 5. Check the box of "Perform hypothesis test."
- 6. Enter the hypothesized value "7" into the box next to "Perform hypothesis test."

One-Sample t for the Me	an 🛛 📉
	One or more samples, each in a column
	HtBk
	·
	✓ Perform hypothesis test
	Hypothesized mean: 7
Select	Options <u>G</u> raphs
Help	<u>O</u> K Cancel

Fig. 3.34 One-Sample t for the Mean dialog box with data entered

- 7. Click "OK."
- 8. The one-sample t-test result appears automatically in the session window.

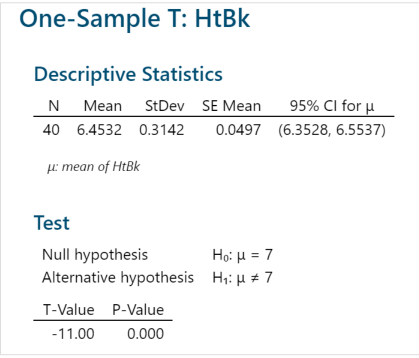


Fig. 3.35 One-Sample t for the Mean output

Since the p-value is smaller than alpha level (0.05), we reject the null hypothesis and claim that average height of our basketball players is statistically different from 7 feet.

What is Two Sample T-Test?

Two sample t-test is a hypothesis test to study whether there is a statistically significant difference between the means of two populations.

- Null Hypothesis (H₀): $\mu_1 = \mu_2$
- Alternative Hypothesis (H_a): $\mu_1 \neq \mu_2$

Where: μ_1 is the mean of one population and μ_2 is the mean of the other population of our interest.

Assumptions of Two Sample T-Tests

- The sample data drawn from both populations are unbiased and representative.
- The data of both populations are continuous.
- The data of both populations are normally distributed.
- The variances of both populations are unknown.
- Two sample t-test is more robust than a z-test when the sample size is small (< 30).

Three Types of Two Sample T-Tests

1. Two sample t-test when the variances of two populations are unknown but equal

Two sample t-test, equal variances; (when $\sigma^{2}_{1} = \sigma^{2}_{2}$)

2. Two sample t-test when the variances of the two population are unknown and unequal

Two sample t-test, unequal variances; (when $\sigma^{2_1} \neq \sigma^{2_2}$)

3. Paired t-test when the two populations are dependent of each other, so each data point from one distribution corresponds to a data point in the other distribution.

Test of Equal Variance

To check whether the variances of two populations of interest are significantly different on a statistical basis, we use the test of equal variance. An F-test is used to test this equality between two normally distributed populations.

- Null Hypothesis (H₀): $\sigma_1^2 = \sigma_2^2$
- Alternative Hypothesis (H₁): $\sigma_1^2 \neq \sigma_2^2$

An *F-test* is a statistic hypothesis test in which the test statistic follows an F-distribution when the null hypothesis is true. The most known F-test is the test of equal variance for two normally distributed populations. The F-test is very sensitive to non-normality. When any one

of the two populations is not normal, we use the Brown–Forsythe test for checking the equality of variances.

Test Statistic

To calculate the F statistic, you divide the squared standard deviation of one population by the other.

$$F_{calc} = \frac{s_1^2}{s_2^2}$$

Where: s_1 and s_2 are the sample standard deviations.

Critical Value

F_{crit} is the F value in a F distribution with the predetermined significance level α and degrees of freedom $(n_1 - 1)$ and $(n_2 - 1)$.

F_{crit} values for a two-sided and a one-sided F-test with the same significance level α and degrees of freedom ($n_1 - 1$) and ($n_2 - 1$) are different.

The critical value is a reference value that you will find in a table, using the prescribed parameters—the significance level alpha, and degrees of freedom for both samples. Based on the sample data, we calculated the test statistic F_{calc} , which is compared against F_{crit} to make a decision of whether to reject the null.

- Null Hypothesis (H₀): $\sigma_1^2 = \sigma_2^2$
- Alternative Hypothesis (H_a): $\sigma_1^2 \neq \sigma_2^2$
- If F_{calc} > F_{crit}, we reject the null and claim there is a statistically significant difference between the variances of the two populations.
- If F_{calc} < F_{crit}, we fail to reject the null and claim there is not any statistically significant difference between the variances of the two populations.

Test Statistic and Critical Value of a Two Sample T-Test when $\sigma_1^2=\sigma_2^2$

Test Statistic

For a two-sample t-test, the *test statistic* is calculated using the sample means for the two populations, the pooled standard deviation, and the sample sizes.

$$t_{calc} = \frac{\overline{Y}_1 - \overline{Y}_2}{s_{pooled} \sqrt{\frac{1}{n_1} + \frac{1}{n_2}}}$$
$$s_{pooled} = \sqrt{\frac{(n_1 - 1)s_1^2 + (n_2 - 1)s_2^2}{(n_1 + n_2 - 2)}}$$

Where:

- $\overline{Y_1}$ and $\overline{Y_2}$ are the sample means of the two populations of our interest.
- n_1 and n_2 are the sample sizes. n_1 is not necessarily equal to n_2 .

Spooled is a pooled estimate of variance. s1 and s2 are the sample standard deviations

Critical Value

- t_{crit} is the *t* value in a Student's t distribution with the predetermined significance level α and degrees of freedom $(n_1 + n_2 2)$.
- t_{crit} values for a two-sided and a one-sided t-test with the same significance level α and degrees of freedom $(n_1 + n_2 2)$ are different

Pooled variance is a method for estimating variance given several different samples taken in different circumstances where the mean may vary between samples but the true variance is assumed to remain the same. The critical t value is pulled from the Student's t distribution using the desired significance level and the degrees of freedom calculated by $(n_1 + n_2 - 2)$.

Test Statistic and Critical Value of a Two Sample T-Test when $\sigma_1^2\neq\sigma_2^2$

Test Statistic

For a two-sample t-test with unknown variances or with known unequal variances, the test statistic is not calculated using the pooled standard deviation.

$$t_{calc} = \frac{\overline{Y}_1 - \overline{Y}_2}{\frac{S_1^2}{n_1} + \frac{S_2^2}{n_2}}$$
$$\left(S_1^2 - S_2^2\right)^2$$

$$df = \frac{\left(\frac{s_1}{n_1} + \frac{s_2}{n_2}\right)}{\frac{\left(\frac{s_1}{n_1}\right)^2}{n_1 - 1} + \frac{\left(\frac{s_2}{n_2}\right)^2}{n_2 - 1}}$$

Where:

- $\overline{Y_1}$ and $\overline{Y_2}$ are the sample means of the two populations of our interest
- n_1 and n_2 are the sample sizes. n_1 is not necessarily equal to n_2 .

 s_{pooled} is a pooled estimate of variance. s_1 and s_2 are the sample standard deviations

Critical Value

• t_{crit} is the t value in a Student's t distribution with the predetermined significance level α and degrees of freedom *df* calculated using the formula above.

• *t_{crit}* values for a two-sided and a one-sided t-test with the same significance level *α* and degrees of freedom *df are different*.

Pooled variance is a method for estimating variance given several different samples taken in different circumstances where the mean may vary between samples but the true variance is assumed to remain the same. When we cannot assume equal variances or if we know they are unequal we cannot make the same assertions by using pooled variances. Therefore, we use the standard deviation of each sample separately in our t_{calc} equation.

Test Statistic and Critical Value of a Paired T-Test

Test Statistic

When using a paired t-test, the test statistic is calculated using the average difference between the data pairs, the standard deviation of the differences, and the sample size of either population.

$$t_{calc} = \frac{\overline{d}}{\frac{S_d}{\sqrt{n}}}$$

Where:

- *d* is the difference between each pair of data
- \overline{d} is the average of d
- *n* is the sample size of either population of interest.

Critical Value

- t_{crit} is the t value in a Student's t distribution with the predetermined significance level α and degrees of freedom (n 1).
- *t*_{crit} values for a two-sided and a one-sided t-test with the same significance level α and degrees of freedom (*n* 1) are different.

The critical t is looked up using the significance level alpha and degrees of freedom (n – 1).

Decision Rules of a Two-Sample T-Test

Based on the sample data, we calculated the test statistic t_{calc} , which is compared against t_{crit} to make a decision of whether to reject the null.

- Null Hypothesis (H₀): $\mu_1 = \mu_2$
- Alternative Hypothesis (H_a): $\mu_1 \neq \mu_2$

If $|t_{calc}| > t_{crit}$, we reject the null and claim there is a statistically significant difference between the means of the two populations.

If $|t_{calc}| < t_{crit}$, we fail to reject the null and claim there is not any statistically significant difference between the means of the two populations.

Use Minitab to Run a Two-Sample T-Test

Case study: Compare the average retail price of a product in state A and state B.

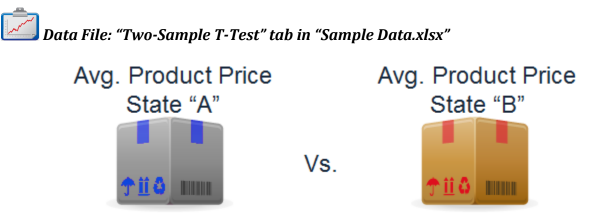


Fig. 3.36 Two-Sample T-Test comparison

- Null Hypothesis (H₀): $\mu_1 = \mu_2$
- Alternative Hypothesis (H_a): $\mu_1 \neq \mu_2$

In this example, we will be comparing the average price of a product in two different states. The null hypothesis is that the price in state A is equal to the price in state B.

Step 1: Test the normality of the retail price for both state A and B.

- 1. Click Stat \rightarrow Basic Statistics \rightarrow Graphical Summary.
- 2. A window named "Graphical Summary" pops up.
- 3. Select "Retail Price" as the "Variables."
- 4. Click in the blank box right below "By variables (optional)" and "State" appears in the list box on the left.

5. Select the "State" as the "By variables (optional)."

Graphical Summary	— X —
C1 State C2 Retail Price	Variables: 'Retail Price'
	Confidence level: 95.0
Select	
Help	OK Cancel

Fig. 3.37 Graphical Summary dialog box with chosen variables

- 6. Click "OK."
- 7. The normality test results would appear in the new windows.

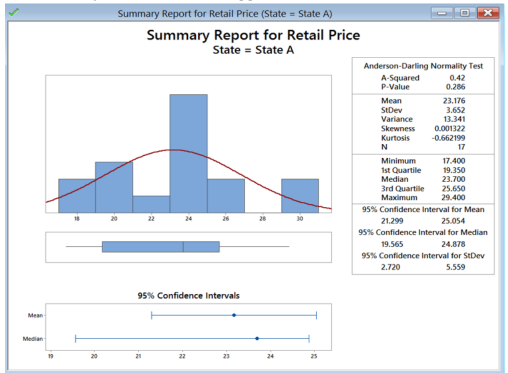


Fig. 3.38 Graphical Summary output for State A

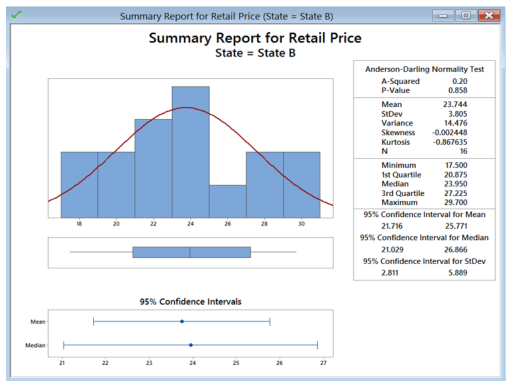


Fig. 3.39 Graphical Summary output for State B

- Null Hypothesis (H₀): The data are normally distributed.
- Alternative Hypothesis (H_a): The data are not normally distributed.

Both retail price data of state A and B are normally distributed since the p-values are both greater than alpha level (0.05). If any of the data series is not normally distributed, we need to use other hypothesis testing methods other than the two-sample t-test.

By following the instructions on the previous page, we first determine if the data follow a normal distribution. In this case, you can see that the p-value for both is higher than 0.05, so we fail to reject the null hypothesis that the data are normally distributed. If the data are not normally distributed, we must use a different test.

Step 2: Test whether the variances of the two data sets are equal.

- Null Hypothesis (H₀): $\sigma_1^2 = \sigma_2^2$
- Alternative Hypothesis (H_a): $\sigma_1^2 \neq \sigma_2^2$
- 1. Click Basic Statistics \rightarrow 2 Variances.
- 2. A new window named "Two-Sample Variance" pops up.
- 3. Click in the blank box of "Samples" and "Retail Price" appears in the list box on the left.
- 4. Select "Retail Price" as the "Samples."
- 5. Click in the blank box of "Sample IDs" and "State" appears in the list box on the left.

6. Select "State" as the "Sample IDs."

Two-Sample Variance	×
C1 State C2 Retail Price	Both samples are in one column Samples: 'Retail Price' Sample IDs: State
Select	Options <u>G</u> raphs <u>R</u> esults
Help	<u>O</u> K Cancel

Fig. 3.40 Two Sample Variance dialog box with sample IDs selection

7. Click "Options" and select

"Use test and confidence intervals based on normal distribution"

Two-Sample Variance: Options	X
Ratio: (sample 1 standard deviation) /	(sample 2 standard deviation) 💌
Confidence level: 95.0	
Hypothesized ratio:	
Alternative hypothesis: Ratio≠ hypoth	esized ratio
✓ Use test and confidence intervals ba	sed on normal distribution
Help	<u>Q</u> K Cancel

Fig. 3.41 Two Sample Variance Options

- 8. Click "OK." in the "Options" window.
- 9. Click "OK" in the 2-Sample t window

cot un	d C	I for 1	Two Va	ariar	nces: Re	etail P	rice	vs S	tate
Metho	k								
σ₁: stand	lard d	leviation	of Retail F	rice w	hen State =	State A			
σ ₂ : stand	lard d	leviation	of Retail F	rice w	hen State =	State B			
Ratio: σ ₁									
F metho	d was	used. Th	is method	d is acc	urate for n	ormal dat	a only.		
Descrip	tivo	Statict	ice						
State	N	StDev	Variance	05%	CI for a				
State A	17	3.652	13.341		0, 5.559)				
State B		3.805		-	1, 5.889)				
		95%							
	itio	Ratio F	:						
Ra 0.9599	itio		:						
Ra 0.9599 Test	ntio 986	(0.570,	1.603)	: σ ₁ /	σ ₂ = 1				
Ra 0.9599 Test Null hy	ntio 986 poth	(0.570, nesis	: 1.603) H₀						
Ra 0.9599 Test Null hy	ntio 986 poth	(0.570, nesis hypothe	1.603)	: σ ₁ /	σ ₂ ≠ 1				
Ra 0.9599 Test Null hy Alterna	ntio 986 poth	(0.570, nesis hypothe		: σ ₁ /	σ ₂ ≠ 1				
Ra 0.9599 Test Null hy Alterna	poth tive	(0.570, nesis hypothe e level	H ₀ H ₀ esis H ₁	: σ ₁ /	σ ₂ ≠ 1				

Fig. 3.42 Two Sample Variance output

Because the retail prices at state A and state B are both normally distributed, an F test is used to test their variance equality. The p-value of F test is 0.870, greater than the alpha level (0.05), so we fail to reject the null hypothesis and we claim that the variances of the two data sets are equal. We will use the two-sample t-test (when $\sigma^{2}_{1} = \sigma^{2}_{2}$) to compare the means of the two groups. If $\sigma^{2}_{1} \neq \sigma^{2}_{2}$, we will use the two-sample t-test (when $\sigma^{2}_{1} \neq \sigma^{2}_{2}$) to compare the means of the two groups.

Step 3: Run two-sample t-test to compare the means of two groups.

- 1. Click Stat \rightarrow Basic Statistics \rightarrow 2-Sample t.
- 2. A new window named "Two-Sample t for the Mean" pops up.
- 3. Click in the blank box right next to "Samples" and the "Retail Price" appears in the list box on the left.
- 4. Select "Retail Price" as the "Samples."

- 5. Click in the blank box right next to "Sample IDs" and the "State" appears in the list box on the left.
- 6. Select "State" as the "Subscripts."

Two-Sample t for the Me	ean 🗾
C1 State C2 Retail Price	Both samples are in one column
C2 RetairFrice	Samples: Retail Price'
	Sample IDs: State
Select	Options
Help	<u>Q</u> K Cancel

Fig. 3.43 Two-Sample t for the mean dialog box with variable selections

- 7. Click "Options"
- 8. Check the box "Assume equal variances."

Two-Sample t: Options	X
Difference = (sample 1 me	ean) - (sample 2 mean)
<u>C</u> onfidence level:	95.0
Hypothesized difference:	0.0
Alternative hypothesis:	Difference ≠ hypothesized difference 💽
Assume equal variance	25
Help	<u>O</u> K Cancel

Fig. 3.44 Two-Sample t: Options dialog box

- 9. Click "OK." in the "Options" window.
- 10. Click "OK" in the 2-Sample t window

The results of the two-sample t-test (when $\sigma_1 = \sigma_2$) appear in the session window.

Descrip	tive	Statis	stics: R	etail Prie
State	Ν	Mean	StDev	SE Mean
State A	17	23.18	3.65	0.89
State B	16	23.74	3.80	0.95
Estimat	ion	for Di	fferen	ce
	F	Pooled	95% CI	for
Differenc	e	StDev	Differe	nce
-0.5	7	3.73	(-3.21, 2	2.08)
Test				
Test				
Null hyp	othes	sis	H₀: μ₁	1 - μ ₂ = 0
Alternati	ve hy	pothesi	s H₁: μ₁	ı - μ₂ ≠ 0
T-Value	DF	P-Valu	le	
-0.44	31	0.66	55	

Fig. 3.45 Two-Sample t output

Since the p-value of the t-test (assuming equal variance) is 0.665, greater than the alpha level (0.05), we fail to reject the null hypothesis and we claim that the means of the two data sets are equal. If the variances of the two groups do not equal, we will need to use the two-sample t-test (when $\sigma_1 \neq \sigma_2$) to compare the means of the two groups. In the window of "2-Sample t: Options)", uncheck the box next to "Assume equal variances" and run the 2-sample t-test again.

Two-Sample t: Options	— X —
Difference = (sample 1 me	an) - (sample 2 mean)
Confidence level:	95.0
Hypothesized difference:	0.0
Alternative hypothesis:	Difference ≠ hypothesized difference 💌
Assume equal variance	15
· · · ·	
Help	<u>Q</u> K Cancel

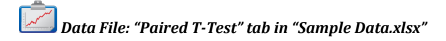
Fig. 3.46 Two-Sample t: Options dialog box

State N Mean StDev SE Mean State A 17 23.18 3.65 0.89 State B 16 23.74 3.80 0.99 Estimation for Difference 95% Cl for Difference Difference -0.57 (-3.22, 2.09)	Descriptive Statistics: Retail Price			
State B 16 23.74 3.80 0.99 Estimation for Difference 95% Cl for Difference Difference -0.57 (-3.22, 2.09)	n			
Estimation for Difference 95% CI for Difference -0.57 (-3.22, 2.09)	9			
95% CI for Difference -0.57 (-3.22, 2.09)	5			
Difference-0.57(-3.22, 2.09)	Estimation for Difference			
-0.57 (-3.22, 2.09)				
Test	-0.57 (-3.22, 2.09)			
Test				
Null hypothesis $H_0: \mu_1 - \mu_2 = 0$				
Alternative hypothesis $H_1: \mu_1 - \mu_2 \neq 0$				
T-Value DF P-Value				
-0.44 30 0.666				

Fig. 3.47 Two-Sample t output without assuming equal variances

Since the p-value of the t-test (assuming unequal variance) is 0.666, greater than the alpha level (0.05), we fail to reject the null hypothesis and we claim that the means of two groups are equal.

Case study: We are interested to know whether the average salaries (\$1000/yr.) of male and female professors at the same university are the same.



The data were randomly collected from 22 universities. For each university, the salaries of male and female professors were randomly selected.

- Null Hypothesis (H₀): $\mu_{male} \mu_{female} = 0$
- Alternative Hypothesis (H_a): $\mu_{male} \mu_{female} \neq 0$

Use Minitab to Run a Paired T-Test

Step 1: Create a new column for the difference between data sets, name it "Difference."

- 1. Right click on the newly-generated column, "Difference."
- 2. Select "Formulas" and then "Assign Formula to Column."
- 3. A new window named "Assign Formula to C3 (Difference)" pops up.
- 4. Enter "'MALES'-'FEMALES' " into the box right below "Expression."

Assign Formula to C3	(Difference)	x
C1 MALES C2 FEMALES	Expression: 'MALES'-'FEMALES'	*
		Ŧ
	Functions:	
	7 8 9 + = ≠ All functions	-
	4 5 6 - < > Absolute value	*
	1 2 3 * ≤ ≥ Antilog Any	
	0 . [] / And Arccosh Arccosine	
1	Or Arcsine	-
Select	() Not Select	
Help	OK Can	icel

Fig. 3.48 Assign Formula dialog box with formula

5. Click "OK."

6. The values would appear in the column "Difference."

Ŧ	C1	C2	C3
	MALES	FEMALES	DIFFERENCE
1	34.5000	33.9000	0.60000
2	30.5000	31.2000	-0.70000
3	35.1000	35.0000	0.10000
4	35.7000	34.2000	1.50000
5	31.5000	32.4000	-0.90000
6	34.4000	34.1000	0.30000
7	32.1000	32.7000	-0.60000
8	30.7000	29.9000	0.80000
9	33.7000	31.2000	2.50000
10	35.3000	35.5000	-0.20000
11	30.7000	30.2000	0.50000
12	34.2000	34.8000	-0.60000
13	39.6000	38.7000	0.90000
14	30.5000	30.0000	0.50000
15	33.8000	33.8000	0.00000
16	21 7000	20 1000	0 70000

Fig. 3.49 Formula output

Step 2: Test whether the difference is normally distributed.

- Null Hypothesis (H₀): The difference between two data sets is normally distributed.
- Alternative Hypothesis (H_a): The difference between two data sets is not normally distributed.
- 1. Click Stat \rightarrow Basic Statistics \rightarrow Normality Test.
- 2. A new window named "Normality Test" pops up.
- 3. Select "Difference" as the "Variable."

Normality Test	
	Variable: Difference
	Percentile Lines None
	O At data values:
	Tests for Normality
	Anderson-Darling Ryan-Joiner (Similar to Shapiro-Wilk)
	C Kolmogorov-Smirnov
Select	Title:
Help	OK Cancel

Fig. 3.50 Normality Test dialog box with variable

4. Click "OK."

5. The normality test results would appear in the new window.

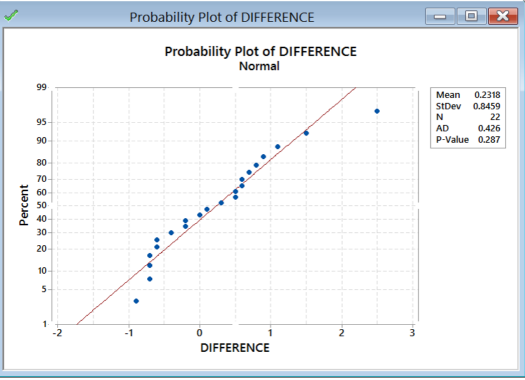


Fig. 3.51 Normality Test output

The p-value of the normality test is 0.287, greater than the alpha level (0.05), so we fail to reject the null hypothesis and we claim that the difference is normally distributed. When the difference is not normally distributed, we need other hypothesis testing methods.

Step 3: Run the paired t-test to compare the means of two dependent data sets.

- 1. Click Stat \rightarrow Basic Statistics \rightarrow Paired t.
- 2. A new window named "Paired t (Test and Confidence Interval)" pops up.
- 3. Click in the blank box right next to "First Sample" and the three columns names appear in the list box on the left.
- 4. Select "MALES" as "First Sample."

5. Select "FEMALES" as "Second Sample."

Paired t for the Mean	×
C1 MALES C2 FEMALES C3 Difference	Each sample is in a column Sample 1: MALES Sample 2: FEMALES
Select	Options <u>G</u> raphs
Help	<u>Q</u> K Cancel

Fig. 3.52 Paired t for the Mean dialog box with selected variables

- 6. Click "OK."
- 7. The paired t-test results appear in the session window.

Descript	ve S	Statistic	CS	
Sample	Ν	Mean	StDev	SE Mean
MALES	22	33.432	3.546	0.756
FEMALES	22	33.200	3.601	0.768
Estimatio	on fo	or Paire	ed Diff	erence
			95%	6 CI for
Mean St	Dev	SE Mea	n μ_di	fference
0.232 0.	846	0.18	0 (-0.14	43, 0.607)
μ_difference	$\mu_difference: mean of (MALES - FEMALES)$			
Test				
Null hypot	hesis		H₀: μ_dif	ference = 0
Alternative	e hyp	othesis	H₁: μ_dif	ference ≠ 0
T-Value	P-Val	ue		
1.29	0.2	13		

Fig. 3.53 Paired t output

The p-value of the paired t-test is 0.213, greater than the alpha level (0.05), so we fail to reject the null hypothesis and we claim that there is no statistically significant difference between the

salaries of male and those of female professors. Because the p-value is higher than 0.05, we again fail to reject the null hypothesis, meaning that there is no statistically significant difference between the salaries of male and female professors.

3.3.2 One Sample Variance

What is One Sample Variance Test?

One sample variance test is a hypothesis testing method to study whether there is a statistically significant difference between a population variance and a specified value.

- Null Hypothesis (H₀): $\sigma^2 = \sigma_0^2$
- Alternative Hypothesis (H_a): $\sigma^2 \neq \sigma_0^2$

Where: σ^2 is the variance of a population of our interest and σ_0^2 is the specific value we want to compare against. The null hypothesis is that the population variance is equal to a specified value.

Chi-Square Test

A *chi-square test* is a statistical hypothesis test in which the test statistic follows a chi-square distribution when the null hypothesis is true. A chi-square test can be used to test the equality between the variance of a normally distributed population and a specified value. In a chi-square test, the test statistic follows a chi-square distribution when the null hypothesis is true. The test can be used to test the equality of a variance to a specified value. For the test to be valid, the distribution must be normally distributed.

Test Statistic

The test statistic is calculated using the observed variance of the sample from the population, as well as the sample size and the specified value for variance we are comparing against.

$$\chi_{calc}^{2} = \frac{(n-1)s^{2}}{\sigma_{0}^{2}}$$

Where:

- s^2 is the observed variance and *n* is the sample size
- σ_0^2 is the specified value we compare against.

Critical Value

- χ_{crit^2} is the χ^2 value in a χ^2 distribution with the predetermined significance level α and degrees of freedom (n 1).
- χ_{crit^2} values for a two-sided and a one-sided χ^2 -test with the same significance level α and degrees of freedom (n 1) are different.

Like in other tests we discussed, we compare the calculated chi-square statistic against a critical value, which is a chi-square table value that can be located based on the determined significance level and (n - 1) degrees of freedom. Based on the sample data, we calculated the test statistic χ_{calc}^2 , which is compared against χ_{crit}^2 to make a decision of whether to reject the null.

- Null Hypothesis (H₀): $\sigma^2 = \sigma_0^2$
- Alternative Hypothesis (H_a): $\sigma^2 \neq \sigma_0^2$
- If $|\chi_{calc}| > \chi_{crit}^2$, we reject the null and claim there is a statistically significant difference between the population variance and the specified value.

• If $|\chi_{calc}| < \chi_{crit}^2$, we fail to reject the null and claim there is not any statistically

significant difference between the population variance and specified value.

In other words, if the absolute value of the statistic is greater that the critical value, then we reject the null and claim that the population variance is different than the specified value. Otherwise, we fail to reject the null.

Use Minitab to Run a One-Sample Variance Test

Case study: We are interested in comparing the variance of the height of basketball players with zero.

Data File: "One-Sample T-Test" tab in "Sample Data.xlsx"

- Null Hypothesis (H₀): $\sigma^2 = 0$
- Alternative Hypothesis (H_a): $\sigma^2 \neq 0$

In this case study, we will use the software to determine if the variance in the height of basketball players is zero or different from zero.

Step 1: Test whether the data are normally distributed

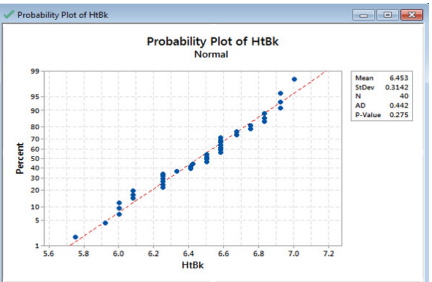
- 1. Click Stat \rightarrow Basic Statistics \rightarrow Normality Test.
- 2. A new window named "Normality Test" pops up.

3. Select "HtBk" as the variable.

Normality Test	— ×—
	Variable: HtBk Percentile Lines (None At Y values: At data values: Tests for Normality (Anderson-Darling C Ryan-Joiner (Similar to Shapiro-Wilk)
1	C Kolmogorov-Smirnov
Select	Title:
Help	OK Cancel

Fig. 3.54 Normality Test dialog box with variable

- 4. Click "OK."
- 5. A new window named "Probability Plot of HtBk" appears, which covers the results of the normality test.





- Null Hypothesis (H₀): The data are normally distributed.
- Alternative Hypothesis (H_a): The data are not normally distributed.

Since the p-value of the normality is 0.275, which is greater than alpha level (0.05), we fail to reject the null and claim that the data are normally distributed. If the data are not normally distributed, you need to use other hypothesis tests. Recall that the data must be normally distributed in order for a one-sample variance test to be valid. As you can see from the results, the p-value is greater than 0.05, so we fail to reject the null hypothesis that the data are normally distributed.

Step 2: Check whether the population variance is equal to the specified value (zero).

- 1. Click Stat \rightarrow Basic Statistics \rightarrow 1 Variance.
- 2. A new window named "One-Sample Variance" appears.
- 3. Click in the blank box right below "Columns" and the "HtBk" appears in the list box on the left.
- 4. Check the box of "Perform hypothesis test."
- 5. Select "Hypothesized variance" from the dropdown menu.
- 6. Enter "0" as the "Hypothesized variance."

One-Sample Variance	
	One or more samples, each in a column
	HtBk
	Perform hypothesis test Value: Hypothesized standard deviation I
Select	Optio <u>n</u> s
Help	<u>Q</u> K Cancel

Fig. 3.56 One-Sample Variance dialog box with selected variables

- 7. Click "OK."
- 8. An alert window pops up saying "Invalid hypothesized value. The following value is too small: "0". Please specify: A single numeric constant. Value > 0."

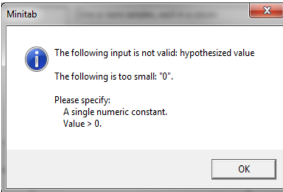


Fig. 3.57 Minitab error alert

9. Click "OK" in the alert message window.

10. Enter "0.000000001" as the "Hypothesized variance" instead.

One-Sample Variance	
	One or more samples, each in a column
	HtBk
	✓ Perform hypothesis test Value: Hypothesized standard deviation ✓ 0.000000001
Select	Optio <u>n</u> s
Help	QK Cancel

Fig. 3.58 One-Sample Variance dialog box with new hypothesized variance value

11. Click "OK."

12. The one-sample variance test appears in new window.

	-		• •			
Test	and	CI for C	One Vari	iance:	HtBk	
Me	thod					
		deviation o				
			alid for any o			
The	e chi-squa	are method	is valid only	for the n	ormal distril	oution.
Des	scriptiv	e Statist	ics			
				95%	CI for σ	
			95% CI for	σι	using	
Ν	StDev	Variance	using Bone	ett Chi	-Square	
40	0.314	0.0987	(0.266, 0.39	90) (0.25	7, 0.403)	
Tes	t					
Nu	ll hypoth	esis	$H_0: \sigma^2 = 0.0$	00000000	01	
Alt	ernative ł	nypothesis	H ₁ : σ ² ≠ 0.	.00000000	001	
Me	thod	Test Stati	stic DF P	-Value		
Bo	nett			0.000		
Ch	-Square	3.85068E	+10 39	0.000		

Fig. 3.59 One-Sample Variance output

Since the p-value of the one-sample variance test is smaller than alpha level (0.05), we reject the null and claim that the population variance is statistically different from zero.

Another way to see whether the population variance is statistically different from zero is to check whether zero stays between the upper and lower confidence interval boundaries of the variance. If yes, we fail to reject the null hypothesis and claim that the population variance is not statistically different from zero. Since the p-value is less than the alpha level, we must reject the null hypothesis that the population variance is equal to zero.

Another way to interpret the results: we can check to see if the specified variance falls within the confidence interval (CI) for the population variance. If it does, then the population variance is statistically the same as the specified value.

3.3.3 ONE-WAYANOVA

What is One-Way ANOVA?

One-way ANOVA is a statistical method to compare means of two or more populations.

- Null Hypothesis (H₀): $\mu_1 = \mu_2 = ... = \mu_k$
- Alternative Hypothesis (H_a): At least one μ_i is different, where *i* is any value from 1 to k.

It is a generalized form of the two-sample t-test since a two-sample t-test compares two population means and one-way ANOVA compares k population means where $k \ge 2$.

Assumptions of One-Way ANOVA

- The sample data drawn from *k* populations are unbiased and representative.
- The data of *k* populations are continuous.
- The data of *k* populations are normally distributed.
- The variances of *k* populations are equal.

How ANOVA Works

ANOVA compares the means of different groups by analyzing the variances between and within groups. Let us say we are interested in comparing the means of three normally distributed populations. We randomly collected one sample for each population of our interest.

- Null Hypothesis (H₀): $\mu_1 = \mu_2 = \mu_3$
- Alternative Hypothesis (H_a): One of the μ is different from the others.

Based on the sample data, the means of the three populations might look different because of two variation sources.

- 1. Variation between groups there are non-random factors leading to the variation between groups.
- 2. Variation within groups there are random errors resulting in the variation within each individual group.

What we care about the most is the variation between groups since we are interested in whether the groups are statistically different from each other. Variation between groups is the *signal* we want to detect and variation within groups is the *noise* which corrupts the signal.

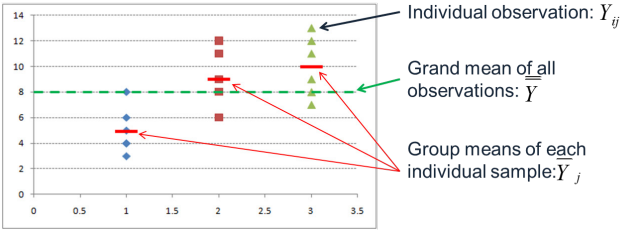


Fig. 3.60 How ANOVA works

Total Variation = SS(Total) =
$$\sum_{j=1}^{k} \sum_{i=1}^{n_i} (Y_{ij} - \overline{\overline{Y}})^2$$

Between Variation = SS(Between) = $\sum_{j=1}^{k} n_j (\overline{Y_j} - \overline{\overline{Y}})^2$
Within Variation = SS(Within) = $\sum_{j=1}^{k} \sum_{i=1}^{n_i} (Y_{ij} - \overline{Y_j})^2$

Using this visual, the purpose of using the ANOVA is to determine if there is a difference between the data points (the group means of the individual samples). The variation within the groups can make it difficult to detect the difference between the group means.

Variation Components

Total Variation = Variation Between Groups + Variation Within Groups

$$\sum_{j=1}^k \sum_{i=1}^{n_i} \bigl(Y_{ij} \overline{\overline{Y}}\bigr)^2 = \sum_{j=1}^k n_j \bigl(Y_j \overline{\overline{Y}}\bigr)^2 + \sum_{j=1}^k \sum_{i=1}^{n_i} \bigl(Y_{ij} \overline{Y_j}\bigr)^2$$

Total Variation = Sums of squares of the vertical difference between the individual observation and the grand mean Variation Between Groups = Sums of squares of the vertical difference between the group mean and the grand mean. Variation Within Groups = Sum of squares of the vertical difference between the individual observation and the group mean

Degrees of Freedom (DF)

In statistics, the degrees of freedom are the number of unrestricted values in the calculation of a statistic.

Degrees of Freedom Components

 $DF_{total} = DF_{between} + DF_{within}$ $DF_{total} = n-1$ $DF_{between} = k-1$ $DF_{within} = n-k$

Where:

- n is the total number of observations
- k is the number of groups.

Signal-to-Noise Ratio (SNR)

Earlier, the terms signal and noise were introduced. The *signal* is the variation between groups that we are trying to detect. The *noise* is the variation within a group that makes it difficult to see the signal.

SNR denotes the ratio of a signal to the noise corrupting the signal. It measures how much a signal has been corrupted by the noise. When it is higher than 1, there is more signal than noise. The higher the SNR, the less the signal has been corrupted by the noise.

F-ratio is the SNR in ANOVA

$$F = \frac{MS_{between}}{MS_{within}} = \frac{\frac{SS_{between}}{DF_{between}}}{\frac{SS_{within}}{DF_{within}}} = \frac{\sum_{j=1}^{k} (\overline{Y}_{j} - \overline{Y})^{2} / (k-1)}{\sum_{j=1}^{k} \sum_{i=1}^{n_{j}} (Y_{ij} - \overline{Y}_{j})^{2} / (n-k)}$$

In ANOVA, we use the F-test to compare the means of different groups. The F-ratio calculated as above is the test statistic F_{calc} .

The critical value (F_{cri}) in an F-test can be derived from the F table with predetermined significance level (α) and with (k – 1) degrees of freedom in the numerator and (n – k) degrees of freedom in the denominator.

How ANOVA Works

- Null Hypothesis (H₀): $\mu_1 = \mu_2 = ... = \mu_k$
- Alternative Hypothesis (H_a): At least one μ_i is different, where *i* is any value from 1 to *k*.

- If $|F_{calc}| < F_{crit}$, we fail to reject the null and claim that the means of all the populations of our interest are the same.
- If $|F_{calc}| > F_{crit}$, we reject the null and claim that there is at least one mean different from the others.

Using the F-ratio, we can compare the calculated value against the critical value. If the absolute value of the calculated value is less than the critical value, then we fail to reject the null hypothesis. If the absolute value of the calculated value is greater than the critical value, then we reject the null hypothesis.

Model Validation

ANOVA is a modeling procedure, which means we are using a model to try to predict results. To make sure the conclusions made in ANOVA are reliable, we need to perform residuals analysis.

Good residuals:

- Have a mean of zero
- Are normally distributed
- Are independent of each other
- Have equal variance

The difference between the actual and predicted result is called a *residual* or unexplained variation.

Use Minitab to Run an ANOVA

Case study: We are interested in comparing the average startup costs of five kinds of business.



Data File: "One-Way ANOVA" tab in "Sample Data.xlsx"

- Null Hypothesis (H₀): $\mu_1 = \mu_2 = \mu_3 = \mu_4 = \mu_5$
- Alternative Hypothesis (H_a): At least one of the five means is different from others.

Step 1: Test whether the data for each level are normally distributed.

- 1. Click Stat \rightarrow Basic Statistics \rightarrow Graphical Summary.
- 2. A new window named "Graphical Summary" pops up.
- 3. Select the "Cost" as the variable.
- 4. Click in the blank box right next to "By variables (optional)" and the "Business" appears in the list box on the left.

5. Select the "Business" as the "By variables (optional)."

Graphical Summary	— X —
C1 Business C2 Cost	Variables: Cost By variables (optional): Business T
	Confidence level: 95.0
Select	
Help	OK Cancel

Fig. 3.61 Graphical Summary dialog box with additional selected variables

- 6. Click "OK."
- 7. The normality results appear in the new window.

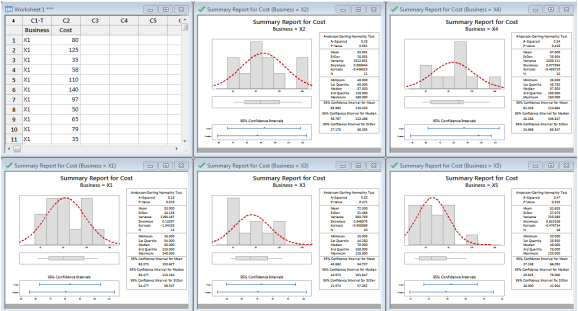


Fig. 3.62 Graphical Summary output

Notice all of the p-values are greater than 0.05; therefore, we fail to reject the null hypothesis that the data are normally distributed.

- Null Hypothesis (H₀): The data are normally distributed.
- Alternative Hypothesis (H_a): The data are not normally distributed.

Since the p-values of normality tests for the five data sets are higher than alpha level (0.05), we fail to reject the null hypothesis and claim that the startup costs for any of the five businesses are normally distributed. If any of the five data sets are not normally distributed, we need to use other hypothesis testing methods other than one-way ANOVA. In this example, all five data sets are normally distributed; however, if *any* of them were not normally distributed, we would need to use another hypothesis test.

Step 2: Test whether the variance of the data for each level is equal to the variance of other levels.

- Null Hypothesis (H₀): $\sigma_1^2 = \sigma_2^2 = \sigma_3^2 = \sigma_4^2 = \sigma_5^2$
- Alternative Hypothesis (H_a): at least one of the variances is different from others.
- 1. Click Stat \rightarrow ANOVA \rightarrow Test for equal variances.
- 2. A new window named "Test for Equal Variances" pops up.
- 3. Select the "Cost" as the "Response."
- 4. Select the "Business" as the "Factors."

Test for Equal Variances	
C1 Business C2 Cost	Response data are in one column for all factor levels Response: Cost Factors: Business •<
Select	Results Storage
Help	<u></u> Cancel

Fig. 3.63 Test for Equal Variances dialog box with response and factors variables

- 5. Click "Options."
- 6. Select "Use test based on normal distribution"

Test for Equal Variances: Options	
Confidence level: 95.0	
Se test based on normal distribut	tion
Help	<u>Q</u> K Cancel

Fig. 3.64 Test for Equal Variances: Options dialog box

7. Click "OK" to close the "Options" window

- 8. Click "OK" to run the test
- 9. The results show up in a new window.

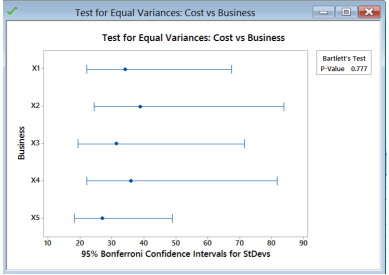


Fig. 3.65 Test for Equal Variances output

Use the Bartlett's test for testing the equal variances between five levels in this case since there are more than two levels in the data and the data of each level are normally distributed. The p-value of Bartlett's test is 0.777, greater than the alpha level (0.05), so we fail to reject the null hypothesis and we claim that the variances of five groups are equal. If the variances are not all equal, we need to use other hypothesis testing methods other than one-way ANOVA. If this test suggested that at least one variance was different, then we would need to use a different hypothesis test to evaluate the group means.

Step 3: Test whether the mean of the data for each level is equal to the means of other levels.

- Null Hypothesis (H₀): $\mu_1 = \mu_2 = \mu_3 = \mu_4 = \mu_5$
- Alternative Hypothesis (H_a): at least one of the means is different from others.
- 1. Click Stat \rightarrow ANOVA \rightarrow One-way.
- 2. A new window named "One-Way Analysis of Variance" pops up.
- 3. Select "Cost" as "Response."

4. Select "Business" as "Factor."

One-Way Analysis of Vari	ance 🗾
C1 Business C2 Cost	Response data are in one column for all factor levels Response: Cost Eactor: Business
Select	Options Comparisons Graphs Results Storage QK Cancel

Fig. 3.66 One-Way Analysis of Variance dialog box with response and factor variables selected

- 5. Click "Storage."
- 6. Check the boxes next to "Fits" and "Residuals"

One-Way Analysis of V	ariance: Storage	. X
Eits		
Help	<u>О</u> К	Cancel

Fig. 3.67 One-Way Analysis of Variance: Storage dialog box

- 7. Click "OK" to close the "Storage" window
- 8. Click "OK."
- 9. The ANOVA results appear in the session window. The fitted response and the residuals are stored in the data table.

One-way ANOVA: Cost versus Business

Method

Null hypothesis	All means are equal
Alternative hypothesis	Not all means are equal
Significance level	α = 0.05

Equal variances were assumed for the analysis.

Factor Information

Factor	Levels	Values
Business	5	X1, X2, X3, X4, X5

Analysis of Variance

Source	DF	Adj SS	Adj MS	F-Value	P-Value
Business	4	14298	3575	3.25	0.018
Error	55	60561	1101		
Total	59	74859			

Model Summary

S	R-sq	R-sq(adj)	R-sq(pred)
33.1829	19.10%	13.22%	3.18%

Means

Business	Ν	Mean	StDev	95% CI
X1	13	83.00	34.13	(64.56, 101.44)
X2	11	92.1	38.9	(72.0, 112.1)
X3	10	72.30	31.37	(51.27, 93.33)
X4	10	87.0	35.9	(66.0, 108.0)
X5	16	51.63	27.07	(35.00, 68.25)
Pooled StDev = 33.1829				

Fig. 3.68 One-Way Analysis of Variance output

Since the p-value of the F test is 0.018, lower than the alpha level (0.05), the null hypothesis is rejected and we conclude that the at least one of the means of the five groups is different from others.

Step 4: Test whether the residuals are normally distributed with mean equal zero. The residuals have been stored in the data table in step 4.

- 1. Click Stat \rightarrow Basic Statistics \rightarrow Graphical Summary.
- 2. A new window named "Graphical Summary" appears.
- 3. Select "RESI" as the "Variables."

Graphical Summary	×
C1 Business C2 Cost C3 FITS C4 RESI	Variables:
	By variables (optional):
	95.0
Select	
Help	OK Cancel

Fig. 3.69 Graphical Summary dialog box with variable selection

4. Click "OK."

5. The normality test results show up in a new window.

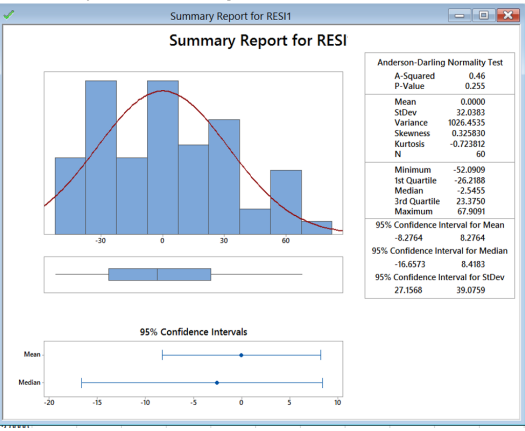


Fig. 3.70 Graphical Summary output

The p-value of the normality test is 0.255, greater than the alpha level (0.05), and we conclude that the residuals are normally distributed. The mean of the residuals is 0.0000.

Step 5: Check whether the residuals are independent of each other. If the residuals are in time order, we can plot IR charts to check the independence. When no data points on the IR charts fail any tests, the residuals are independent of each other. If the residuals are not in time order, the IR charts cannot deliver reliable conclusion on the independence.

- 1. Click Stat \rightarrow Control Charts \rightarrow Variables Charts for Individuals \rightarrow I-MR.
- 2. A new window named "Individuals Moving Range Chart" pops up.

3. Select "RESI1" as the "Variables."

Individuals-Moving Range	Chart		×
C2 Cost C3 FITS C4 RESI	<u>V</u> ariables:		×
Select	<u>S</u> cale <u>M</u> ultiple Graphs	Labels Data Options	I-MR Options
Help		<u>о</u> к	Cancel

Fig. 3.71 Individuals-Moving Range Chart dialog box with variables 4. Click "OK."

Next, we check to determine if the residuals are independent of each other. In Minitab, we use an Individuals–Moving Range control chart (called IR here) to determine independence. If the data points are in control (not failing any of the control chart tests) we can conclude that the residuals are independent.

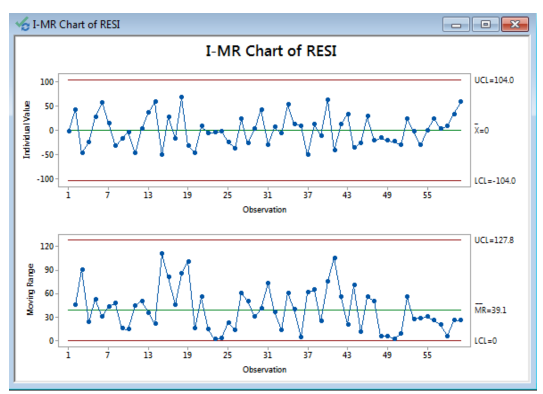


Fig. 3.72 Individuals-Moving Range Chart output

No tests failed; therefore, the residuals are independent.

Step 6: Plot residuals versus fitted values and check whether there is any systematic pattern.

- 1. Click Graph \rightarrow Scatter Plot.
- 2. A new window named "Scatterplots" pops up.
- 3. Click "OK."
- 4. A new window named "Scatterplot Simple" pops up.
- 5. Select the "RESI" as the "Y variables."
- 6. Select the "FITS" as "X variables."

Scatt	erplot: Simple				×
C2 C3 C4	Cost FITS RESI	 Y variables RESI ale	X variables FITS	•	Data View
	Select				
	Help		<u>о</u> к		Cancel

Fig. 3.73 Scatterplot: Simple dialog box with selected variables

- 7. Click "OK."
- 8. The charts appear in a new window.

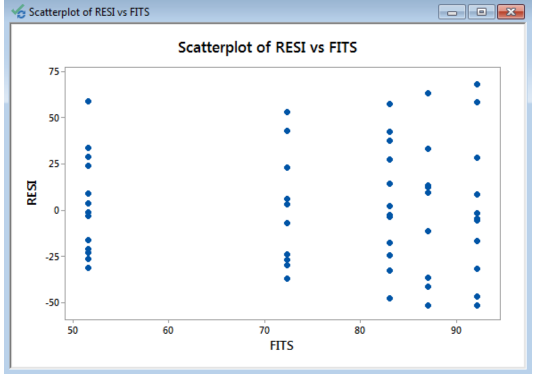


Fig. 3.74 Scatterplot output

If the data points spread out evenly at any of the five levels, we claim that the residuals have equal variances across the five levels.

3.4 Hypothesis Testing Non-Normal Data

The tests we have discussed up to this point all apply when we are working with data that follow normal distributions. So, what do we do if that is not the case? The following pages describe tests available to us when the data do not follow the normal distribution.

3.4.1 MANN-WHITNEY

What is the Mann-Whitney Test?

The *Mann–Whitney test* (also called Mann–Whitney U test or Wilcoxon rank-sum test) is a statistical hypothesis test to compare the medians of two populations that are not normally distributed. In a non-normal distribution, the median is the better representation of the center of the distribution.

- Null Hypothesis (H₀): $\eta_1 = \eta_2$
- Alternative Hypothesis (H_a): $\eta_1 \neq \eta_2$

Where:

- η_1 is the median of one population
- η_2 is the median of the other population.

The null hypothesis is that the medians are equal, and the alternative is that they are not.

Mann-Whitney Test Assumptions

- The sample data drawn from the populations of interest are unbiased and representative.
- The data of both populations are continuous or ordinal when the spacing between adjacent values is not constant. (Reminder: Ordinal data—A set of data is said to be ordinal if the values can be ranked or have a rating scale attached. You can count and order, but not measure, ordinal data.)
- The two populations are independent to each other.
- The Mann–Whitney test is robust for the non-normally distributed population.
- The Mann–Whitney test can be used when shapes of the two populations' distributions are different.

How Mann-Whitney Test Works

Step 1: Group the two samples from two populations (sample 1 is from population 1 and sample 2 is from population 2) into a single data set and then sort the data in ascending order ranked from 1 to n, where n is the total number of observations.

Step 2: Add up the ranks for all the observations from sample 1 and call it R_1 . Add up the ranks for all the observations from sample 2 and call it R_2 .

Step 3: Calculate the test statistics

$$U = \min(U_1, U_2)$$

Where:

$$U_1 = n_1 n_2 + \frac{n_1 (n_1 + 1)}{2} - R_1$$

$$U_2 = n_1 n_2 + \frac{n_2 (n_2 + 1)}{2} - R_2$$

and where:

- *n*¹ and *n*² are the sample sizes
- R_1 and R_2 are the sum of ranks for observations from sample 1 and 2 respectively.

Step 4: Make a decision on whether to reject the null hypothesis.

- Null Hypothesis (H₀): $\eta_1 = \eta_2$
- Alternative Hypothesis (H_a): $\eta_1 \neq \eta_2$

If both of the sample sizes are smaller than 10, the distribution of U under the null hypothesis is tabulated.

- The test statistic is U and, by using the Mann–Whitney table, we would find the p-value.
- If the p-value is smaller than alpha level (0.05), we reject the null hypothesis.
- If the p-value is greater than alpha level (0.05), we fail to reject the null hypothesis.

If both sample sizes are greater than 10, the distribution of U can be approximated by a normal distribution. In other words, $\frac{U-\mu}{\sigma}$ follows a standard normal distribution.

$$Z_{calc} = \frac{U - \mu}{\sigma}$$

Where:

$$\mu = \frac{n_1 n_2}{2}$$

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$$\sigma = \sqrt{\frac{\sqrt{n_1 n_2 (n_1 + n_2 + 1)}}{12}}$$

If the sample sizes are greater than 10, then the distribution of U can be approximated by a normal distribution. The U value is then plugged into the formula seen here to calculate a Z statistic.

When $|Z_{calc}|$ is greater than Z value at $\alpha/2$ level (e.g., when $\alpha = 5\%$, the z value we compare $|Z_{calc}|$ to is 1.96), we reject the null hypothesis.

Use Minitab to Run a Mann-Whitney Test

Case study: We are interested in comparing customer satisfaction between two types of customers using a nonparametric (i.e., distribution-free) hypothesis test: Mann–Whitney test.



Fig. 3.75 Mann-Whitney Test

- Null Hypothesis (H₀): $\eta_1 = \eta_2$
- Alternative Hypothesis (H_a): $\eta_1 \neq \eta_2$

Steps to run a Mann–Whitney Test in Minitab:

- 1. Un-stack the data into two separate columns:
 - "Overall Satisfaction 1" for customer type = 1
 - "Overall Satisfaction 2" for customer type = 2
- 2. Click Stat \rightarrow Nonparametrics \rightarrow Mann–Whitney.
- 3. A new window named "Mann–Whitney" pops up.
- 4. Select "Overall Satisfaction 1" as the "First Sample."

5. Select "Overall Satisfaction 2" as the "Second Sample."

Mann-Whitney	×
	First Sample: all Satisfaction_1'
	Second Sample: all Satisfaction_2
	Confidence level: 95.0
	Alternative: not equal
Select	
Help	OK Cancel

Fig. 3.76 Mann-Whitney dialog box with selected variables

- 6. Click "OK."
- 7. The Mann–Whitney test results appear in the session window.

Mann-Whitney: Overall Satisfaction_1, Overall Sat

Method

 $\begin{array}{l} \eta_1: \mbox{ median of Overall Satisfaction_1} \\ \eta_2: \mbox{ median of Overall Satisfaction_2} \\ \mbox{ Difference: } \eta_1 - \eta_2 \end{array}$

Descriptive Statistics

Sample	Ν	Median
Overall Satisfaction_1	31	3.56
Overall Satisfaction_2	42	4.34

Estimation for Difference

Difference	CI for Differen		hieved idence	
-0.81	(-1.17, -0	.45) 9	5.11%	
Test				
Null hypoth	esis	H₀: η₁ - ι	$\eta_2 = 0$	
Alternative	hypothesis	H ₁ : η ₁ - ι	η₂ ≠ 0	
Method		W-Value	P-Value	
Not adjuste	d for ties	772.50	0.000	
Adjusted fo	r ties	772.50	0.000	

Fig. 3.77 Mann-Whitney Test output

The p-value of the test is lower than alpha level (0.05); so we reject the null hypothesis and conclude that there is a statistically significant difference between the overall satisfaction medians of the two customer types.

3.4.2 Kruskal-Wallis

Kruskal-Wallis One-Way Analysis of Variance

The *Kruskal–Wallis one-way analysis of variance* is a statistical hypothesis test to compare the medians among more than two groups.

- Null Hypothesis (H₀): $\eta_1 = \eta_2 = ... = \eta_k$
- Alternative Hypothesis (H_a): at least one of the medians is different from others.

Where:

- η_i is the median of population i
- k is the number of groups of our interest.

It is an extension of the Mann–Whitney test. While the Mann–Whitney test allows us to compare the samples of two populations, the Kruskal–Wallis test allows us to compare the samples of more than two populations.

One key difference between this test and the Mann–Whitney test is the robustness of the test when the populations are not identically shaped. If this is the case, there is a different test, called Mood's median, which is more appropriate.

Kruskal–Wallis One-Way Analysis of Variance: Assumptions

- The sample data drawn from the populations of interest are unbiased and representative.
- The data of k populations are continuous or ordinal when the spacing between adjacent values is not constant.
- The k populations are independent to each other.
- The Kruskal–Wallis test is robust for the non-normally distributed population.

How Kruskal-Wallis One-Way ANOVA Works

The Kruskal–Wallis test works very similarly to the Mann–Whitney Test.

Step 1: Group the k samples from k populations (sample i is from population i) into one single data set and then sort the data in ascending order ranked from 1 to N, where N is the total number of observations across k groups.

Step 2: Add up the ranks for all the observations from sample i and call it r_i , where i can be any integer between 1 and k.

$$T = (N-1)\frac{\sum_{i=1}^{k} n_i (\bar{r}_i - \bar{r})^2}{\sum_{i=1}^{k} \sum_{j=1}^{n_i} (\bar{r}_{ij} - \bar{r})^2}$$

Where:

- *k* is the number of groups
- *n_i* is the sample size of sample *i*
- *N* is the total number of all the observations across *k* groups
- *r*_{*ij*} is the rank (among all the observations) of observation *j* from group *i*.

Step 4: Make a decision of whether to reject the null hypothesis.

- Null Hypothesis (H₀): $\eta_1 = \eta_2 = ... = \eta_k$
- Alternative Hypothesis (H_a): at least one of the medians is different from others.

The test statistic follows chi-square distribution when the null hypothesis is true. If T is greater than χ^2_{k-1} (the critical chi-square statistic), we reject the null and claim there is at least one median statistically different from other medians. If T is smaller than χ^2_{k-1} , we fail to reject the null and claim the medians of k groups are equal.

Use Minitab to Run a Kruskal-Wallis One-Way ANOVA

Case study: We are interested in comparing customer satisfaction among three types of customers using a nonparametric (i.e., distribution-free) hypothesis test: Kruskal–Wallis one-way ANOVA.

Data File: "Kruskal–Wallis" tab in "Sample Data.xlsx"

Customer Satisfaction Comparison



Fig. 3.78 Kruskal-Wallis Test

• Null Hypothesis (H₀): $\eta_1 = \eta_2 = \eta_3$

• Alternative Hypothesis (H_a): at least one of the customer types has different overall satisfaction levels from the others.

Steps to run a Kruskal-Wallis One-Way ANOVA in Minitab

- 1. Click Stat \rightarrow Nonparametrics \rightarrow Kruskal–Wallis.
- 2. A new window named "Kruskal–Wallis" pops up.
- 3. Select "Overall Satisfaction" as the "Response."
- 4. Select "Customer Type" as the "Factor."

Kruskal-Wallis	×
	Response: erall Satisfaction'
	Factor: 'Customer Type'
Select	
Help	OK Cancel

Fig. 3.79 Kruskal-Wallis dialog box with selected response and factor variables

5. Click "OK."

6. The Kruskal–Wallis test results appear in the session window.

(ruskal-	ruskal-Wallis Test: Overall Satisfaction versus Customer Type						
Descript	ive St	atistics					
Customer							
Туре	N	Median	Mean Rank	Z-Value			
1	31	3.56	36.0	-3.34			
2	42	4.34	65.9	4.53			
3	27	3.51	43.1	-1.56			
Overall	100		50.5				
Test Null hypo Alternative		-	: All medians : At least one		different		
Method		DF	H-Value P	-Value			
Not adjus	ted for	ties 2	21.36	0.000			
Adjusted f	for ties	2	21.36	0.000			

Fig. 3.80 Kruskal-Wallis Test output

The p-value of the test is lower than alpha level (0.05), and we reject the null hypothesis and conclude that at least the overall satisfaction median of one customer type is statistically different from the others.

3.4.3 Mood's Median

What is Mood's Median Test?

Mood's median test is a statistical test to compare the medians of two or more populations.

- Null Hypothesis (H₀): $\eta_1 = ... = \eta_k$
- Alternative Hypothesis (H_a): At least one of the medians is different from the others.

The symbol *k* is the number of groups of our interest and is equal to or greater than two. Mood's median is an alternative to Kruskal–Wallis. For the data with outliers, Mood's median test is more robust than the Kruskal–Wallis test. It is the extension of one sample sign test. In a one-sample sign test, the null hypothesis is that the probability of a random value from the population being above the specified value is equal to the probability of a random value being below the specified value.

Mood's Median Test Assumptions

- The sample data drawn from the populations of interest are unbiased and representative.
- The data of k populations are continuous or ordinal when the spacing between adjacent values is not constant.

- The k populations are independent to each other.
- The distributions of k populations have the same shape.
- Mood's median test is robust for non-normally distributed populations.
- Mood's median test is robust for data with outliers.

How Mood's Median Test Works

Step 1: Group the k samples from k populations (sample i is from population i) into one single data set and get the median of this combined data set. Step 2: Separate the data in each sample into two groups. One consists of all the observations with values higher than the grand median and the other consists of all the observations with values lower than the grand median.

Step 3: Run a Pearson's chi-square test to determine whether to reject the null hypothesis.

- Null Hypothesis (H₀): $\eta_1 = ... = \eta_k$
- Alternative Hypothesis (H_a): At least one of the medians is different from the others.
- If χ^2_{calc} is greater than χ^2_{crit} , we reject the null hypothesis and claim that at least one median is different from the others.
- If χ^2_{calc} is smaller than χ^2_{crit} , we fail to reject the null hypothesis and claim that the medians of k populations are not statistically different.

Use Minitab to Run a Mood's Median Test

Case study: We are interested in comparing customer satisfaction among three types of customers using a nonparametric (i.e., distribution-free) hypothesis test: Mood's median test.

_____ Data File: "Median Test" tab in "Sample Data.xlsx"

- Null Hypothesis (H₀): $\eta_1 = \eta_2 = \eta_3$
- Alternative Hypothesis (H_a): At least one customer type has different overall satisfaction from the others.

Steps to run a Mood's median test in Minitab

- 1. Click Stat \rightarrow Nonparametrics \rightarrow Mood's Median Test.
- 2. A new window named "Mood's Median Test" pops up.
- 3. Select "Overall Satisfaction" as the "Response."

4. Select "Customer Type" as the "Factor."

Mood's Median Test	x
	Response: erall Satisfaction'
	Factor: 'Customer Type'
	Store residuals
	🗖 Store fits
Select	
Help	OK Cancel

Fig. 3.81 Mood's Median Test with response and factor variables

- 5. Click "OK."
- 6. The Mood's median test results appear in the session window.

Mood's	Mood's Median Test: Overall Satisfaction versus Customer Type							
Descripti	ive Stati	stics						
Customer Type	Median	N <= Overall Median	N > Overall Median	Q3 – Q1	95% Median Cl			
1	3.560	21	10	1.2100	(2.95350, 3.93624)			
2	4.340	12	30	0.8975	(4.09464, 4.51841)			
3	3.510	17	10	0.9300	(3.28911, 4.02267)			
Overall	3.945							
Test Null hypot	thesis	H₀: The population (medians are all equal					
Alternative	Alternative hypothesis H1: The population medians are not all equal							
DF Chi-S 2		Value 0.001						

Fig. 3.82 Mood's Median Test output

The p-value of the test is lower than alpha level (0.05), and we reject the null hypothesis and conclude that at least the overall satisfaction median of one customer type is statistically different from the others.

3.4.4 Friedman

What is the Friedman Test?

The *Friedman test* is a hypothesis test used to detect the differences in the medians of various groups across multiple attempts. It is a non-parametric test developed by economist Milton Friedman.

- Null Hypothesis (H₀): The treatments have identical effects (i.e., $\eta_1 = ... = \eta_k$).
- Alternative Hypothesis (H_a): At least one of the treatments has different effects from the others (i.e., at least one of the medians is statistically different from others).

It is used as an alternative of the parametric repeated measures ANOVA when the assumption of normality or variance equality is not met.

Friedman Test Assumptions

- The sample data drawn from the populations of interest are unbiased and representative.
- The data are continuous or ordinal when the spacing between adjacent values is not constant.
- Results in one block are independent of the results in another.
- The Friedman test is robust for the non-normally distributed population.
- The Friedman test is robust for populations with unequal variances.

How Friedman Test Works

Step 1: Organize the data into a tabular view with n rows indicating the blocks and k columns indicating the treatments. Each observation x_{ij} is filled into the intersection of a specific block i and a specific treatment j.

Step 2: Calculate the ranks of each observation within each block.

Step 3: Replace the values in the table created in step 1 with the order r_{ij} (within a block) of the corresponding observation x_{ij} .

Step 4: Calculate the test statistic

$$Q = \frac{n \sum_{j=1}^{k} (\bar{r}_{j} - \bar{r})^{2}}{\frac{1}{n(k-1)} \sum_{i=1}^{n} \sum_{j=1}^{k} (r_{ij} - \bar{r})^{2}}$$

Where:

$$\bar{r}_j = \frac{1}{n} \sum_{i=1}^n r_{ij}$$

$$\overline{r} = \frac{1}{nk} \sum_{i=1}^{n} \sum_{j=1}^{k} r_{ij}$$

Step 5: Make a decision of whether to reject the null hypothesis.

When n > 15 or k > 4, the test statistic Q follows a chi-square distribution if the null hypothesis is true and the p-value is $P(\chi^2_{k-1} \ge Q)$.

When n < 15 and k < 4, the test statistic Q does not approximate a chi-square distribution and the p-value can be obtained from tables of Q for the Friedman test. If p-value > alpha level (0.05), we fail to reject the null.

Friedman Test Examples

A number of n water testers judge the quality of k different water samples, each of which is from a distinct water source. We will apply a Friedman test to determine whether the water qualities of the k sources are the same. There are n blocks and k treatments in this experiment. One tester's decision would not have any influence on other testers. When running the experiment, each tester judges the water in a random sequence.

Use Minitab to Run a Friedman Test

Case study: We are interested in comparing the effect of a drug treatment on enzyme activity. Three different drug therapies were given to four animals and each animal belongs to a different litter. (*Note: we will be using data from a difference source than we have been to this point.*)

Data File: "Friedman.MTW"

- Null Hypothesis (H₀): η₁ = η₂ = η₃
- Alternative Hypothesis (H_a): At least one treatment effect is statistically different from the others.

Steps to run a Friedman test in Minitab:

- 1. Click Stat \rightarrow Nonparametrics \rightarrow Friedman.
- 2. A new window named "Friedman" pops up.
- 3. Select "EnzymeActivity" as the "Response."
- 4. Select "Therapy" as the "Treatment."

5. Select "Litter" as the "Blocks."

Friedman		×
	Response:	EnzymeActivity
	Treatment:	Therapy
	Blocks:	Litter
	Store re	
Select Help	ОК	Cancel

Fig. 3.83 Friedman dialog box with response, treatment, and blocks variables selected

- 6. Click "OK."
- 7. The Friedman test results appear in the session window.

Friedma	Friedman Test: EnzymeActivity vs Therapy, Litter							
Method								
Treatmen Block = L		herapy						
Descript	tive	Statistic	s					
Therapy	Ν	Mediar	I	Sum of Ranks	_			
1	4	0.245000)	6.5	-			
2	4	0.311667	,	7.0				
3	4	0.578333		10.5				
Overall	12	0.378333						
Test								
Null hypo	thesi	is	H ₀	: All treatment	effects are	e zero		
Alternativ	Alternative hypothesis H_1 : Not all treatment effects are zero							
Method		D	F	Chi-Square	P-Value			
Not adjus	Not adjusted for ties 2 2.38 0.305							
Adjusted	for ti	es	2	3.80	0.150			

Fig. 3.84 Friedman Test output

In this example, the p-value is 0.305, which is obviously greater than 0.05. Therefore, we fail to reject the null hypothesis and we conclude that there is not a statistically significant difference between treatments.

3.4.5 ONE SAMPLE SIGN

What is the One Sample Sign Test?

The *one sample sign test* is a hypothesis test to compare the median of a population with a specified value. However, with this test we are comparing the median of the population instead of the mean.

- Null Hypothesis (H₀): $\eta = \eta_0$
- Alternative Hypothesis (H_a): $\eta \neq \eta_0$

It is an alternative test of one sample t-test when the distribution of the data is non-normal. It is robust for the data with non-symmetric distribution.

One Sample Sign Test Assumptions

- The sample data drawn from the population of interest are unbiased and representative.
- The data are continuous or ordinal when the spacing between adjacent values is not constant.
- The one sample sign test is robust for the non-normally distributed population.
- The one sample sign test does not have any assumptions on the distribution. It is a distribution-free test.

Assumptions for this test are very similar to those of other non-parametrics; however, a key difference is that there are no assumptions about the shape of the distribution.

How the One Sample Sign Test Works

Step 1: Separate the sample set of data into two groups: one with values greater than and the other with values less than the hypothesized median η_0 . Count the number of observations in each group.

Step 2: Calculate the test statistic.

If the null hypothesis is true, the number of observations in each group should not be significantly different from half of the total sample size. The test statistic follows a binomial distribution when the null is true. When n is large, we use the normal distribution to approximate the binomial distribution.

$$Z_{calc} = \frac{n_{+} - np}{\sqrt{np(1-p)}}$$

Where:

- n_{+} is the number of observations with values greater than the hypothesized median
- *n* is the total number of observations
- *p* is 0.5.

Step 3: Make a decision on whether to reject the null hypothesis. If the $|Z_{calc}|$ is smaller than Z_{crit} , we fail to reject the null hypothesis and claim that there is no significant difference between the population median and the hypothesized median.

Use Minitab to Run a One Sample Sign Test

Case study: We are interested in comparing the overall satisfaction of customer type 1 against a specified benchmark satisfaction (3.5) using a nonparametric (i.e., distribution-free) hypothesis test: one sample sign test.



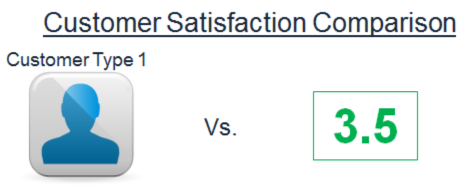


Fig. 3.85 One Sample Sign Test

- Null Hypothesis (H₀): $\eta_1 = 3.5$
- Alternative Hypothesis (H_a): $\eta_1 \neq 3.5$

Steps to run a one sample sign test in Minitab:

- 1. Click Stat \rightarrow Nonparametrics \rightarrow 1-Sample Sign.
- 2. A new window named "1-Sample Sign" pops up.
- 3. Select "Overall Satisfaction" as the "Variables."
- 4. Click the box of "Test median."

5. Enter "3.5" in the box next to "Test median."

1-Sample Sign	×
	Variables: 'Overall Satisfaction'
	 Confidence interval Level: 95.0 Test median: 3.5 Alternative: not equal ▼
Select	OK Cancel

Fig. 3.86 1-Sample Sign dialog box with selected variables

- 6. Click "OK."
- 7. The one sample sign test results appear in session window.

Sign Test for Median: Overall Satisfaction					
Method					
η: median of Overall Satisfaction					
Descriptive Statisti	Descriptive Statistics				
Sample N	N Median				
Overall Satisfaction 3	1 3.56				
Test					
Null hypothesis	H₀: η = 3.5				
Alternative hypothesis	H₁: η ≠ 3.5				
Sample N	umber < 3.5	Number = 3.5	Number > 3.5	P-Value	
Overall Satisfaction	15	0	16	1.000	

Fig. 3.87 1-Sample Sign output

The p-value of the one sample sign test is 1.0000, higher than the alpha level (0.05), and we fail to reject the null hypothesis. There is not any statistically significant difference between the overall satisfaction of customer type 1 and the benchmark satisfaction level.

3.4.6 ONE SAMPLE WILCOXON

What is the One Sample Wilcoxon Test?

The *one sample Wilcoxon test* is a hypothesis test to compare the median of one population with a specified value.

- Null Hypothesis (H₀): $\eta = \eta_0$
- Alternative Hypothesis (H_a): $\eta \neq \eta_0$

It is an alternative test of one sample t-test when the distribution of the data is non-normal. It is more powerful than one sample sign test but it assumes the distribution of the data is symmetric.

One Sample Wilcoxon Test Assumptions

- The sample data drawn from the population of interest are unbiased and representative.
- The data are continuous or ordinal when the spacing between adjacent values is not constant.
- The distribution of the data is symmetric about a median.
- The one sample Wilcoxon test is robust for the non-normally distributed population.

The assumptions are similar to other non-parametric tests, again the difference about the symmetry of the distribution.

How the One Sample Wilcoxon Test Works

Step 1: Create the following columns one by one:

- Column 1: All the raw observations (X)
- Column 2: The differences between each observation value and the hypothesized median $(X \eta_0)$
- Column 3: The signs (+ or –) of column 2
- Column 4: The absolute value of column 2
- Column 5: The ranks of each item in column 4 in ascending order
- Column 6: The product of column 3 and column 5

Step 2: Calculate the test statistic W_{calc} , which is the sum of all the non-negative values in column 6.

Step 3: Make a decision on whether to reject the null hypothesis. Use the table of critical values for the Wilcoxon test to get the W_{crit} with predetermined alpha level and number of observations.

If the W_{calc} is smaller than the W_{crit} , we fail to reject the null hypothesis and claim that there is no significant difference between the population median and the hypothesized median. To calculate the W statistic, sum all of the non-negative values in column 6. To assess the hypothesis, the W statistic is compared against the critical W value.

Use Minitab to Run a One Sample Wilcoxon Test

Case study: We are interested in comparing the overall satisfaction of customer type 1 against a specified benchmark satisfaction (3.5) using a nonparametric (i.e., distribution-free) hypothesis test: one sample Wilcoxon test.



Fig. 3.88 One Sample Wilcoxon

- Null Hypothesis (H₀): $\eta_1 = 3.5$
- Alternative Hypothesis (H_a): $\eta_1 \neq 3.5$

Steps to run a one sample Wilcoxon test in Minitab:

- 1. Click Stat \rightarrow Nonparametrics \rightarrow 1-Sample Wilcoxon.
- 2. A new window named "1-Sample Wilcoxon" pops up.
- 3. Select "Overall Satisfaction" as the "Variables."
- 4. Click the radio button "Test median."
- 5. Enter "3.5" in the box next to "Test median."

1-Sample Wilcoxon	
	Variables:
	'Overall Satisfaction'
	C Confidence interval Level: 95.0
	Test median: 3.5 Alternative: not equal
Select	
Help	OK Cancel

Fig. 3.89 1-Sample Wilcoxon dialog box with selected variables

- 6. Click "OK."
- 7. The one sample Wilcoxon test results appear in the session window.

Wilcoxon Signed Rank Test: Overall Satisfaction				
Method				
η: median of Overall	Satisfac	tion		
Descriptive Statistics				
Sample	N N	1edian		
Overall Satisfaction	31 3	3.4075		
Test				
Null hypothesis	H₀:	η = 3.5		
Alternative hypothesis $H_1: \eta \neq 3.5$				
	N for	Wilcoxon		
Sample	Test	Statistic	P-Value	
Overall Satisfaction	31	217.50	0.557	

Fig. 3.90 1-Sample Wilcoxon output

The p-value of the one sample Wilcoxon test is 0.557, higher than the alpha level (0.05), and we fail to reject the null hypothesis. There is not any statistically significant difference between the overall satisfaction of customer type 1 and the benchmark satisfaction level.

3.4.7 One and Two Sample Proportion

What is the One Sample Proportion Test?

One sample proportion test is a hypothesis test to compare the proportion of one certain outcome (e.g., the number of successes per the number of trials, or the number of defects per the total number of opportunities) occurring in a population following the binomial distribution with a specified proportion.

- Null Hypothesis (H₀): *p* = *p*₀
- Alternative Hypothesis (H_a): $p \neq p_0$

One Sample Proportion Test Assumptions

• The sample data drawn from the population of interest are unbiased and representative.

- There are only two possible outcomes in each trial: success/failure, yes/no, and defective/non-defective etc.
- The underlying distribution of the population is binomial distribution.
- When np ≥ 5 and np(1 p) ≥ 5, the binomial distribution can be approximated by the normal distribution.

How the One Sample Proportion Test Works

When $np \ge 5$ and $np(1 - p) \ge 5$, we use normal distribution to approximate the underlying binomial distribution of the population.

Test Statistic

$$Z_{calc} = \frac{\hat{p} - p_0}{\sqrt{\frac{p_0(1 - p_0)}{n}}}$$

Where:

- \hat{p} is the observed probability of one certain outcome occurring?
- p₀ is the hypothesized probability
- *n* is the number of trials.

When $|Z_{calc}|$ (absolute value of the calculated test statistic) is smaller than Z_{crit} (critical value), we fail to reject the null hypothesis and claim that there is no statistically significant difference between the population proportion and the hypothesized proportion.

Use Minitab to Run a One Sample Proportion Test

Case study: We are interested in comparing the exam pass rate of a high school this month against a specified rate (70%) using a nonparametric (i.e., distribution-free) hypothesis test: one sample proportion test.



Data File: "One Sample Proportion" tab in "Sample Data.xlsx"

- Null Hypothesis (H₀): p = 70%
- Alternative Hypothesis (H_a): $p \neq 70\%$

Steps to run a one sample proportion test in Minitab:

- 1. Click Stat \rightarrow Basic Statistics \rightarrow 1 Proportion.
- 2. A new window named "One-Sample Proportion" appears.
- 3. Click on the drop-down menu and choose "Summarized data."
- 4. Enter "77" in the box of "Number of events."
- 5. Enter "105" in the box of "Number of trials."
- 6. Check the box "Perform hypothesis test."

7. Enter "0.70" as the "Hypothesized proportion."

One-Sample Proportion	
	Summarized data
	Number of events: 77
1	Number of trials: 105
	-
	Perform hypothesis test <u>Hypothesized proportion</u> : 0.70
Select	Optio <u>n</u> s
Help	<u>O</u> K Cancel

Fig. 3.91 One-Sample Proportion dialog box with hypothesis test choice

- 8. Click "OK."
- 9. The one sample proportion test results appear in the session window.

Test and CI for One Proportion					
Method					
p: event proportion Exact method is used for this analysis.					
Descriptive Statistics					
Ν	Event	Sample p	95% CI for p		
105	77	0.733333	(0.638143, 0.814925)		
Test					
Null hypothesis			$H_0: p = 0.7$		
Alternative hypothesis $H_1: p \neq 0.7$					
P-Val	ue				
0.4	60				

Fig. 3.92 One-Sample Proportion output

The p-value of the one sample proportion test is 0.460, greater than the alpha level (0.05), and we fail to reject the null hypothesis. We conclude that the exam pass rate of the high school this month is not statistically different from 70%.

What is the Two Sample Proportion Test?

The *two-sample proportion test* is a hypothesis test to compare the proportions of one certain event occurring in two populations following the binomial distribution.

- Null Hypothesis (H₀): p₁ = p₂
- Alternative Hypothesis (H_a): $p_1 \neq p_2$

Two-Sample Proportion Test Assumptions

- The sample data drawn from the populations of interest are unbiased and representative.
- There are only two possible outcomes in each trial for both populations: success/failure, yes/no, and defective/non-defective etc.
- The underlying distributions of both populations are binomial distribution.
- When np ≥ 5 and np(1 p) ≥ 5, the binomial distribution can be approximated by the normal distribution.

How the Two-Sample Proportion Test Works

When $np \ge 5$ and $np(1 - p) \ge 5$, we use normal distribution to approximate the underlying binomial distributions of the populations.

Test Statistic

$$Z = \frac{\hat{p}_1 - \hat{p}_2}{\sqrt{\hat{p}_0 \left(1 - \hat{p}_0\right) \left(\frac{1}{n_1} + \frac{1}{n_2}\right)}}$$

Where:

$$\hat{p}_0 = \frac{x_1 + x_2}{n_1 + n_2}$$

and where:

- \hat{p}_1 and \hat{p}_2 are the observed proportions of events in the two samples
- n_1 and n_2 is the number of trials in the two samples respectively
- X_1 and X_2 is the number of events in the two samples respectively.

When $|Z_{calc}|$ is smaller than Z_{crit} , we fail to reject the null hypothesis.

Use Minitab to Run a Two-Sample Proportion Test

Case study: We are interested in comparing the exam pass rates of a high school in March and April using a nonparametric (i.e., distribution-free) hypothesis test: two sample proportion test.

Data File: "Two-Sample Proportion" tab in "Sample Data.xlsx"

- Null Hypothesis (H₀): p_{March} = p_{April}
- Alternative Hypothesis (H_a): p_{March} ≠ p_{April}

Steps to run a two-sample proportion test in Minitab:

- 1. Click Stat \rightarrow Basic Statistics \rightarrow 2 Proportions.
- 2. A new window named "Two-Sample Proportion" pops up.
- 3. Click the drop-down menu and choose "Summarized data."
- 4. Enter "89" in the box intersecting "First" and "Events."
- 5. Enter "112" in the box intersecting "First" and "Trials."
- 6. Enter "102" in the box intersecting "Second" and "Events."
- 7. Enter "130" in the box intersecting "Second" and "Trials."

Two-Sample Proportion			×
	Summarized data		_
	Number of events:	Sample 1 89	Sample 2 102
	Number of trials:	112	130
Select			Optio <u>n</u> s
Help		<u>0</u> K	Cancel

Fig. 3.93 Two-Sample Proportion dialog box with data entered 8. Click "OK."

9. The two-sample proportion test results appear in the session window.

est and CI for T	wo Proportior						
Method							
p_1 : proportion where S	•						
p ₂ : proportion where S Difference: p ₁ - p ₂	ample 2 = Event						
Descriptive Statist	ics						
Sample N Even	t Sample p						
Sample 1 112 89	0.794643						
Sample 2 130 102	0.784615						
95% CI for Difference Difference							
Difference Diffe							
Difference Diffe	erence 4, 0.112939)						
DifferenceDiffe0.0100275(-0.092884CI based on normal approx	erence 4, 0.112939)						
Difference Diffe 0.0100275 (-0.092884 CI based on normal approx Test	erence 4, 0.112939)						
Difference Diffe 0.0100275 (-0.092884 <i>CI based on normal approx</i> Test Null hypothesis	erence 4, 0.112939) <i>kimation</i> H ₀ : p ₁ - p ₂ = 0						
Difference Diffe 0.0100275 (-0.092884 CI based on normal approx Test	erence 4, 0.112939) <i>kimation</i> H ₀ : p ₁ - p ₂ = 0						
Difference Diffe 0.0100275 (-0.092884 <i>CI based on normal approx</i> Test Null hypothesis	erence 4, 0.112939) <i>kimation</i> H ₀ : p ₁ - p ₂ = 0						
DifferenceDiffe0.0100275(-0.092884CI based on normal approxTestNull hypothesisAlternative hypothesis	erence 4, 0.112939) kimation $H_0: p_1 - p_2 = 0$ $H_1: p_1 - p_2 \neq 0$ Z-Value P-Value						

Fig. 3.94 Two-Sample Proportion Test output

The p-value of the two-sample proportion test is 0.849, greater than the alpha level (0.05), and we fail to reject the null hypothesis. We conclude that the exam pass rates of the high school in March and April are not statistically different.

3.4.8 CHI-SQUARED (CONTINGENCY TABLES)

What is the Chi-Square Test?

We have looked at hypothesis tests to analyze the proportion of one population vs. a specified value, and the proportions of two populations, but what do we do if we want to analyze more than two populations? A *chi-square test* is a hypothesis test in which the sampling distribution of the test statistic follows a chi-square distribution when the null hypothesis is true. There are

multiple chi-square tests available and in this module, we will cover the Pearson's chi-square test used in contingency analysis.

- Null Hypothesis (H₀): p₁ = p₂ = ... = p_k
- Alternative Hypothesis (H_a): At least one of the proportions is different from others.

The symbol *k* is the number of populations of our interest; $k \ge 2$.

What is the Chi-Square Test?

The chi-square test can also be used to test whether two factors are independent of each other. In other words, it can be used to test whether there is any statistically significant relationship between two discrete factors.

- Null Hypothesis (H₀): Factor 1 is independent of factor 2.
- Alternative Hypothesis (H_a): Factor 1 is not independent of factor 2.

Chi-Square Test Assumptions

- The sample data drawn from the populations of interest are unbiased and representative.
- There are only two possible outcomes in each trial for an individual population: success/failure, yes/no, and defective/non-defective etc.
- The underlying distribution of each population is binomial distribution.
- When np ≥ 5 and np(1 p) ≥ 5, the binomial distribution can be approximated by the normal distribution.

How Chi-Square Test Works

Test Statistic

$$\chi^{2}_{calc} = \sum_{i=1}^{n} \frac{(O_{i} - E_{i})^{2}}{E_{i}}$$

Where:

- *O_i* is an observed frequency
- *E_i* is an expected frequency
- *N* is the number of cells in the contingency table.

If χ^2_{calc} (calculated chi-square statistic) is smaller than χ^2_{crit} (critical value), we fail to reject the null hypothesis. The test statistic is calculated with the observed and expected frequency.

Use Minitab to Run a Chi-Square Test

Case study 1: We are interested in comparing the product quality exam pass rates of three suppliers A, B, and C using a nonparametric (i.e., distribution-free) hypothesis test: chi-square test.



- Null Hypothesis (H₀): p_A = p_B = p_C
- Alternative Hypothesis (H_a): At least one of the suppliers has different pass rates from the others.

Use Minitab to Run a Chi-Square Test:

Steps to run a chi-square test in Minitab:

- 1. Click Stat \rightarrow Tables \rightarrow Cross Tabulation and Chi-Square.
- 2. A new window named "Cross Tabulation and Chi-Square" pops up.
- 3. Select "Results" as "For rows."
- 4. Select "Supplier" as "For columns."
- 5. Select "Count" as "Frequencies."

Cross Tabulation and Chi	-Square
C3 Count	Raw data (categorical variables)
	Results
	Columns: Supplier
	Layers:
	Erequencies: Count (optional)
	Display
	Column percents
	Total percents
Select	Chi-Square Other <u>S</u> tats Options
Help	QK Cancel

Fig. 3.95 Cross Tabulation and Chi-Square with selected variables

- 6. Click the "Chi-Square" button.
- 7. A new window named "Cross Tabulation Chi-Square" pops up.
- 8. Check the boxes of "Chi-square analysis", "Expected cell counts," and "Each cell's contribution to the Chi-Square statistic."

Cross Tabulation: Chi-Square	-	×
Chi-square test		
Statistics to display in each cell		
Expected cell counts		
Raw residuals		
Standardized residuals		
Adjusted residuals		
Each cell's contribution to chi-squar	e	
Help	<u>o</u> ĸ	Cancel

Fig. 3.96 Cross Tabulation: Chi-Square dialog box with proper selections

9. Click "OK" in the window named "Cross Tabulation – Chi-Square."

10. Click "OK" in the window named "Cross Tabulation and Chi-Square."

11. The Chi-square test results appear in the session window.

Tabula	Tabulated Statistics: Results, Supplier									
Using fre	Using frequencies in Count									
Rows:	Rows: Results Columns: Supplier									
	Supplier A Supplier B Supplier C All									
Fail	2 21.1 0.065	-	3 20.0 5.000		10 18.82 4.1360	60				
Pass	16 158.8 0.008	2	14 150.0 0.666	0	150 141.18 0.5515	450				
All	18	0	17	0	160	510				
Cell Contents Count Expected count Contribution to Chi-square										
Chi-Square Test										
Chi-Square DF P-Value										
	Pearson	10.428	-	2						
Likeliho	od Ratio	10.622	2	2	0.005					

Fig. 3.97 Cross Tabulation and Chi-Square output

Counts are based on the sample observation. Expected counts are based on the assumption that the null hypothesis is true. Since the p-value is smaller than alpha level (0.05), we reject the null hypothesis and claim that at least one supplier has different pass rate from others.

Case study 2: We are trying to check whether there is a relationship between the suppliers and the results of the product quality exam using nonparametric (i.e., distribution-free) hypothesis test: chi-square test.



Data File: "Chi-Square Test2" tab in "Sample Data.xlsx"

- Null Hypothesis (H₀): Product quality exam results are independent of the suppliers.
- Alternative Hypothesis (H_a): Product quality exam results depend on the suppliers.

Use Minitab to Run a Chi-Square Test:

Steps to run a chi-square test in Minitab:

- 1. Click Stat \rightarrow Tables \rightarrow Cross Tabulation and Chi-Square.
- 2. A new window named "Cross Tabulation and Chi-Square" pops up.
- 3. Select "Results" as "For rows."
- 4. Select "Supplier" as "For columns."
- 5. Select "Count" as "Frequencies are in."

Cross Tabulation and Chi	-Square
C3 Count	Raw data (categorical variables)
	Results
	Columns: Supplier
	Layers:
	Erequencies: Count (optional)
	Display
	Counts
	Column percents
	Total percents
Select	Chi-Square Other <u>S</u> tats Options
Help	<u>Q</u> K Cancel

Fig. 3.98 Cross Tabulation and Chi-Square dialog box with selected variables

- 6. Click the "Chi-Square" button.
- 7. Check the boxes of "Chi-square analysis", "Expected cell counts," and "Each cell's contribution to the Chi-Square statistic."

Cross Tabulation: Chi-Square		×
☑ Chi-square test		
Statistics to display in each cell		
Expected cell counts		
Raw residuals		
Standardized residuals		
Adjusted residuals		
Each cell's contribution to chi-squar	e	
Help	<u>O</u> K	Cancel

Fig. 3.99 Cross Tabulation: Chi-Square dialog box with selections

- 8. Click "OK" in the window named "Cross Tabulation Chi-Square."
- 9. Click "OK" in the window named "Cross Tabulation and Chi-Square."
- 10. The Chi-square test results appear in the session window.

Tabulated Statistics: Results, Supplier

Using frequencies in Count

Rows: Results Columns: Supplier

	Supplier A	Supplier B	Supplier (C All
Fail	20	30	1	0 60
	20	20	20	D
	0.000	5.000	5.00	D
Marginal	20	30	4	0 90
	30	30	30	D
	3.333	0.000	3.33	3
Pass	160	140	15	0 450
	150	150	15	D
	0.667	0.667	0.00	D
All	200	200	20	0 600
Chi-Squ	iare Test	t		
	(Chi-Square	DF P-	Value
F	Pearson ?	18.000	4 0.0	001
Likelihoo	Likelihood Ratio		4 0.0	001

Fig. 3.100 Cross Tabulation and Chi-Square Test output

The p-value is smaller than the alpha level (0.05) and we reject the null hypothesis. The product quality exam results are not independent of the suppliers. These results indicate the danger that we can get into when using discrete data. Not everything is as simple as yes/no or pass/fail. Even though supplier C has a lower fail rate of 10, you can see that the number of marginal results is higher. However, the p-value tells us that we must reject the null hypothesis and claim that the quality exam results *are* dependent on the suppliers.

3.4.9 TESTS OF EQUAL VARIANCE

What are Tests of Equal Variance?

Tests of equal variance are a family of hypothesis tests used to check whether there is a statistically significant difference between the variances of two or more populations.

- Null Hypothesis (H₀): $\sigma_1^2 = \sigma_2^2 = ... = \sigma k^2$
- Alternative Hypothesis (H_a): At least the variance of one population is different from others.

k is the number of populations of interest; $k \ge 2$.

A test of equal variance can be used alone but most of the time it is used with other statistical methods to verify or support the assumption about the variance equality.

F-Test

The *F-test* is used to compare the variances between two normally distributed populations. It is extremely sensitive to non-normality and serves as a preliminary step for two sample t-test. Therefore, it is crucial to validate that the data is normal before using this test.

Test Statistic

$$F_{calc} = \frac{{S_1}^2}{{S_2}^2}$$

Where:

- s1 and s2 are the sample standard deviations
- The sampling distribution of the test statistic follows F distribution when the null is true.

Bartlett's Test

Bartlett's test is used to compare the variances among two or more normally distributed populations. It is sensitive to non-normality and it serves as a preliminary step for ANOVA.

Test Statistic

$$\chi^{2} = \frac{(N-k)\ln(S_{p}^{2}) - \sum_{i=1}^{k} (n_{i}-1)\ln(S_{i}^{2})}{1 + \frac{1}{3(k-1)} \left(\sum_{i=1}^{k} \left(\frac{1}{n_{i}-1}\right) - \frac{1}{N-k}\right)}$$

Where:

$$N = \sum_{i=1}^{k} n_i$$
 and $S_p^2 = \frac{1}{N-k} \sum_{i} (n_i - 1) S_i^2$

The sampling distribution of test statistic follows chi square distribution when the null is true.

Brown-Forsythe Test

The *Brown–Forsythe test* is used to compare the variances between two or more populations with any distributions. It is not as sensitive to non-normality as Bartlett's test. The test statistic is the model F statistic from the ANOVA on the transformed response:

$$z_{ij} = \left| y_{ij} - \widetilde{y}_i \right|$$

Where: \tilde{y}_i is the median response at ith level.

Levene's Test

Levene's test is used to compare the variances between two or more populations with any distributions. It is not as sensitive to non-normality as Bartlett's test. The test statistic is the model F statistic from the ANOVA on the transformed response:

$$z_{ij} = \left| y_{ij} - \overline{y}_i \right|$$

Where: \overline{y}_i is the mean response at ith level.

Brown-Forsythe Test vs. Levene's Test

$$F = \frac{(N-k)}{(k-1)} \frac{\sum_{i=1}^{k} N_i (Z_{i.} - Z_{..})^2}{\sum_{i=1}^{k} \sum_{j=1}^{N_i} (Z_{ij} - Z_{i.})^2}$$

Where:

- *N* is the total number of observations.
- *k* is the number of groups.

- *N_i* is the number of observations in the ith group
- *Z_i* is the group mean of the ith group
- *Z*. is the grand mean of all the observations.

The F statistic for both the Brown–Forsythe and Levene's test are calculated the same way; however, you can see at the bottom of the slide how Z represents a different value for each.

In Brown–Forsythe test, $Z_{ij} = |Y_{ij} - \widetilde{Y}_{ij}|$, where \widetilde{Y}_{ij} is the group median of the ith group. In Levene's test, $Z_{ij} = |Y_{ij} - \overline{Y}_{ij}|$, where \overline{Y}_{ij} is the group mean of the ith group.

Use Minitab to Run Tests of Equal Variance

Case study: We are interested in comparing the variances of the retail price of a product in state A and state B.

Data File: "Two-Sample T-Test" tab in "Sample Data.xlsx"

- Null Hypothesis (H₀): $\sigma_A^2 = \sigma_B^2$
- Alternative Hypothesis (H_a): $\sigma_A^2 = \sigma_B^2$

Step 1: Run the normality test to check whether all levels of data are normally distributed

- 1. Click Stat \rightarrow Basic Statistics \rightarrow Graphical Summary.
- 2. A new window named "Graphical Summary" pops up.
- 3. Select "Retail Price" as the "Variables."
- 4. Click in the blank box right below "By variables" and the "State" appears in the list box on the left.

5. Select "State" as "By variables."

Graphical Summary	X
C1 State	Variables:
C2 Retail Price	'Retail Price'
	τ
	By variables (optional):
	State 🔺
	-
	95.0
Select	
Help	OK Cancel

Fig. 3.101 Graphical Summary dialog box with selected variable

- 6. Click "OK."
- 7. The normality test results appear in the new windows.

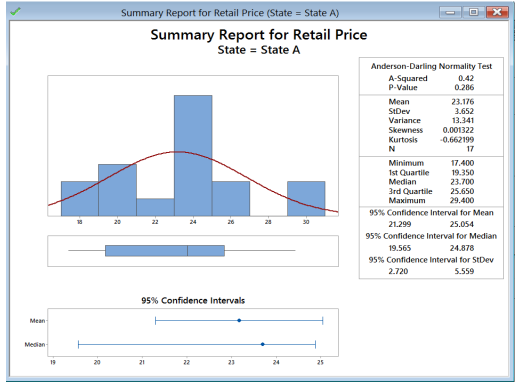


Fig. 3.102 Graphical Summary output for State A

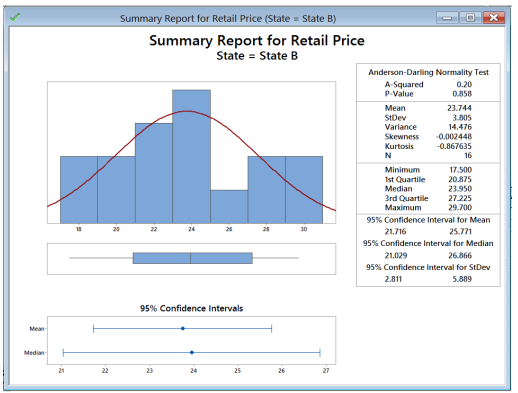


Fig. 3.103 Graphical Summary output for State B

- Null Hypothesis (H₀): The data are normally distributed.
- Alternative Hypothesis (H_a): The data are not normally distributed.

Both retail price data of state A and B are normally distributed since the p-values are both greater than alpha level (0.05). When the p-value is greater than the alpha value, we fail to reject the null hypothesis and claim that the distributions are normal. Both samples from state A and state B are normally distributed.

Step 2: Run tests of equal variance in Minitab

- 1. Click Stat \rightarrow ANOVA \rightarrow Test for Equal Variances.
- 2. A new window named "Test for Equal Variances" pops up.
- 3. Select "Retail Price" as the "Response."

4. Select "State" as "Factors."

Test f	for Equal Variances		x
C1 C2	State Retail Price	Response data are in one column for all factor levels <u>Response</u> : 'Retail Price' Factors:	•
		State	*
		Options	is
	Select	R <u>e</u> sults <u>S</u> torag	je
	Help	<u>O</u> K Can	cel

Fig. 3.104 Test for Equal Variances dialog box with selected variables

- 5. Click "Options"
- 6. Check "Use test based on normal distribution"

Test for Equal Variances: Options		×
Confidence level: 95.0		
☑ Use test based on normal distribut	tion	
Help	<u>о</u> к	Cancel

Fig. 3.105 Test for Equal Variances: Options dialog box

- 7. Click "OK" to close the Options window
- 8. Click "OK."

9. The results of variances equality tests appear in the new window.

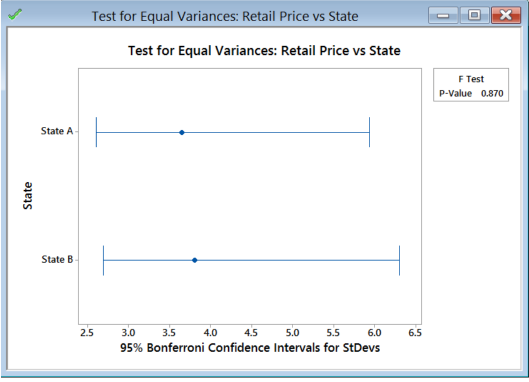


Fig. 3.106 Test for Equal Variances output

If all the groups of data are normally distributed, use the F-test or Bartlett's test in Minitab to test the equality of the variances. If at least one of the groups is not normally distributed, use Levene's test in Minitab to test the equality of the variances. If the p-value of the variances equality test is greater than the alpha level (0.05), we fail to reject the null hypothesis and conclude that the variances of different groups are identical.

Hypothesis Testing Roadmap: Putting it all together

Hypothesis Testing Roadmap

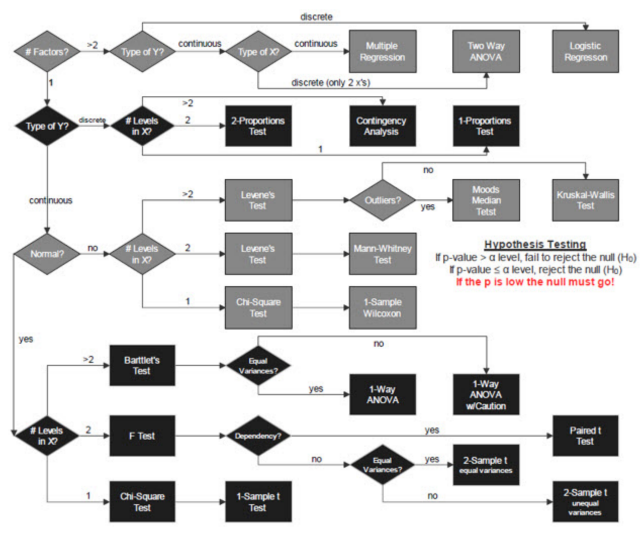


Fig. 3.107 Hypothesis Testing Roadmap

The roadmap is a great way to quickly decide what type of hypothesis test fits a given situation by evaluating the type of data (discrete vs. continuous), the normality of the data, the number of levels, and other factors.

4.0 IMPROVE PHASE

4.1 SIMPLE LINEAR REGRESSION

After identifying and validating the critical Xs during the Analyze phase, we will use some statistical tools during the Improve phase to define the settings needed to achieve the desired output. One such tool is simple linear regression.

4.1.1 CORRELATION

We will start with correlation before regression, because it is a tool that helps us understand whether a relationship exists between two variables without having to know causality (which variable causes a response in the other). There are many similarities between correlation and simple linear regression.

What is Correlation?

Correlation is a statistical technique that describes whether and how strongly two or more variables are related.

Correlation analysis helps to understand the direction and degree of association between variables, and it suggests whether one variable can be used to predict another. Of the different metrics to measure correlation, *Pearson's correlation coefficient* is the most popular. It measures the linear relationship between two variables.

Pearson's Correlation Coefficient

Pearson's correlation coefficient is also called Pearson's r or coefficient of correlation and Pearson's product moment correlation coefficient (r), where r is a statistic measuring the linear relationship between two variables.

Correlation coefficients range from -1 to 1.

• If r = 0, there is no linear relationship between the variables.

The *sign* of r indicates the *direction* of the relationship:

- If r < 0, there is a negative linear correlation.
- If r > 0, there is a positive linear correlation.

The *absolute value* of r describes the *strength* of the relationship:

- If $|r| \le 0.5$, there is a weak linear correlation.
- If |r| > 0.5, there is a strong linear correlation.
- If |r| = 1, there is a perfect linear correlation.

When the correlation is *strong*, the data points on a scatter plot will be close together (tight). The closer r is to -1 or 1, the stronger the relationship.

- -1 Strong inverse relationship
- +1 Strong direct relationship

When the correlation is *weak*, the data points are spread apart more (loose). The closer the correlation is to 0, the weaker the relationship.

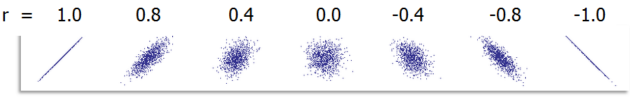


Fig. 4.1 Examples of Types of Correlation

This Figure above demonstrates the relationships between variables as the Pearson r value ranges from 1 to 0 and to -1. Notice that at -1 and 1 the points form a perfectly straight line.

- At 0 the data points are completely random.
- At 0.8 and –0.8, notice how you can see a directional relationship, but there is some noise around where a line would be.
- At 0.4 and -0.4, it looks like the scattering of data points is leaning to one direction or the other, but it is more difficult to see a relationship because of all the noise.

Pearson's correlation coefficient is only sensitive to the *linear* dependence between two variables. It is possible that two variables have a perfect non-linear relationship when the correlation coefficient is low. Notice the scatter plots below with correlation equal to 0. There are clearly *relationships* but they are not linear and therefore cannot be determined with Pearson's correlation coefficient.

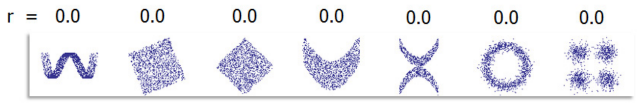


Fig. 4.2 Examples of Types of Relationships

Correlation and Causation

Correlation *does not* imply causation.

If variable A is highly correlated with variable B, it does not necessarily mean A causes B or vice versa. It is possible that an unknown third variable C is causing both A and B to change. For example, if ice cream sales at the beach are highly correlated with the number of shark

attacks, it does not imply that increased ice cream sales causes increased shark attacks. They are triggered by a third factor: summer.

This example demonstrates a common mistake that people make: assuming causation when they see correlation. In this example, it is hot weather that is a common factor. As the weather is hotter, more people consume ice cream *and* more people swim in the ocean, making them susceptible to shark attacks.

Correlation and Dependence

If two variables are independent, the correlation coefficient is zero.

WARNING! If the correlation coefficient of two variables is zero, it does not imply they are independent. The correlation coefficient only indicates the linear dependence between two variables. When variables are non-linearly related, they are not independent of each other but their correlation coefficient could be zero.

Correlation Coefficient and X-Y Diagram

The correlation coefficient indicates the direction and strength of the linear dependence between two variables but it does not cover all the existing relationship patterns. With the same correlation coefficient, two variables might have completely different dependence patterns. A scatter plot or X-Y diagram can help to discover and understand additional characteristics of the relationship between variables. The correlation coefficient is not a replacement for examining the scatter plot to study the variables' relationship.

The correlation coefficient by itself does not tell us everything about the relationship between two variables. Two relationships could have the same correlation coefficient, but completely different patterns.

Statistical Significance of the Correlation Coefficient

The correlation coefficient could be high or low by chance (randomness). It may have been calculated based on two small samples that do not provide good inference on the correlation between two populations.

To test whether there is a statistically significant relationship between two variables, we need to run a hypothesis test to determine whether the correlation coefficient is statistically different from zero.

Hypothesis Test Statements

- H₀: r = 0: Null Hypothesis: There is *no* correlation.
- H_1 : $r \neq 0$: Alternate Hypothesis: There is a correlation.

Hypothesis tests will produce p-values as a result of the statistical significance test on r. When the p-value for a test is low (less than 0.05), we can reject the null hypothesis and conclude that r is significant; there is a correlation. When the p-value for a test is > 0.05, then we fail to reject the null hypothesis; there is no correlation.

We can also use the t statistic to draw the same conclusions regarding our test for significance of the correlation coefficient. To use the t-test to determine the statistical significance of the Pearson correlation, calculate the t statistic using the Pearson r value and the sample size, n.

Test Statistic

$$t = \frac{r}{\sqrt{\frac{1 - r^2}{n - 2}}}$$

Critical Statistic

Is the t-value in t-table with (n - 2) degrees of freedom.

If the absolute value of the calculated t value is less than or equal to the critical t value, then we fail to reject the null and claim no statistically significant linear relationship between X and Y.

- If |t| ≤ t_{critical}, we fail to reject the null. There is no statistically significant linear relationship between X and Y.
- If |t| > t_{critical}, we reject the null. There is a statistically significant linear relationship between X and Y.

Using Software to Calculate the Correlation Coefficient

We are interested in understanding whether there is linear dependence between a car's MPG and its weight and if so, how they are related. The MPG and weight data are stored in the "Correlation Coefficient" tab in "Sample Data.xlsx." We will discuss three ways to get the results.

Use Excel to Calculate the Correlation Coefficient

The formula CORREL in Excel calculates the sample correlation coefficient of two data series. The correlation coefficient between the two data series is -0.83, which indicates a strong negative linear relationship between MPG and weight. In other words, as weight gets larger, gas mileage gets smaller.

		9 • (* •)	÷							Boo
C	Hor	ne Insert	Page	Layout	Formulas	Data	Review	View		
No	rmal Page Layou	ut Preview	Views	Full Screen	Ruler Gridlines Message I	✓ H Bar	ormula Bar leadings	Q Zoom	100%	Zoom to Selection
		Workbook V	iews		Sł	now/Hide			Zoom	1
	SUM	1	- (• × ·	✓ f _x	=CORREL(\$A2	:\$A399,I				
	А	В	С			D		CORREL(a	rray1, a	rray2)
1	MPG	weight		=CORR	EL(\$A2:\$A399	,B2:B39	9)			
2	18	3504								
3	15	3693								
4	18	3436								
5	16	3433								
6	17	3449								
7	15	4341								
8	14	4354								
0	14	4212								

Fig. 4.3 Correlation coefficient in Excel

Use Minitab to Calculate the Correlation Coefficient

- Step 1: Stat \rightarrow Basic Statistics \rightarrow Correlation
- Step 2: Select the two variables of interest in the pop-up window "Correlation" and click "OK."

Correl	ation	×
C1	MPG	Variables:
C2	weight	MPG weight
		T
		Method: Pearson correlation 💌
		✓ Display p-values
	Select	
	Help	OK Cancel

Fig. 4.4 Correlation dialog box

ght

Fig. 4.5 Correlation output

The correlation coefficient result (-0.832) appears in the session window. The p-value (0.000) is lower than the alpha level (0.05), indicating the linear correlation is statistically significant. We can claim that there is a linear relationship between mileage and weight.

Interpreting Results

How do we interpret results and make decisions based Pearson's correlation coefficient (r) and p-values?

Let us look at a few examples:

- r = -0.832, p = 0.000 (previous example). The two variables are inversely related and the linear relationship is strong. Also, this conclusion is significant as supported by p-value of 0.00.
- r = -0.832, p = 0.71. Based on r, you should conclude the linear relationship between the two variables is strong and inversely related. However, with a p-value of 0.71, you should then conclude that r is not significant and that your sample size may be too small to accurately characterize the relationship.
- r = 0.5, p = 0.00. Moderately positive linear relationship, r is statistically significant.
- r = 0.92, p = 0.61. Strong positive linear relationship but r is not statistically significant. Get more data.
- r = 1.0, p = 0.00. The two variables have a perfect linear relationship and r is significant.

Correlation Coefficient Calculation

Population Correlation Coefficient (ρ)

$$\rho_{X,Y} = \frac{\text{cov}(X,Y)}{\sigma_X \sigma_Y} = \frac{\text{E}[(X-\mu_X)(Y-\mu_Y)]}{\sigma_X \sigma_Y}$$

Sample Correlation Coefficient (r)

$$r = \frac{\sum_{i=1}^{n} (X_i - \overline{X}) (Y_i - \overline{Y})}{\sqrt{\sum_{i=1}^{n} (X_i - \overline{X})^2} \sqrt{\sum_{i=1}^{n} (Y_i - \overline{Y})^2}}$$

It is only defined when the standard deviations of both X and Y are non-zero and finite. When covariance of X and Y is zero, the correlation coefficient is zero.

4.1.2 X-Y DIAGRAM

What is an X-Y Diagram?

An *X-Y diagram* is a scatter plot depicting the relationship between two variables (i.e., X and Y). Each point on the X-Y diagram represents a pair of X and Y values, with X plotted on the horizontal axis and Y plotted on the vertical axis.

With an X-Y diagram, you can qualitatively assess both the strength and direction of the relationship between X and Y. To quantitatively measure the relationship between X and Y, you may need to calculate the correlation coefficient.

Example 1: Perfect Linear Correlation

In the chart below, there are 10 data points depicted (10 pairs of X and Y values), and they were created using the equation Y = 2X. As a result, all the data points fall onto the straight line of Y = 2X.

The chart demonstrates a perfect positive linear correlation between X and Y since the relationship between X and Y can be perfectly described by a linear equation in a format of $Y = a \times X + b$ where $a \neq 0$. If they were negatively correlated, the line would slope down to the right, meaning as X increases, Y decreases.

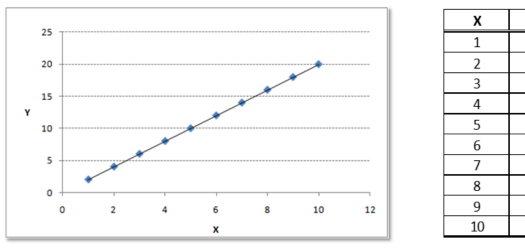


Fig. 4.6 Perfect Positive Linear Correlation

Example 2: Strong Linear Correlation

In this chart, the data points scatter closely around a straight line. When X increases, Y increases accordingly.

This chart demonstrates a strong positive linear correlation between X and Y. The straight line is the best fitting trend that shows approximately how Y changes with X.

Υ

2

4

6

8

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18

20

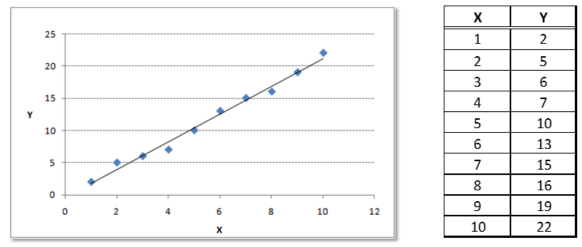


Fig. 4.7 Strong Positive Linear Correlation

In this example, there is a strong correlation, but not perfect. Notice how the points do not all line up perfectly.

Example 3: Weak Linear Correlation

In this chart, the data points scatter remotely around a straight line. When X increases, Y increases accordingly.

This chart demonstrates a weak positive linear correlation between X and Y since the distance between the data points and the trend line is relatively far on average.

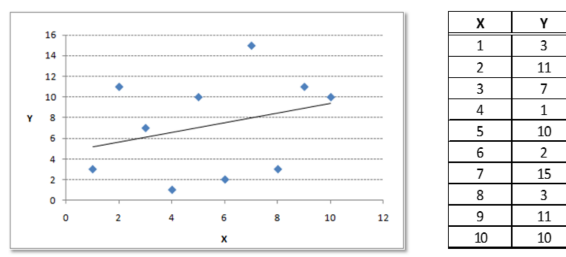


Fig. 4.8 Weak Positive Linear Correlation

The distance between the data points and the best fitting line is much higher than on the previous chart, which is means that the line does not describe the relationship between the X and Y as tightly.

Example 4: Non-Linear Correlation

The X-Y diagram also helps to identify any nonlinear relationship between X and Y.

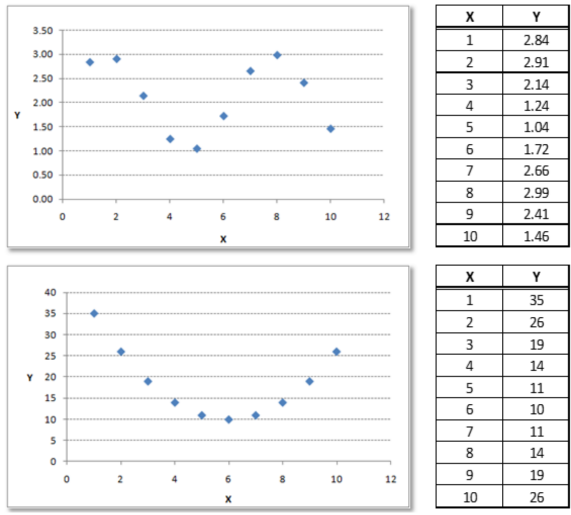


Fig. 4.9 Non-Linear Correlation

The examples show some non-linear correlations. Notice there is a pattern for each relationship, but it is not a line that describes it.

Example 5: Uncorrelated

In this chart, the Y value of each data point is a constant regardless of what the X value is.

Changes in X do not show any relative impact on Y. As a result, there is no correlation between X and Y.

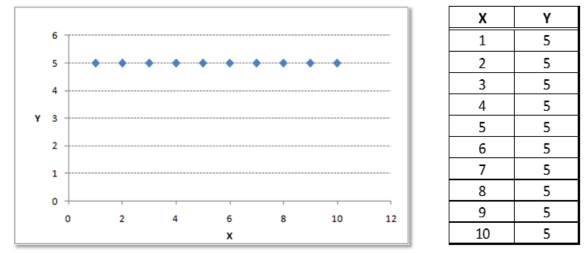
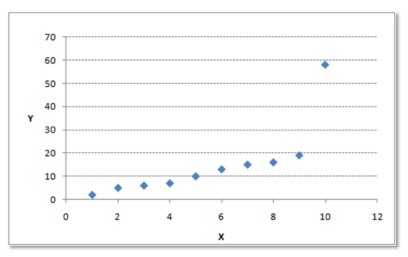


Fig. 4.10 Uncorrelated Correlation

Example 6: Outlier Identification

Another feature of the X-Y diagram is that it allows the user to identify an outlier (or a data point that does not follow the same trend as the rest of the other data points).

In this chart, the last data point does not seem to follow the trend of other data points. When an outlier is spotted, it is good practice to investigate and understand what is causing it.



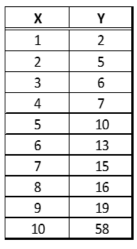


Fig. 4.11 Outlier Identification

Benefits to Using an X-Y Diagram

- An X-Y diagram graphically demonstrates the relationship between two variables (no relationship, linear relationship, nonlinear relationship).
- It suggests whether two variables are associated and helps to identify the linear or nonlinear correlation between X and Y.
- It captures the strength and direction of the relationship between X and Y.
- It helps identify any outliers in the data.

Limitations of the X-Y Diagram

Although the X-Y diagram helps to "spot" interesting features in the data, it does not provide any quantitative conclusions about the data and further statistical analysis is needed to:

- Assess whether the association between variables is statistically significant.
- Measure the strength of the relationship between variables.
- Determine whether outliers exist in the data.
- Quantitatively describe the pattern of the data.

4.1.3 Regression Equations

Correlation and Regression Analysis

The correlation coefficient answers the following questions:

- 1. Are two variables correlated?
- 2. How strong is the relationship between two variables?
- 3. When one variable increases, does the other variable increase or decrease?
- 4. The correlation coefficient *cannot* address the following questions:
- 5. How much does one variable changes when the other variable changes by one unit?
- 6. How can we set the value of one variable to obtain a targeted value of the other variable?
- 7. How can we use the relationship between two variables to make predictions?
- 8. The simple linear regression analysis helps to answer these questions.

What is Simple Linear Regression?

Simple linear regression is a statistical technique to fit a straight line through the data points. It models the quantitative relationship between two variables. It is simple because only one predictor variable is involved. It describes how one variable changes according to the change of another variable. Both variables need to be continuous; there are other types of regression to model discrete data.

Simple Linear Regression Equation

The simple linear regression analysis fits the data to a regression equation in the form

$$Y = \alpha \times X + \beta + e$$

Where:

- *Y* is the dependent variable (the response) and *X* is the single independent variable (the predictor)
- *α* is the slope describing the steepness of the fitting line. β is the intercept indicating the *Y* value when *X* is equal to 0

• *e* stands for error (residual). It is the difference between the actual *Y* and the fitted *Y* (i.e., the vertical difference between the data point and the fitting line).

Ordinary Least Squares

The *ordinary least squares* is a statistical method used in linear regression analysis to find the best fitting line for the data points. It estimates the unknown parameters of the regression equation by minimizing the sum of squared residuals (i.e., the vertical difference between the data point and the fitting line).

In mathematical language, we look for α and β that satisfy the following criteria:

$$\min_{\alpha,\beta} Q(\alpha,\beta) \text{ where } Q(\alpha,\beta) = \sum_{i=1}^{n} e_i^2 = \sum_{i=1}^{n} \left(y_i \text{-}\alpha \text{-}\beta_{x_i} \right)^2$$

The actual value of the dependent variable:

$$Y_i = \alpha \times X_i + \beta + e_i$$

Where: *i* = 1, 2 . . . *n*.

The fitted value of the dependent variable:

$$\hat{y}_i = \alpha \times X_i + \beta$$

Where: *i* = 1, 2 . . . *n*.

By using calculus, it can be shown the sum of squared error is minimal when

$$\beta = \frac{\sum_{i=1}^{n} (X_i - \bar{X})(Y_i - \bar{Y})}{\sum_{i=1}^{n} (X_i - \bar{X})^2}$$

and

$$\alpha = \bar{Y} - \beta \, \bar{X}$$

ANOVA in Simple Linear Regression

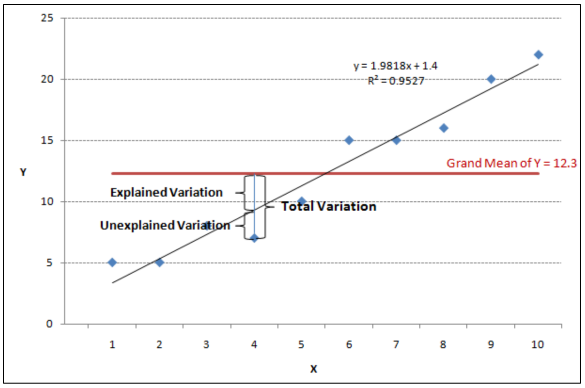


Fig. 4.12 ANOVA in Simple Linear Regression

X: the independent variable that we use to predict;

Y: the dependent variable that we want to predict.

The variance in simple linear regression can be expressed as a relationship between the actual value, the fitted value, and the grand mean—all in terms of Y.

Total Variation = Total Sums of Squares =
$$\sum_{i=1}^{n} (Y_i - \overline{Y})^2$$

Explained Variation = Regression Sums of Squares = $\sum_{i=1}^{n} (\hat{Y}_i - \bar{Y})^2$

Unexplained Variation = Error Sums of Squares = $\sum_{i=1}^{n} (Y_i - Y)^2$

Regression follows the same methodology as ANOVA and the hypothesis tests behind it use the same assumptions.

Variation Components

Total Variation = Explained Variation + Unexplained Variation

i.e., Total Sums of Squares = Regression Sums of Squares + Error Sums of Squares

Degrees of Freedom Components

Total Degrees of Freedom = Regression Degrees of Freedom + Residual Degrees of Freedom i.e., n - 1 = (k - 1) + (n - k), where n is the number of data points, k is the number of predictors Whether the overall model is statistically significant can be tested by using F-test of ANOVA.

- H₀: The model is not statistically significant.
- H_a: The model is statistically significant.

Test Statistic

$$F = \frac{MSR}{MSE} = \frac{\sum_{i=1}^{n} (\hat{Y}_{i} - \bar{Y})^{2} / (k-1)}{\sum_{i=1}^{n} (\hat{Y}_{i} - \hat{Y}_{i})^{2} / (n-k)}$$

Critical Statistic

Is represented by F value in F table with (k – 1) degrees of freedom in the numerator and (n – k) degrees of freedom in the denominator.

- If $F \le F_{critical}$ (calculated F is less than or equal to the critical F), we fail to reject the null. There is no statistically significant relationship between X and Y.
- If F > F_{critical}, we reject the null. There is a statistically significant relationship between X and Y.

Coefficient of Determination

R-squared or R² (also called coefficient of determination) measures the proportion of variability in the data that can be explained by the model.

 R^2 ranges from 0 to 1. The higher R^2 is, the better the model can fit the actual data.

R² can be calculated with the formula:

$$R^{2} = \frac{SS_{regression}}{SS_{total}} = 1 - \frac{SS_{error}}{SS_{total}} = 1 - \frac{\sum_{i=1}^{n} (Y_{i} - \bar{Y})^{2}}{\sum_{i=1}^{n} (Y_{i} - \bar{Y})^{2}}$$

Use Minitab to Run a Simple Linear Regression

Case study: We want to see whether the score on exam one has any statistically significant relationship with the score on the final exam. If yes, how much impact does exam one have on the final exam?

Data File: "Simple Linear Regression" tab in "Sample Data.xlsx"

Step 1: Determine the dependent and independent variables. Both should be continuous variables.

- Y (dependent variable) is the score of final exam.
- X (independent variable) is the score of exam one.

Step 2: Create a scatter plot to visualize whether there seems to be a linear relationship between X and Y.

- 1. Click Graph \rightarrow Scatterplot.
- 2. A new window named "Scatterplots" pops up.
- 3. Click "OK."
- 4. A new window named "Scatterplot Simple" pops up.
- 5. Select "FINAL" as "Y variables" and "EXAM1" as "X variables."

Scatterplot: Simple			X
C1 EXAM1 C2 FINAL	Y variables 1 FINAL 2 3 4 5 6 7 Scale Scale	X variables	<u>D</u> ata View
Select			
Help		<u>0</u> K	Cancel

Fig. 4.13 Scatterplot: Simple dialog box with selected variables

6. Click "OK."

7. A scatter plot is generated in a new window.

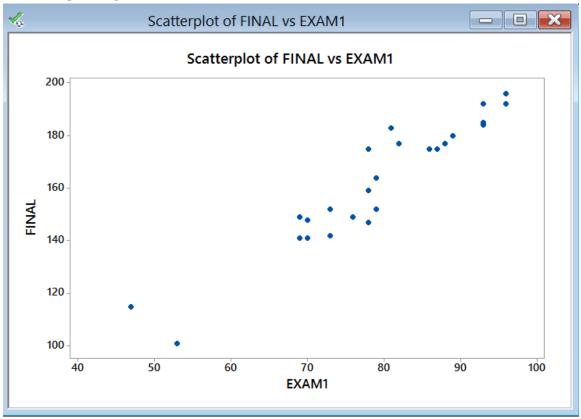


Fig. 4.14 Scatterplot output

Based on the scatter plot, the relationship between exam one and final seems linear. The higher the score on exam one, the higher the score on the final. It appears you could "fit" a line through these data points.

Step 3: Run the simple linear regression analysis.

- 1. Click Stat \rightarrow Regression \rightarrow Regression \rightarrow Fit Regression Model.
- 2. A new window named "Regression" pops up.

3. Select "FINAL" as "Response" and "EXAM1" as "Continuous Predictors."

Regre	ssion	
C1 C2	EXAM1 FINAL	Responses:
		Continuous predictors:
		Categorical predictors:
		Model Optio <u>n</u> s <u>Cod</u> ing <u>S</u> tepwise
	Select	<u>G</u> raphs <u>R</u> esults Storage
	Help	<u>Q</u> K Cancel

Fig. 4.15 Regression dialog box with selected variables

- 4. Click the "Storage" button.
- 5. Check the box of "Residuals" so that the residuals can be saved automatically in the last column of the data table.

Regression: Storage		x
E Ets	Co <u>e</u> fficients	
Residuals	Design <u>m</u> atrix	
<u>S</u> tandardized residuals		
Deleted residuals		
Leverages		
Cook's distance		
Help	<u>O</u> K Ca	ncel

Fig. 4.16 Regression: Storage dialog box

6. Click "OK."

7. The regression analysis results appear in the new window.

Anal	ysis of	Var	ianc	e				Model S	Summa	ry			
Sour	ce	DF	Ad	j ss	Adj MS	F-Value	P-Value	S	R-sq	R-sq(adj) R-sq(p	red)	
Regr	ession	1	1241	18.8	12418.8	196.15	0.000	7.95689	89.51%	89.05%	6 86.	61%	
EXA	M1	1	1241	18.8	12418.8	196.15	0.000						
Error		23	145	56.2	63.3								
Lac	k-of-Fit	14	83	37.0	59.8	0.87	0.607	Coeffici	ents				
Pur	e Error	9	61	19.2	68.8			Term	Coef	SE Coef	T-Value	P-Value	VIF
Total		24	1387	75.0				Constant	15.6	10.6	1.48	0.153	
its a	and Dia	igno	ostic	s for	Unusu	al Obse	rvations	EXAM1	1.852	0.132	14.01	0.000	1.00
					Std								
Obs	FINAL		Fit	Resid	Resid			Regress	ion Equ	uation			
8	115.00	102	.69	12.31	1.88	Х		FINAL =	= 15.6 +	- 1.852 EX/	۵M1		
18	183.00	165	6.67	17.33	2.22	R			- 15.0 1	1.052 L/0			



Step 4: Check whether the model is statistically significant. If not significant, we will need to reexamine the predictor or look for new predictors before continuing. R² measures the percentage of variation in the data set that can be explained by the model. 89.5% of the variability in the data can be accounted for by this linear regression model. "Analysis of Variance" section provides an ANOVA table covering degrees of freedom, sum of squares, and mean square information for total, regression and error. The p-value of the F-test is lower than the α level (0.05), indicating that the model is statistically significant.

The p-value is 0.0001; therefore, we reject the null and claim the model is statistically significant. The R square value says that 89.5% of the variability can be explained by this model.

Step 5: Understand regression equation

The estimates of slope and intercept are shown in "Parameter Estimate" section. In this example, $Y = 15.6 + 1.85 \times X$, where X is the score on Exam 1 and Y is the final exam score. One unit increase in the score of Exam1 would increase the final score by 1.85.

Regressio	n Eo	quatio	n				
FINAL =	15.6	5 + 1.852	EXAM1				
Analysis of	Vari	ance				I	
Source	DF	Adj SS		F-Value	P-Value		
Regression	1	12418.8	12418.8	196.15	0.000		
EXAM1	1		12418.8	196.15	0.000		
Error	23	1456.2	63.3	0.07	0.007		
Lack-of-Fit		837.0	59.8	0.87	0.607		
Pure Error Total	9 24	619.2 13875.0	68.8				
Model Sun s	nmaı <u>R-sq</u> .51%	ry R-sq(adj) 89.05%		red)61%			
Model Sun s 7.95689 89	<u>R-sq</u> .51%	R-sq(adj)					
Model Sun s 7.95689 89 Coefficient	<u>R-sq</u> .51%	<u>R-sq(adj)</u> 89.05%	5 86.0		VIF		
Model Sun S 7.95689 89 Coefficient	<u>R-sq</u> .51%	<u>R-sq(adj)</u> 89.05%	5 86.0	61%	VIF		
Model Sun S 7.95689 89 Coefficient Term C Constant	<u>R-sq</u> .51% ts Coef	R-sq(adj) 89.05% SE Coef	5 86.6 T-Value	61% P-Value	VIF 1.00		
Model Sun S 7.95689 89 Coefficient Term C Constant	R-sq .51% S Coef 15.6 .852	R-sq(adj) 89.05% SE Coef 10.6 0.132	5 86.6 <u>T-Value</u> 1.48 14.01	P-Value 0.153 0.000	1.00	5	
S 7.95689 89 Coefficient Term C Constant 1	<u>R-sq</u> .51% ts Coef 15.6 .852 agno	R-sq(adj) 89.05% SE Coef 10.6 0.132	<u>T-Value</u> 1.48 14.01 r Unusu Std	P-Value 0.153 0.000	1.00	s	
S 7.95689 89 Coefficient Term Constant EXAM1 1 Tits and Dia	<u>R-sq</u> .51% ts Coef 15.6 .852 agno	<u>R-sq(adj)</u> 89.05% <u>SE Coef</u> 10.6 0.132 ostics fo Fit Resid	<u>T-Value</u> 1.48 14.01 r Unusu Std Resid	P-Value 0.153 0.000	1.00	s	

Fig. 4.18 Regression Analysis: Final versus EXAM1

Interpreting the Results

Let us say you are the professor and you want to use this prediction equation to estimate what two of your students might get on their final exam.

Rsquare Adj = 89.0%

• 89% of the variation in FINAL can be explained by EXAM1

P-value of the F-test = 0.000

• We have a statistically significant model

Prediction Equation: 15.6 + 1.85 × EXAM1

- 15.6 is the Y intercept, all equations will start with 15.6
- 1.85 is the EXAM1 Coefficient: multiply it by EXAM1 score

Because the model is significant, and it explains 89% of the variability, we can use the model to predict final exam scores based on the results of Exam1.

Let us assume the following:

- Student "A" exam 1 results were: 79
- Student "B" exam 1 results were: 94

Remember our prediction equation 15.6 + 1.85 × Exam1?

Now apply the equation to each student:

Student "A" Estimate: 15.6 + (1.85 × 79) = 161.8

Student "B" Estimate: 15.6 + (1.85 × 94) = 189.5

By simply replacing exam 1 scores into the equation we can predict their final exam scores. But the key thing about the model is whether or not it is useful. In this case, the professor can use the results to Figure out where to spend his time helping students.

4.1.4 RESIDUALS ANALYSIS

What are Residuals?

Residuals are the vertical differences between actual values and the predicted values or the "fitted line" created by the regression model.

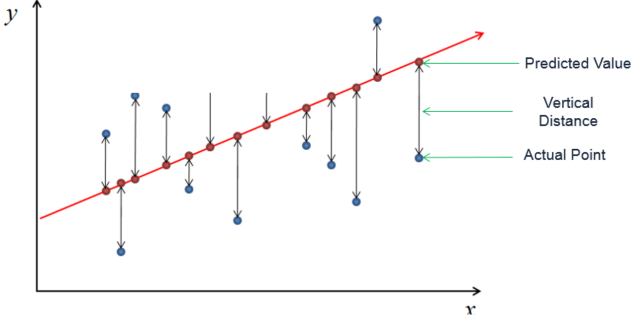


Fig. 4.19 Understanding Residuals

A residual is the vertical (y-axis) distance between an actual value and a value predicted by a regression model.

Why Perform Residuals Analysis?

Regression equations are generated on the basis of certain statistical assumptions. *Residuals analysis* helps to determine the validity of these assumptions.

The assumptions are:

- The residuals are normally distributed, mean equal to zero.
- The residuals are independent.
- The residuals have a constant variance.
- The underlying population relationship is linear.
- If residuals performance does not meet the requirements, we will need to rebuild the model by replacing the predictor with a new one, adding new predictors, building non-linear models, and so on.

A linear regression model is not always appropriate for the data; therefore, assess the appropriateness of the model by analyzing the residuals.

Use Minitab to Perform Residuals Analysis

The residuals of the model are saved in the last column of the following data table.

🖽 Wo	orksheet 9 **	ŕ	
Ŧ	C1	C2 🗾	C3
	EXAM1	FINAL	RESI
1	73	152	1.1488
2	93	185	-2.9003
3	89	180	-0.4905
4	96	196	2.5424
5	73	142	-8.8512
6	53	101	-12.8021
7	69	149	5.5586
8	47	115	12.3127
9	87	175	-1.7855
10	79	164	2.0341
11	69	141	-2.4414

Fig. 4.20 Residual Data

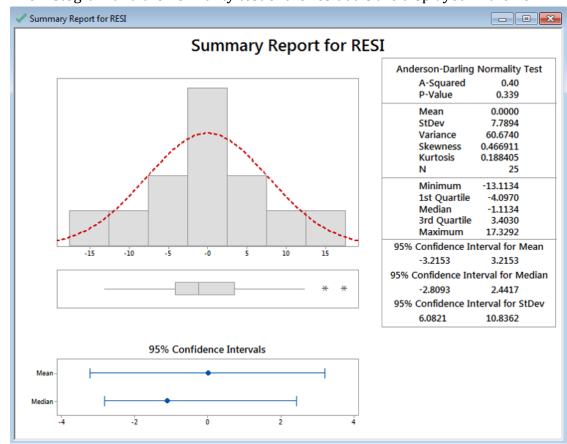
Step 1: Check whether residuals are normally distributed around the mean of zero.

- 1. Click Stat \rightarrow Basic Statistics \rightarrow Graphical Summary.
- 2. A new window named "Graphical Summary" pops up.
- 3. Select "RESI" as "Variables."

Graphical Summary	×
C1 EXAM1 C2 FINAL C3 RESI	Variables:
	By variables (optional):
Select	
Help	OK Cancel

Fig. 4.21 Graphical Summary dialog box with variable

4. Click "OK."



5. The histogram and the normality test of the residuals are displayed in the new window

Fig. 4.22 Graphical Summary output

The mean of residuals is -0.0000.

The first assumption about residuals is that they are normally distributed and have a mean equal to zero. The Anderson–Darling test is used to test the normality.

- H₀: The residuals are normally distributed.
- H₁: The residuals are not normally distributed.

Since the p-value (0.339) is greater than the alpha level (0.05), we fail to reject the null hypothesis; the residuals are normally distributed.

Step 2: If the data are in time order, run the IR chart to check whether residuals are independent.

- 1. Click Stat \rightarrow Control Charts \rightarrow Variable Charts for Individuals \rightarrow I-MR.
- 2. A new window named "Individuals Moving Range Chart" pops up.
- 3. Select "RESI" as "Variables"
- 4. Click the I-MR Options button then the "Tests" tab and select "Perform all tests for causes."

Individuals-Moving Range	Chart		×
C1 EXAM1 C2 FINAL C3 RESI	Variables: RESI Scale Multiple Graphs	Labels Data Options	I-MR Options
Select			
Help		<u>О</u> К	Cancel

Fig. 4.23 Individuals – Moving Range Chart dialog box with variable

- 5. Click "OK." In the options window and again in the I-MR window.
- 6. The control charts are shown automatically in the new window.

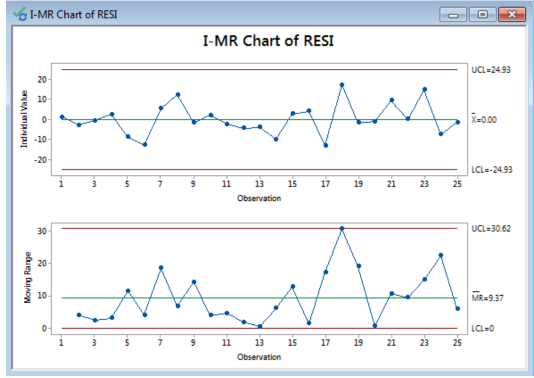


Fig. 4.24 Individuals - Moving Range Chart output

If no data points are out of control in both the I-chart and MR chart, the residuals are independent of each other. If the residuals are not independent, it is possible that some important predictors are not included in the model. In this example, since the IR chart is in control, residuals are independent. An I-MR chart, showing that no data points are out of control, indicates that the data are independent.

Step 3: Check whether residuals have equal variance across the predicted responses.

- 1. Label a new column in your Minitab data table "Fitted".
- 2. Right click on the column labeled "Fitted" in the data table.
- 3. Select Formulas \rightarrow Assign Formula to Column.
- 4. A new window named "Assign Formula to C4" appears.
- 5. Enter the regression equation "15.6+1.85*EXAM1" into the "Expression" box.

A	ssig	n Formula to C4	1							x
ſ	C1	EXAM1	Exp	ressi	on:					
	C2 C3	FINAL RESI	15.	.6+1	.85*'	EXAM	1			*
									Functions:	
			7	8	9	+	=	ŧ	All functions	-
			4	5	6	-	<	>	Absolute value	*
			1	2	3	*	≤	≥	Antilog Any	
			0		[]	1	A	nd	Arccosh Arccosine	
Ľ						^	C	Dr	Arcsine	Ψ.
		Select				()	N	ot	Select	
		Help							ОК Са	ncel

Fig. 4.25 Assign Formula to C4 dialog box with formula

- 6. Click "OK."
- 7. The column of the fitted values is created in the data table, label if "Fitted"
- 8. Create a scatter plot with Y being the residuals and the X being the fitted values.
- 9. Click Graph \rightarrow Scatterplot.
- 10. A new window named "Scatterplots" appears, select OK with Simple selected.
- 11. Select "RESI" as the "Y variables" and "Fitted" as the "X variables."

Scat	terplot: Simple		×
C1 C2 C3 C4	FINAL RESI	X variables	Data View
	Select		
	Help	<u>O</u> K	Cancel

Fig. 4.26 Scatterplots: Simple dialog box with variables selected 12. Click "OK."

13. The scatter plot appears in a new window.

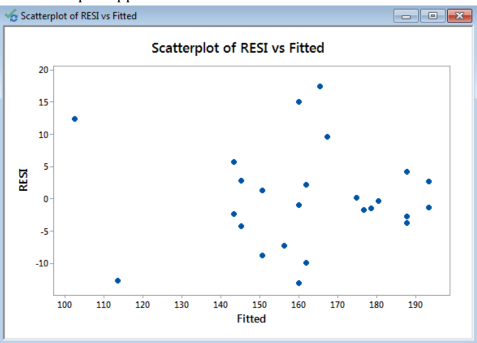


Fig. 4.27 Scatterplot output

We are looking for the pattern in which residuals spread out evenly around zero from the top to the bottom. Finally, we verify that the residuals have equal variance across the predicted responses. In this example, you can see how the residual values disperse evenly around zero.

Heteroscedasticity is the condition where the assumption of equal variance is violated, and can lead us to believe a variable is a predictor when it is not.

4.2 MULTIPLE REGRESSION ANALYSIS

4.2.1 Non-Linear Regression

Linear and Non-Linear

The word linear originally comes from Latin word linearis meaning "created by lines."

A linear function in mathematics follows the following pattern (i.e., the output is proportional to its input):

$$f(x) = \alpha * x + \beta$$
$$f(x_1, x_2, ..., x_n) = \alpha_1 * x_1 + \alpha_2 * x_2 + ... + \alpha_n * x_n + \beta$$

A non-linear function does not follow the above pattern. There are usually exponents, logarithms, power, polynomial components, and other non-linear functions of the independent variables and parameters.

Non-Linear Relationships Using Linear Models

Many non-linear relationships can be transformed into linear relationships, and from there we can use linear regression methods to model the relationship. Some non-linear relationships cannot be transformed to linear ones and we need to apply other methods to build the non-linear models. In this section, we will focus on building non-linear regression models using linear transformation (i.e., transforming the independent or dependent variables or parameters to generate a linear function).

Assumptions in Using Non-Linear Regression

The population relationship is non-linear based on a reliable underlying theory. Across the range of all the possible values of the independent variables, the non-linear relationship applies. It is possible that at some extreme values the relationship between variables changes dramatically.

Non-Linear Functions: Transforming to Linear

Examples of non-linear functions that can be transformed to linear functions:

- Exponential Function
- Inverse Function
- Polynomial Function
- Power Function

The following section illustrates how to mathematically transform several non-linear functions to linear functions.

Exponential Function Exponential Function:

$$Y = a \times b^X$$

Transformation:

$$\log Y = \log a + X \times \log b$$

Inverse Function Inverse Function:

$$Y = a + b \times \frac{1}{X}$$

Transformation:

$$Y = a + b \times Z$$
, where $Z = \frac{1}{x}$

Polynomial Function Polynomial Function:

$$Y = a + b \times X + c \times X^2$$

Transformation:

$$Y = a + b \times X + c \times Z$$
, where $Z = X^2$

Power Function Power Function:

 $Y = a \times X^b$

Transformation:

$$\log Y = \log a + b \times \log X$$

4.2.2 Multiple Linear Regression

What is Multiple Linear Regression?

Multiple linear regression is a statistical technique to model the relationship between one dependent variable and two or more independent variables by fitting the data set into a linear equation.

The difference between simple linear regression and multiple linear regression:

- Simple linear regression only has one predictor.
- Multiple linear regression has two or more predictors.

Multiple Linear Regression Equation

 $Y = \alpha_1 * X_1 + \alpha_2 * X_2 + ... + \alpha_p * X_p + \beta + e$

Where:

- *Y* is the dependent variable (response)
- $X_1, X_2 \dots X_p$ are the independent variables (predictors). There are p predictors in total.

Both dependent and independent variables are continuous.

• β is the intercept indicating the *Y* value when all the predictors are zeros

- *α*₁, *α*₂... *α*_p are the coefficients of predictors. They reflect the contribution of each independent variable in predicting the dependent variable.
- *e* is the residual term indicating the difference between the actual and the fitted response value.

Use Minitab to Run a Multiple Linear Regression

Case study: We want to see whether the scores in exam one, two, and three have any statistically significant relationship with the score in final exam. If so, how are they related to final exam score? Can we use the scores in exam one, two, and three to predict the score in final exam?



Data File: "Multiple Linear Regression" tab in "Sample Data.xlsx."

Step 1: Determine the dependent and independent variables, all should be continuous. Y (dependent variable) is the score of final exam. X_1 , X_2 , and X_3 (independent variables) are the scores of exams one, two, and three respectively. All x variables are continuous.

Step 2: Start building the multiple linear regression model

- 1. Click Stat \rightarrow Regression \rightarrow Regression \rightarrow Fit Regression Model
- 2. A new window named "Regression" pops up.
- 3. Select "FINAL" as "Response" and "EXAM1", "EXAM2" and "EXAM3" as "Predictors."

	Regre	ssion		x
	C1 C2 C3 C4	EXAM1 EXAM2 EXAM3 FINAL	Responses: FINAL	*
			Continuous predictors: EXAM1 EXAM2 EXAM3	4
And the second se			Categorical predictors:	*
			Model Options Coding Stepwise	e
		Select	<u>G</u> raphs <u>Results</u> <u>Storage</u>	
		Help	<u>Q</u> K Cance	1

Fig. 4.28 Regression dialog box with variables

4. Click the "Graph" button, select the radio button "Four in one" and click "OK."

Regression: Graphs	×
	Residuals for plots: Regular
	Residuals plots
	○ Individual plots ☐ Histogram of residuals
	☐ Normal probability plot of residuals
	Residuals <u>v</u> ersus fits Residuals versus order
	C Four in one
	Residuals versus the variables:
	×
Select	
Help	<u>O</u> K Cancel

Fig. 4.29 Regression: Graphs dialog box

5. Click the "Storage" button, check the boxes of "Residuals" and "DFITS" and click "OK."

Regression: Storage		×
Eits	Co <u>e</u> ffic	cients
Residuals	🗍 Design	matrix
□ <u>S</u> tandardized res	iduals	
Deleted residuals	i -	
Leverages		
Cook's distance		
Help		OK Cancel

Fig. 4.30 Regression: Storage dialog box

- 6. Click "OK" in the window named "Regression."
- 7. The regression analysis results appear in a session window and the four residual plots appear in another window named "Residual Plots for FINAL."

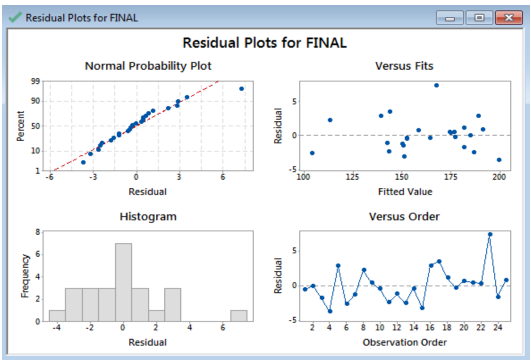


Fig. 4.31 Residual Plots for Final

Step 3: Check whether the whole model is statistically significant. If not, we need to re-examine the predictors or look for new predictors before continuing.

- H₀: The model is not statistically significant (i.e., all the parameters of predictors are not significantly different from zeros).
- H₁: The model is statistically significant (i.e., at least one predictor parameter is significantly different from zero).

In this example, p-value is much smaller than alpha level (0.05), hence we reject the null hypothesis; the model is statistically significant.

Step 4: Check whether multicollinearity exists in the model.

The VIF information is automatically generated in table of Coefficients.

Analysis o	of Va	riance						
Source	DF	Adj SS	Adj MS	F-Value	P-Value			
Regression	3	13731.5	4577.17	670.09	0.000			
EXAM1	1	58.7	58.73	8.60	0.008			
EXAM2	1	197.7	197.67	28.94	0.000			
EXAM3	1	877.3	877.30	128.43	0.000			
Error	21	143.4	6.83					
Total	24	13875.0						
Model Su	imma	arv						
c	P co	P. cg/ad		(rod)				
2 61357 S	R-sq	D I						
2.61357	98.97%			ored) 51%				
	98.97%		% 98.	51%	VIF		I	
2.61357 S	98.97%	98.829 SE Coef	% 98.	51%	VIF		I	
2.61357 S Coefficier Term	98.97%	98.829 SE Coef 3.76	% 98. T-Value	51% P-Value 0.262			I	
2.61357 S Coefficier Term Constant	08.97%	98.829 SE Coef 3.76 0.121	% 98. <u>T-Value</u> -1.15	P-Value 0.262 0.008	7.81		I	
2.61357 S Coefficier Term Constant EXAM1	08.97% nts Coef -4.34 0.356 0.543	98.829 SE Coef 3.76 0.121 0.101	7-Value -1.15 2.93	P-Value 0.262 0.008 0.000	7.81 5.59		I	
2.61357 S Coefficien Term Constant EXAM1 EXAM2 EXAM3	08.97% Coef -4.34 0.356 0.543 1.167	98.829 SE Coef 3.76 0.121 0.101 0.103	T-Value -1.15 2.93 5.38	P-Value 0.262 0.008 0.000	7.81 5.59		I	
2.61357 S Coefficien Term Constant EXAM1 EXAM2	08.97% Coef -4.34 0.356 0.543 1.167	98.829 SE Coef 3.76 0.121 0.101 0.103	T-Value -1.15 2.93 5.38	P-Value 0.262 0.008 0.000	7.81 5.59		I	
2.61357 S Coefficien Term Constant EXAM1 EXAM2 EXAM3	08.97% Coef -4.34 0.356 0.543 1.167 on Eq	98.829 SE Coef 3.76 0.121 0.101 0.103 uation	T-Value -1.15 2.93 5.38 11.33	P-Value 0.262 0.008 0.000 0.000	7.81 5.59 5.16	ХАМЗ	I	
2.61357 S Coefficien Term Constant EXAM1 EXAM2 EXAM3 Regressio	08.97% Coef -4.34 0.356 0.543 1.167 on Eq	98.829 SE Coef 3.76 0.121 0.101 0.103 uation	T-Value -1.15 2.93 5.38 11.33	P-Value 0.262 0.008 0.000 0.000	7.81 5.59 5.16	EXAM3	I	
2.61357 S Coefficien Term Constant EXAM1 EXAM2 EXAM3 Regression FINAL =	-4.34 -4.34 0.356 0.543 1.167 on Eq -4.34	98.829 <u>SE Coef</u> 3.76 0.121 0.101 0.103 uation + 0.356 E	T-Value -1.15 2.93 5.38 11.33	P-Value 0.262 0.008 0.000 0.000	7.81 5.59 5.16 2 + 1.167		Ι	
2.61357 S Coefficien Term Constant EXAM1 EXAM2 EXAM3 Regressio	-4.34 -4.34 0.356 0.543 1.167 on Eq -4.34	98.829 <u>SE Coef</u> 3.76 0.121 0.101 0.103 uation + 0.356 E	T-Value -1.15 2.93 5.38 11.33	P-Value 0.262 0.008 0.000 543 EXAM ual Obs	7.81 5.59 5.16 2 + 1.167		I	
2.61357 S Coefficien Term Constant EXAM1 EXAM2 EXAM3 Regression FINAL =	08.97% 14.34 0.356 0.543 1.167 0n Eq -4.34 Diagn	98.829 <u>SE Coef</u> 3.76 0.121 0.101 0.103 uation + 0.356 E	T-Value -1.15 -2.93 5.38 11.33	P-Value 0.262 0.008 0.000 543 EXAM ual Obs	7.81 5.59 5.16 2 + 1.167		Ι	

Fig. 4.32 Regression output

We use the VIF (Variance Inflation Factor) to determine if multicollinearity exists.

Multicollinearity

Multicollinearity is the situation when two or more independent variables in a multiple regression model are correlated with each other. Although multicollinearity does not necessarily reduce the predictability for the model as a whole, it may mislead the calculation for individual independent variables. To detect multicollinearity, we use VIF (Variance Inflation Factor) to quantify its severity in the model.

Variance Inflation Factor (1)

VIF quantifies the degree of multicollinearity for each individual independent variable in the model.

VIF calculation:

Assume we are building a multiple linear regression model using p predictors.

$$Y = \alpha_1 \times X_1 + \alpha_2 \times X_2 + \dots + \alpha_p \times X_p + \beta$$

Two steps are needed to calculate VIF for X₁.

Step 1: Build a multiple linear regression model for X_1 by using $X_2, X_3 \dots X_p$ as predictors.

$$X_1 = a_2 \times X_2 + a_3 \times X_3 + \dots + a_p \times X_p + b$$

Step 2: Use the R^2 generated by the linear model in step 1 to calculate the VIF for X_1 .

$$VIF = \frac{1}{1 - R^2}$$

Apply the same methods to obtain the VIFs for other X's. The VIF value ranges from one to positive infinity.

Variance Inflation Factor (2)

Rules of thumb to analyze variance inflation factor (VIF):

- If VIF = 1, there is no multicollinearity.
- If 1 < VIF < 5, there is small multicollinearity.
- If VIF \geq 5, there is medium multicollinearity.
- If VIF \geq 10, there is large multicollinearity.

How to Deal with Multicollinearity

- 1. Increase the sample size.
- 2. Collect samples with a broader range for some predictors.
- 3. Remove the variable with high multicollinearity and high p-value.
- 4. Remove variables that are included more than once.
- 5. Combine correlated variables to create a new one.

In this section, we will focus on removing variables with high VIF and high p-value.

Step 5: Deal with multicollinearity:

- 1. Identify a list of independent variables with VIF higher than 5. If no variable has VIF higher than 5, go to Step 6 directly.
- 2. Among variables identified in Step 5.1, remove the one with the highest p-value.
- 3. Run the model again, check the VIFs and repeat Step 5.1.

Note: we only remove one independent variable at a time.

In this example, all three predictors have VIF higher than 5. Among them, EXAM1 has the highest p-value. We will remove EXAM1 from the equation and run the model again.

Analysis c	of Va	riance					
Source	DF	Adj SS	Adj MS	F-Value	P-Value		
Regression	3	13731.5	4577.17	670.09	0.000		
EXAM1	1	58.7	58.73	8.60	0.008		
EXAM2	1	197.7	197.67	28.94	0.000		
EXAM3	1	877.3	877.30	128.43	0.000		
Error	21	143.4	6.83				
Total	24	13875.0					
Aodel Su	mma	ary					
		-		b			
S	R-sq	R-sq(ad					
S		R-sq(ad		ored) 51%			
S 2.61357 9	<u>R-sq</u> 8.97%	R-sq(ad					
S 2.61357 9	<u>R-sq</u> 8.97%	R-sq(ad					
s 2.61357 9 C oefficien	<u>R-sq</u> 8.97%	R-sq(ad			VIF		
S 2.61357 9 Coefficien	<u>R-sq</u> 8.97%	R-sq(ad 98.829	% 98.	51%	VIF		
S 2.61357 9 C oefficien Term Constant	<u>R-sq</u> 8.97% nts Coef	R-sq(ad 98.829 SE Coef	% 98. T-Value	51% P-Value	VIF 7.81		
2.61357 9 Coefficien Term Constant EXAM1	<u>R-sq</u> 8.97% nts <u>Coef</u> -4.34	R-sq(ad 98.82 SE Coef 3.76	% 98. <u>T-Value</u> -1.15	51% P-Value 0.262			

Fig. 4.33 Regression Analysis Final vs. Exam 1, 2, and 3

Run the new multiple linear regression with only two predictors (i.e., EXAM2 and EXAM3).

Check the VIFs of EXAM2 AND EXAM3. They are both smaller than 5; hence, there is little multicollinearity existing in the model.

nalysis o	of Var	iance				
ource	DF	Adj SS	Adj MS	F-Value	P-Value	_
egression	2	13672.8	6836.39	743.91	0.000	-
EXAM2	1	555.1	555.11	60.41	0.000	
EXAM3	1	1686.0	1685.97	183.46	0.000	
rror	22	202.2	9.19			
- + - 1	~ 4	42075.0				
	24 mma	13875.0				
otal I <mark>odel Su</mark> S			j) R-sq(p	red)		
odel Su s	mma	iry		<u>red)</u> 18%		
odel Su s .03146 9 pefficier	mma <u>R-sq</u> 18.54%	R-sq(ad) 98.419	% 98.	18%	VIE	
odel Su s .03146 9 cefficier	mma <u>R-sq</u> 18.54% nts Coef	R-sq(ad) 98.419 SE Coef	% 98. F T-Value	18% P-Value		
odel Su s 03146 9 oefficier erm onstant	mma <u>R-sq</u> 18.54%	R-sq(ad 98.419 SE Coef 4.37	6 98. T-Value -0.99	18% P-Value 0.331	1	

Fig. 4.34 Regression Analysis Final vs. Exam 2, and 3

Step 6: Identify the statistically insignificant predictors. Remove one insignificant predictor at a time and run the model again. Repeat this step until all the predictors in the model are statistically significant.

Insignificant predictors are the ones with p-value higher than alpha level (0.05). When p > alpha level, we fail to reject the null hypothesis; the predictor is not significant.

- H₀: The predictor is not statistically significant.
- H₁: The predictor is statistically significant.

As long as the p-value is greater than 0.05, remove the insignificant variables one at a time in the order of the highest p-value. Once one insignificant variable is eliminated from the model, we need to run the model again to obtain new p-values for other predictors left in the new model. In this example, both predictors' p-values are smaller than alpha level (0.05). As a result, we do not need to eliminate any variables from the model.

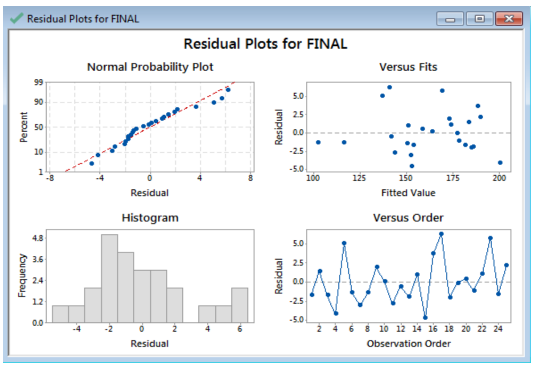


Fig. 4.35 Residual Plots for Final

Step 7: Interpret the regression equation

The multiple linear regression equation appears automatically at the top of the session window. "Parameter Estimates" section provides the estimates of parameters in the linear regression equation.

Analysis o	of Var	iance				
Source	DF	Adj SS	Adj MS	F-Value	P-Value	
Regression	2	13672.8	6836.39	743.91	0.000	
EXAM2	1	555.1	555.11	60.41	0.000	
EXAM3	1	1686.0	1685.97	183.46	0.000	
Error	22	202.2	9.19			
Total	24	13875.0				
3.03146 S	98.54%	98.419	6 98. ⁻	18%		
Term	Coef	SE Coef	T-Value	P-Value	VIF	
Constant	-4.34					
EXAM2	0.7222	0.0929	7.77	0.000	3.53	
EXAM3	1.3375	0.0987	13.54	0.000	3.53	
Regressic	on Equ	uation				

Fig. 4.36 Interpreting the Regression Equation

Now that we have removed multicollinearity and all the insignificant predictors, we have the parameters for the regression equation.

Interpreting the Results Rsquare Adj = 98.4%

• 98% of the variation in FINAL can be explained by the predictor variables EXAM2 & EXAM3.

P-value of the F-test = 0.000

• We have a statistically significant model.

Variables p-value:

• Both are significant (less than 0.05).

VIF

• EXAM2 and EXAM3 are both below 5; we're in good shape!

Equation: -4.34 + 0.722*EXAM2 + 1.34*EXAM3

- -4.34 is the Y intercept, all equations will start with -4.34.
- 0.722 is the EXAM2 coefficient; multiply it by EXAM2 score.
- 1.34 is the EXAM3 coefficient; multiply it by EXAM3 score.

Let us say you are the professor again, and this time you want to use your prediction equation to estimate what one of your students might get on their final exam.

Assume the following:

- Exam 2 results were: 84
- Exam 3 results were: 102

Use your equation: -4.34 + 0.722*EXAM2 + 1.34*EXAM3

Predict your student's final exam score:

-4.34 + (0.722*84) + (1.34*102) = -4.34 + 60.648 + 136.68 = 192.988

Nice work again! Now you can use your "magic" as the smart and efficient professor and allocate your time to other students because this one projects to perform much better than the average score of 162. Now that we know that exams two and three are statistically significant predictors, we can plug them into the regression equation to predict the results of the final exam for any student.

4.2.3 Confidence and Prediction Intervals

Prediction

The purpose of building a regression model is not only to understand what happened in the past but more importantly to *predict* the future based on the past. By plugging the values of independent variables into the regression equation, we obtain the estimation/prediction of the dependent variable.

Uncertainty of Prediction

We build the regression model using the sample data to describe as close as possible the true population relationship between dependent and independent variables. Due to noise in the data, the prediction will probably differ from the true response value. However, the true response value might fall in a range around the prediction with some certainty. To measure the uncertainty of the prediction, we need confidence interval and prediction interval.

Confidence Interval

The *confidence interval* of the prediction is a range in which the population mean of the dependent variable would fall with some certainty, given specified values of the independent variables.

The width of confidence interval is related to:

- Sample size
- Confidence level
- Variation in the data

We build the model based on a sample set $\{y_1, y_2 \dots y_n\}$. The confidence interval is used to estimate the value of the population mean μ of the underlying population. The focus of the confidence interval is represented by the unobservable population parameters. The confidence interval accounts for the uncertainty in the estimates of regression parameters.

Prediction Interval

The *prediction interval* is a range in which future values of the dependent variable would fall with some certainty, given specified values of the independent variables. We build the model based on a sample set $\{y1, y2, ..., yn\}$. The prediction interval is used to estimate the value of future observation y_{n+1} . The focus of the prediction interval is represented by the future observations. Prediction interval is wider than confidence interval because it accounts for the uncertainty in the estimates of regression parameters and the uncertainty of the new measurement.

Use Minitab to Obtain Prediction and Confidence Interval

Steps in Minitab to obtain prediction, confidence interval, and prediction interval:

- 1. Click Stat \rightarrow Regression \rightarrow Regression \rightarrow Predict
- 2. Choose "FINAL" as Response
- 3. Select "Enter columns of values" from second dropdown menu

4. Click on "Exam 2" for the first column and "Exam 3" for the second column.

Predict	
C1 EXAM1 C2 EXAM2 C3 EXAM3 C4 FINAL C5 RESI C6 DFIT C7 RESI_1 C8 DFIT_1	Response: FINAL
Select	Options Results Storage View Model
Help	<u>Q</u> K Cancel

Fig. 4.37 Predict dialog box with variables

- 5. Click "OK"
- *6.* The fitted response (PFIT), the confidence interval (CLIM), and the prediction interval (PLIM) are automatically added to the data table.

C9	C10	C11	C12	C13	C14
PFITS	PSEFITS	CLIM	CLIM_1	PLIM	PLIM_1
153.748	0.91198	151.857	155.640	147.183	160.314
183.601	0.87406	181.788	185.414	177.058	190.144
181.755	0.83474	180.024	183.486	175.234	188.276
200.185	1.16029	197.779	202.591	193.453	206.917
136.951	0.90445	135.075	138.826	130.390	143.511
102.445	1.76442	98.786	106.104	95.171	109.719
152.091	0.66428	150.713	153.468	145.654	158.527

Fig. 4.38 Predict output

4.2.4 RESIDUALS ANALYSIS

Remember what Residuals Are?

Residuals are the vertical difference between actual values and the predicted values or the "fitted line" created by the regression model.

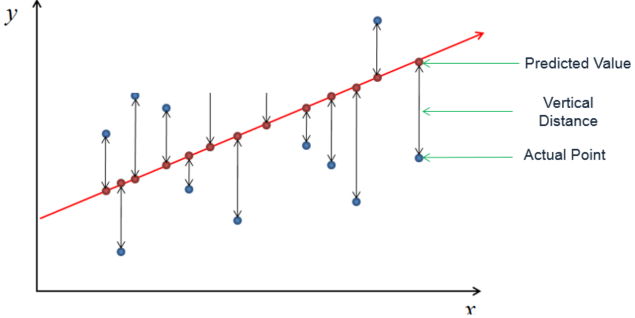


Fig. 4.39 Residuals

Residuals are an important component of evaluating the quality of a regression model. A residual is the vertical (y-axis) distance between an actual value and a value predicted by a regression model.

Why Perform Residuals Analysis?

The *regression equation* generated based on the sample data can make accurate statistical inference only if certain assumptions are met. Residuals analysis can help to validate these assumptions. The following assumptions must be met to ensure the reliability of the linear regression model:

- The errors are normally distributed with mean equal to zero.
- The errors are independent.
- The errors have a constant variance.
- The underlying population relationship is linear.
- If the residuals performance does not meet the requirement, we will need to rebuild the model by replacing the predictors with new ones, adding new predictors, building non-linear models, and so on.

Use Minitab to Perform Residuals Analysis

The residuals of the model are automatically saved in the data table as long as we check the box "Residuals" in the "Regression – Storage" window.

Step 1: Check whether residuals are normally distributed around the mean of zero.

1. Click Stat \rightarrow Basic Statistics \rightarrow Graphical Summary.

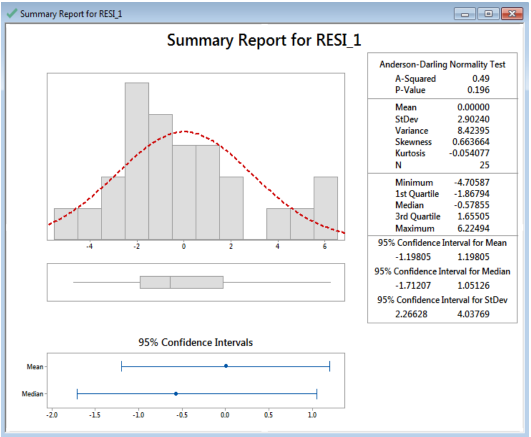
- 2. A new window named "Graphical Summary" pops up.
- 3. Select "RESI_1" as the "Variables."

We are selecting RESI_1 because we ran our regression two times, the first with all Exams which gave us our first set of residuals (RESI), the second model we ran without Exam1 and then we got residuals added to our data table and that column was automatically labeled RESI_1.

Graphical Summary	×
C1 EXAM1 C2 EXAM2 C3 EXAM3 C4 FINAL C5 RESI C6 DFIT C7 RESI_1 C8 DFIT_1 C9 PFITS C10 PSEFITS C10 PSEFITS C11 CLIM C12 CLIM_1 C13 PLIM C14 PLIM_1	Variables: RESI_1 By variables (optional): Confidence level: 95.0
Select	
Help	OK Cancel

Fig. 4.40 Graphical Summary dialog box with variable

- 4. Click "OK."
- 5. The histogram and analysis results are shown automatically in the new window.



The mean of residuals is -0.0000.

The Anderson–Darling test is used to test the normality. Since the p-value (0.196) is greater than the alpha level (0.05), we fail to reject the null hypothesis; the residuals are normally distributed.

- H₀: The residuals are normally distributed.
- H₁: The residuals are not normally distributed.

The residuals are normally distributed and the mean is near zero.

Step 2: If the data are in time order, run the IR chart to check whether residuals are independent.

- 1. Click Stat \rightarrow Control Charts \rightarrow Variable Charts for Individuals \rightarrow I-MR.
- 2. A new window named "Individuals Moving Range Chart" pops up.
- 3. Select "RESI_1" as "Variables"

Individuals-Moving Range	Chart		×
C1 EXAM1 C2 EXAM2 C3 EXAM3 C4 FINAL C5 RESI C6 DFIT C7 RESI_1 C8 DFIT_1 C9 PFITS C10 PSEFITS C10 PSEFITS C10 PSEFITS C11 CLIM_1 C12 CLIM_1 C13 PLIM C14 PLIM_1	Variables: RESI_1 Scale Multiple Graphs	Labels Data Options	I-MR Options
Select			
Help		<u>0</u> K	Cancel

Fig. 4.42 Individuals – Moving Range Chart dialog box with variable

4. Click "I-MR Options"

5. Click on "Tests" tab and select "Perform all tests for special causes"

Individuals-Moving Range Chart: Options	×
Parameters Estimate Limits Tests Stages Box-Cox Display Stora	age
Perform all tests for special causes	к
1 point > K standard deviations from center line	3
K points in a row on same side of center line	9
K points in a row, all increasing or all decreasing	6
K points in a row, alternating up and down	14
K out of K+1 points > 2 standard deviations from center line (same side)	2
K out of K+1 points > 1 standard deviation from center line (same side)	4
K points in a row within 1 standard deviation of center line (either side)	15
K points in a row > 1 standard deviation from center line (either side)	8
<u>QK</u> Can	cel

Fig. 4.43 Individuals - Moving Range Chart: Options

- 6. Click "OK." to close "I-MR" Options
- 7. Click OK
- 8. The control charts are shown automatically in the new window.

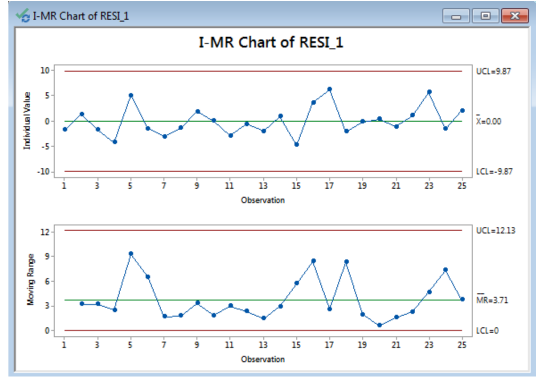


Fig. 4.44 Individual - Moving Range Chart output

If no data points are out of control in both the I-chart and MR chart, the residuals are independent of each other. If the residuals are not independent, it is possible that some important predictors are not included in the model. In this example, since the IR chart is in control, residuals are independent. The residuals are independent.

Step 3: Check whether residuals have equal variance across the predicted responses.

Create a scatterplot with Y being the residuals and the X being the fitted values.

- 1. Click Graph \rightarrow Scatterplot.
- 2. A new window named "Scatterplots" appears, click ok with Simple selected.
- 3. Select "RESI_1" as the "Y variables" and "PFITS" as the "X variables."

Scatterplot: Simple			×
C1 EXAM1 C2 EXAM2 C3 EXAM3 C4 FINAL C5 RESI C6 DFIT C7 RESI_1 C8 DFIT_1 C9 PFITS C10 PSEFITS C11 CLIM C12 CLIM_1 C13 PLIM_1 C14 PLIM_1 Select	Y variables X variables 1 RESI_1' PFITS 2 3 4 3 6 6 7		N
Help		QK Cance	

Fig. 4.45 Scatterplot: Simple dialog box with variables selected 4. Click "OK."

5. The scatterplot appears in a new window.

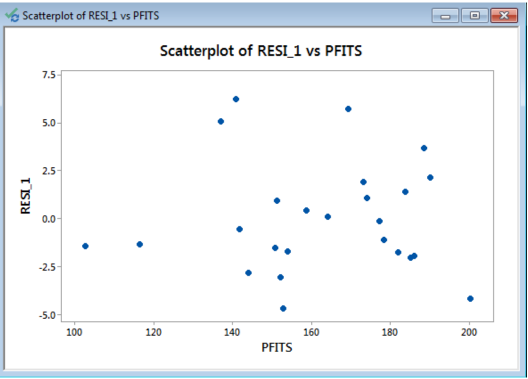


Fig. 4.46 Scatterplot output

We are looking for the pattern in which residuals spread out evenly around zero from the top to the bottom. The residuals are evenly distributed around zero; so, residuals pass the tests.

5.0 CONTROL PHASE

5.1 LEAN CONTROLS

5.1.1 CONTROL METHODS FOR 5S

What is 5S?

5S is a methodology that is applied in a Lean workplace. It is a systematic method to organize, order, clean, and standardize a workplace . . . and keep it that way! 5S is summarized in five Japanese words, all starting with the letter S:

- Seiri (sorting)
- Seiton (straightening)
- Seiso (shining)
- Seiketsu (standardizing)
- *Shisuke* (sustaining)

Originally developed in Japan, 5S is widely used to optimize the workplace to increase productivity and efficiency.

5S Goals

- 1. Reduced waste
- 2. Reduced cost
- 3. Establish a work environment that is:
 - Self-explaining
 - Self-ordering
 - Self-regulating
 - Self-improving.

Where there is/are no more:

- Wandering and/or searching
- Waiting or delays
- Secrets hiding spots for tools
- Obstacles or detours
- Extra pieces, parts, materials, etc.
- Injuries
- Waste

5S Benefits

By meeting the goals of 5S, the following benefits are achieved.

• Reduced changeovers

• Reduced defects

- Reduced waste
- Reduced delays
- Reduced injuries
- Reduced breakdowns
- Reduced complaints
- Reduced red ink

- Higher quality
- Lower costs
- Safer work environment
- Greater associate and equipment capacity

5S Systems Reported Results

Organizations that have implemented 5S systems have reported some very impressive results: reduction in waste, improved safety, reduced costs, and improved performance.

Cut in floor space:	60%
Cut in flow distance:	80%
Cut in accidents:	70%
Cut in rack storage:	68%
Cut in number of forklifts:	45%
Cut in machine changeover time:	62%
Cut in annual physical inventory time:	50%
Cut in classroom training requirements:	55%
Cut in nonconformance in assembly:	96%
Increase in test yields:	50%
Late deliveries:	0%
Increase in throughput:	15%

Sorting (Seiri)

- Go through all the tools, parts, equipment, supply, and material in the workplace.
- Categorize them into two major groups: needed and unneeded.
- Eliminate the unneeded items from the workplace. Dispose of or recycle those items.
- Keep the needed items and sort them in the order of priority. When in doubt . . . throw it out!

Straightening (Seiton)

Straightening in 5S is also called setting in order.

- Label each needed item.
- Store items at their best locations so that the workers can find them easily whenever they needed any item.
- Reduce the motion and time required to locate and obtain any item whenever it is needed.
- Promote an efficient work flow path.

• Use visual aids like the tool board image on this page to remind employees where things belong so the order is sustained.

Shining (Seiso)

Shining in 5S is also called *sweeping*.

- Clean the workplace thoroughly.
- Maintain the tidiness of the workplace.
- Make sure every item is located at the specific location where it should be.
- Create the ownership in the team to keep the work area clean and organized.

Standardizing (Seiketsu)

Standardize the workstation and the layout of tools, equipment, and parts.

Create identical workstations with a consistent way of storing the items at their specific locations so that workers can be moved around to any workstation any time and perform the same task.

Sustaining (Shisuke)

Sustaining in 5S is also called self-discipline.

- Create the culture in the team to follow the first four S's consistently.
- Avoid falling back to the old ways of cluttered and unorganized work environment.
- Keep the momentum of optimizing the workplace.
- Promote innovations of workplace improvement.
- Sustain the first fours S's using:
 - o 5S Maps
 - o 5S Schedules
 - 5S Job cycle charts
 - Integration of regular work duties
 - 5S Blitz schedules
 - Daily workplace scans

Simplified Summary of 5S

- 1. *Sort* "When in doubt, move it out"
- 2. *Set in Order*—Organize all necessary tools, parts, and components of production. Use visual ordering techniques wherever possible.
- 3. *Shine*—Clean machines and/or work areas. Set regular cleaning schedules and responsibilities.
- 4. *Standardize*—Solidify previous three steps, make 5S a regular part of the work environment and everyday life

5. *Sustain*—Audit, manage, and comply with established five-s guidelines for your business or facility

5.1.2 Kanban

What is Kanban?

The Japanese word *Kanban* means signboard. *Kanban system* is a "pull" production scheduling system to determine when to produce, what to produce, and how much to produce based on the demand. It was originally developed by Taiichi Ohno in order to reduce the waste in inventory and increase the speed of responding to the immediate demand.

Kanban system is a demand-driven system.

- The customer demand is the signal to trigger or pull the production.
- Products are made only to meet the immediate demand. When there is no demand, there is no production.

It is designed to minimize the in-process inventory and to have the right material with the right amount at the right location at the right time.

Principles of the Kanban System:

- Only produce products with exactly the same amount that customers consume.
- Only produce products when customers consume.

The production is driven by the *actual* demand from the customer side instead of the *forecasted* demand planned by the staff. The goal is to match production to demand.

Kanban Card

The *Kanban card* is the ticket or signal to authorize the production or movement of materials. It is the message of asking for more. The Kanban card is sent from the end customer up to the chain of production. Upon receiving of a Kanban card, the production station would start to produce goods. The Kanban card can be a physical card or an electronic signal.

Kanban System Example

The simplest example of a Kanban system is the supermarket operation.

- Customers visit the supermarkets and buy what they need.
- The checkout scanners send electronic Kanban cards to the local warehouse asking for more when the items are sold to customers.
- When the warehouse receives the Kanban cards, it starts to replenish the exact goods being sold.

It the warehouse prepares more than what Kanban cards require, the goods would become obsolete. If it prepares less, the supermarket would not have the goods available when customers need them.

Kanban System Benefits

- Minimize in-process inventory
- Free up space occupied by unnecessary inventory
- Prevent overproduction
- Improve responsiveness to dynamic demand
- Avoid the risk of inaccurate demand forecast
- Streamline the production flow
- Visualize the work flow

5.1.3 Рока-Үоке

What is Poka-Yoke?

The Japanese term *poka-yoke* means mistake-proofing. It is a mechanism to eliminate defects as early as possible in the process. It was originally developed by Shigeo Shingo and was initially called "baka-yoke" (fool-proofing). Poka-yoke is an extremely effective control method because it means to prevent, correct, or draw attention to a mistake as it occurs.

Two Types of Poka-Yoke

- Prevention
 - Preventing defects from occurring
 - \circ $\,$ Removing the possibility that an error could occur $\,$
 - Making the occurrence of an error impossible
- Detection
 - $\circ \quad \text{Detecting defects once they occur}$
 - Highlighting defects to draw workers' attention immediately
 - Correcting defects so that they would not reach the next stage

The best way to drive quality is to prevent defects before they occur. The next best solution is to detect defects as they occur so they can be corrected. Poka-yoke solutions can aim to either prevent or detect defects.

Three Methods of Poka-Yoke

Shigeo Shingo identified three methods of poka-yoke for detecting and preventing errors in a mass production system.

1. Contact Method—Use of shape, color, size, or any other physical attributes of the items

- 2. Constant Number Method—Use of a fixed number to make sure a certain number of motions are completed
- 3. Sequence Method—Use of a checklist to make sure all the prescribed process steps are followed in the right order

Poka-Yoke Devices

We are surrounded by poka-yoke devices daily. Prevention Devices (Example: The dishwasher does not start to run when the door is open). Detection Devices (Example: The car starts to beep when the passengers do not buckle their seatbelts)

Poka-yoke devices can be in any format that can quickly and effectively prevent or detect mistakes: visual, electrical, mechanical, procedural, human etc.

Other examples:

- Tether on a car's gas cap to keep driver from leaving it behind.
- Inability to remove the key from car ignition until transmission is in park.
- Safety bar on lawn mower must be engaged to start engine.
- Hole near the top of a bathroom sink to keep it from overflowing.
- Clothes dryer stops when the door is opened.

Steps to Apply Poka-Yoke

Step 1: Identify the process steps in need of mistake proofing.

Step 2: Use the 5-why's to analyze the possible mistakes or failures for the process step.

Step 3: Determine the type of poka-yoke: prevention or detection.

Step 4: Determine the method of poka-yoke: contact, constant number, or sequence.

Step 5: Pilot the poka-yoke approach and make any adjustments if needed.

Step 6: Implement poka-yoke in the operating process and maintain the performance.

5.2 STATISTICAL PROCESS CONTROL

5.2.1 DATA COLLECTION FOR SPC

What is SPC?

Statistical process control (SPC) is a statistical method to monitor the performance of a process using control charts in order to keep the process in statistical control. It can be used to distinguish between special cause variation and common cause variation in the process. SPC presents the voice of the process.

Common Cause Variation

Common cause variation (also called chance variation) is the inherent natural variation in any processes. It is the random background noise, which cannot be controlled or eliminated from the process. Its presence in the process is expected and acceptable due to its relatively small influence on the process. The goal of SPC is to distinguish between *common cause* variation and *special cause* variation to minimize tampering or inaction to real process changes.

Special Cause Variation

Special cause variation (also called assignable cause variation) is the unnatural variation in the process. It is the cause of process instability and leads to defects of the products or services. It is also the signal of unanticipated change (either positive or negative) in the process. It is possible to eliminate the special cause variation from the process.

Special cause variation is unnatural variation that has an assignable cause. In some cases, special cause variation is unwanted, but in other cases it can represent something desirable to replicate.

Process Stability

A process is *stable* when:

- There is not any special cause variation involved in the process
- The process is in statistical control
- The future performance of the process is predictable within certain limits
- The changes happening in the process are all due to random inherent variation
- There are not any trends, unnatural patterns, and outliers in the control chart of the process.

When a process performs predictably within certain limits, we say it is "in control." A process that is in control is stable.

SPC Benefits

- Statistical process control can be used in different phases of Six Sigma projects to:
- Understand the stability of a process
- Detect the special cause variation in the process
- Identify the statistical difference between two phases
- Eliminate or apply the unnatural change in the process
- Improve the quality and productivity.

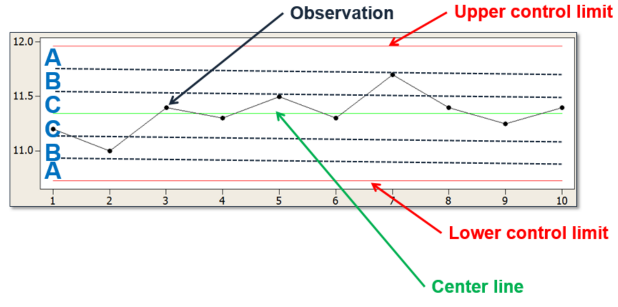
Control Charts

Control charts are graphical tools to present and analyze the process performance in statistical process control. Control charts are used to detect special cause variation and determine

whether the process is in statistical control (stable). They can differentiate *assignable* (special) sources of variation from *common* sources.

Variation solutions:

- Minimize the common cause variation
- Eliminate the special cause variation when it leads to unanticipated negative changes in the outcome
- Implement the special cause variation when it leads to unanticipated positive changes in the outcome.



Control Charts Elements

Fig. 5.1 Control Chart Elements

At its basic level, the control chart is a visual tool to illustrate when a process is stable or not. While there are other rules that will tell us whether or not a process is in control, the basic rule is to observe if the data points fall between the upper and lower control limits.

Control Charts Elements

Control charts can work for both continuous data and discrete or count data. Control limits are approximately three-sigma away from the process mean. A process is in statistical control when all the data points on the control charts fall within the control limits and have random patterns only. Otherwise, the process is out of control and we need to investigate the special cause variation in the process.

Possible Errors in SPC

Two types of errors can be made when interpreting and acting (or not acting) on a control chart.

- *Type I (False Positive)*—Tampering, acting to correct variation when you should not, because it is common cause variation.
- *Type II (False Negative)*—Not acting to correct variation when you should, because it is special cause variation.

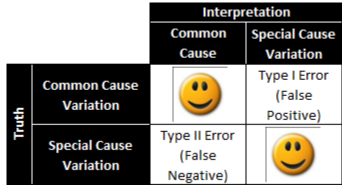


Fig. 5.2 Types of Errors

A control chart is a form of a hypothesis test where the null hypothesis is that the process is stable, and the alternative is that the process is unstable.

- Null Hypothesis (H₀): The process is stable (i.e., in statistical control).
- Alternative Hypothesis (H_a): The process is unstable (i.e., out of statistical control).

Type I Error

- False positive
- False alarm
- Considering true common cause variation as special cause variation

Type I errors waste resources spent on investigation.

Type II Error

- False negative
- Miss
- Considering true special cause variation as common cause variation

Type II errors neglect the need to investigate critical changes in the process.

Data Collection Considerations

To collect data for plotting control charts, we need to consider:

- 1. What is the measurement of interest?
- 2. Are the data discrete or continuous?

- 3. How many samples do we need?
- 4. How often do we sample?
- 5. Where do we sample?
- 6. What is the sampling strategy?
- 7. Do we use the raw data collected or transfer them to percentages, proportions, rates, etc.?

Subgroups and Rational Subgrouping

When sampling, we randomly select a group of items (i.e., a subgroup) from the population of interest. The *subgroup size* is the count of samples in a subgroup. It can be constant or variable. Depending on the subgroup sizes, we select different control charts accordingly.

Rational subgrouping is the basic sampling scheme in SPC. The goal of rational subgrouping is to maximize the likelihood of detecting special cause variation. In other words, the control limits should only reflect the variation *between* subgroups.

The numbers of subgroups, subgroup size, and frequency of sampling have great impact on the quality of control charts.

Sampling is randomly selecting a group of items from a population in which all items are produced under the conditions where only random effects are responsible for the observed variation. This maximizes the likelihood of detecting a special cause when it occurs.

Impact of Variation

The rational subgrouping strategy is designed to minimize the opportunity of having special cause variation *within* subgroups. If there is only random variation (background noise) within subgroups, all the special cause variation would be reflected between subgroups. It is easier to detect an out-of-control situation. Random variation is inherent in any process. We are more interested in identifying and taking actions on special cause variation.

Frequency of Sampling

The *frequency of sampling* in SPC depends on whether we have sufficient data to signal the changes in a process with reasonable time and costs. The more frequently we sample, the higher costs we incur. We need the subject matter experts' knowledge on the nature and characteristics of the process to make good decisions on sampling frequency.

5.2.2 I-MR CHART

I-MR Chart

The *I-MR chart* (also called individual-moving range chart or IR chart) is a popular control chart for continuous data with subgroup size equal to one.

- The I chart plots an individual observation as a data point.
- The MR chart plots the absolute value of the difference between two consecutive observations in individual charts as a data point.

If there are n data points in the I chart, there are n-1 data points in the MR chart. The I chart is valid only if the MR chart is in control. The underlying distribution of the I-MR chart is normal distribution.

I Chart Equations

I Chart (Individuals Chart) Data Point: \mathcal{X}_i

Center Line: $\frac{\sum_{i=1}^{n} x_i}{n}$

Control Limits:
$$\frac{\sum_{i=1}^{n} x_i}{n} \pm 2.66 \times \overline{MR}$$

Where: *n* is the number of observations.

MR-Chart Equations

MR Chart (Moving Range Chart) Data Point: $|x_{i+1} - x_i|$

Center Line: $\frac{\sum |x_{i+1} - x_i|}{n-1}$

Upper Control Limit:
$$3.267 \times \frac{\sum |x_{i+1} - x_i|}{n-1}$$

Lower Control Limit: 0

Where: *n* is the number of observations.

Use Minitab to Plot I-MR Charts



Data File: "IR" tab in "Sample Data.xlsx"

Steps to plot IR charts in Minitab:

1. Click Stat \rightarrow Control Charts \rightarrow Variable Charts for Individuals \rightarrow I-MR.

- 2. A new window named "Individuals Moving Range Chart" appears.
- 3. Select "Measurement" as the "Variables."

Ir	dividuals-Moving Range	Chart		— X
	C1 Measurement	<u>V</u> ariables:		
		Measurement		*
J				-
		<u>S</u> cale	Labels	
		Multiple Graphs	D <u>a</u> ta Options	I-MR Options
l	Select			
	Help		<u>O</u> K	Cancel

Fig. 5.3 Individuals-Moving Range Chart dialog box with variable

- 4. Click "I-MR Options" button and a new window "Individual Moving Range Chart Options" appears.
- 5. Click on the tab "Tests."
- 6. Select the item "Perform all tests for special causes" in the dropdown box.

Individuals-Moving Range Chart: Options							
Parameters Estimate Limits Tests Stages Box-Cox Display Store							
Perform all tests for special causes							
1 point > K standard deviations from center line							
K points in a row on same side of center line	9						
K points in a row, all increasing or all decreasing	6						
K points in a row, alternating up and down							
K out of K+1 points > 2 standard deviations from center line (same side)	2						
K out of K+1 points > 1 standard deviation from center line (same side)							
K points in a row within 1 standard deviation of center line (either side)	15						
K points in a row > 1 standard deviation from center line (either side)	8						
Help <u>Q</u> K Can	cel						

Fig. 5.4 Individuals-Moving Range Chart: Options Test Tab with selection

- 7. Click "OK" in the window "Individuals Moving Range Chart Options."
- 8. Click "OK."

9. The IR charts appear in the newly-generated window.

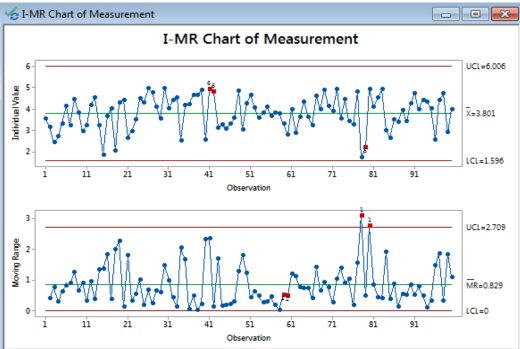


Fig. 5.5 Individuals-Moving Range Chart output

I-MR Charts Diagnosis

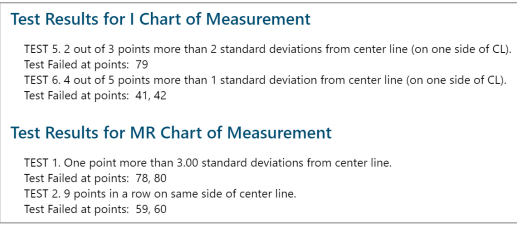


Fig. 5.6 I-MR Charts Diagnosis

I Chart (Individuals' Chart): Since the MR chart is out of control, the I chart is invalid.

MR Chart (Moving Range Chart): Two data points fall beyond the upper control limit. This indicates the MR chart is out of control (i.e., the variations between every two contiguous individual samples are not stable over time). We need to further investigate the process, identify the root causes that trigger the outliers, and correct them to bring the process back in control.

Xbar-R Chart

The *Xbar-R chart* is a control chart for continuous data with a constant subgroup size between two and ten.

- The Xbar chart plots the average of a subgroup as a data point.
- The R chart plots the difference between the highest and lowest values within a subgroup as a data point.

The Xbar chart monitors the process mean and the R chart monitors the variation within subgroups. The Xbar is valid only if the R chart is in control.

The underlying distribution of the Xbar-R chart is normal distribution.

Xbar Chart Equations

Xbar chart

Data Point:
$$\overline{X_i} = \frac{\sum_{j=1}^m x_{ij}}{m}$$

Center Line:
$$\overline{\overline{X}} = \frac{\sum_{i=1}^{k} \overline{X}_{i}}{k}$$

Control Limits: $\overline{\overline{X}} \pm A_2 \overline{R}$

Where:

- *m* is the subgroup size
- *k* is the number of subgroups.

A₂ is a constant depending on the subgroup size.

R Chart Equations

R chart (Rage Chart)
Data Point:
$$R_i = \underset{j \in [1,m]}{Max}(x_{ij}) - \underset{j \in [1,m]}{Min}(x_{ij})$$

Contact Line $\overline{R} = \sum_{i=1}^{k} R_i$

Center Line: $\overline{R} = \frac{i=1}{k}$

Upper Control Limit: $D_4 \times \overline{R}$

Lower Control Limit: $D_3 \times \overline{R}$

Where:

- *m* is the subgroup size and *k* is the number of subgroups
- D₃ and D₄ are constants depending on the subgroup size.

Use Minitab to Plot Xbar-R Charts

Data File: "Xbar-R" tab in "Sample Data.xlsx"

Steps to plot Xbar-R charts in Minitab:

- 1. Click Stat \rightarrow Control Charts \rightarrow Variable Charts for Subgroups \rightarrow Xbar-R.
- 2. A new window named "Xbar-R Chart" appears.
- 3. Click in the blank box right below "All observations for a chart are in one column" and the variables appear in the list box on the left.
- 4. Select the "Measurement" into the box below "All observations for a chart are in one column."

Xbar-R Chart			×
C1 Measurement C2 Subgroup ID	All observations for a d Measurement Subgroup sizes: 'Subg Scale Multiple Graphs		a number or ID column)
Select			
Help		<u>O</u> K	Cancel

Fig. 5.7 Xbar-R Chart with Measurement variable selection

- 5. Select the "Subgroup ID" as the "Subgroup size (enter a number or ID column).
- 6. Click "Xbar-R Options" button and a new window "Xbar-R Chart Options" appears.
- 7. Click on the tab "Tests."

8. Select the item "Perform all tests for special causes" in the dropdown menu.

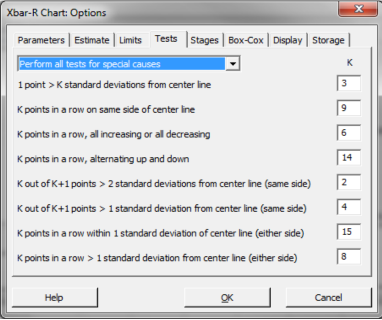


Fig. 5.8 Xbar-R Chart: Options Test Tab with test selection

- 9. Click "OK" in the window "Xbar-R Chart Options."
- 10. Click "OK."
- 11. The Xbar-R charts appear in the newly-generated window.

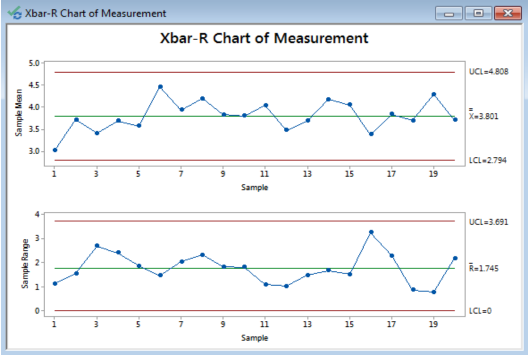


Fig. 5.9 Xbar-R Chart output

Xbar-R Charts Diagnosis

Xbar-R Charts:

Since the R chart is in control, the Xbar chart is valid. In both charts, there are not any data points failing any tests for special causes (i.e., all the data points fall between the control limits and spread around the center line with a random pattern). We conclude that the process is in control.

5.2.4 U CHART

The *U chart* is a type of control chart used to monitor discrete (count) data where the sample size is greater than one, typically the average number of defects per unit.

Defect vs. Defective

Remember the difference between defect and defective?

A *defect* of a unit is the unit's characteristic that does not meet the customers' requirements.

A *defective* is a unit that is not acceptable to the customers.

One defective might have multiple defects. One unit might have multiple defects but still be usable to the customers.

U Chart

The *U chart* is a control chart monitoring the average defects per unit. It plots the count of defects per unit of a subgroup as a data point. It considers the situation when the subgroup size of inspected units for which the defects would be counted is not constant. The underlying distribution of the U chart is Poisson distribution.

U Chart Equations

U chart Data Point: $u_i = \frac{x_i}{n_i}$ Center Line: $\overline{u} = \frac{\sum_{i=1}^k u_i}{k}$ Control Limits: $\overline{u} \pm 3 \times \sqrt{\frac{u}{n_i}}$

Where:

- n_i is the subgroup size for the ith subgroup
- *k* is the number of subgroups

• x_i is the number of defects in the ith subgroup.

Use Minitab to Plot a U Chart



Data File: "U" tab in "Sample Data.xlsx"

Steps to plot a U chart in Minitab:

- 1. Click Stat \rightarrow Control Charts \rightarrow Attributes Charts \rightarrow U.
- 2. A new window named "U Chart" appears.
- 3. Select "Count of Defects" as the "Variables."
- 4. Select "Count of Units Inspected" as the "Subgroup Sizes."

U Cha	rt			×
C1 C2 C3	Day Count of Defects Count of Units Inspe	Variables: 'Count of Defects'		A
		Subgroup sizes: Jf Un (enter a number or colum	its Inspected' nn containing the sizes)	*
		<u>S</u> cale	Labels	
	Select	Multiple Graphs	Data Options	U Chart Options
	Help		<u>0</u> K	Cancel

Fig. 5.10 U Chart dialog box with variable selections

- 5. Click the button "U Chart Options" to open a window named "U Chart Options".
- 6. Click the tab "Tests."

7. Select the item "Perform all tests for special causes" in the dropdown menu.

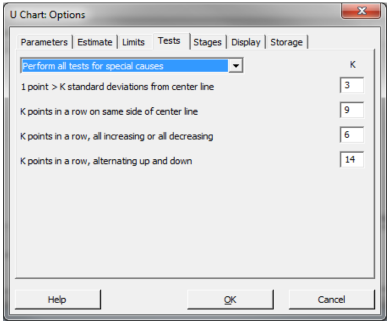


Fig. 5.11 U Chart: Options dialog box on Test Tab with test selection

- 8. Click "OK" in the window "U Chart Options."
- 9. Click "OK" in the window "U Chart."
- 10. The U chart appears in the newly-generated window.

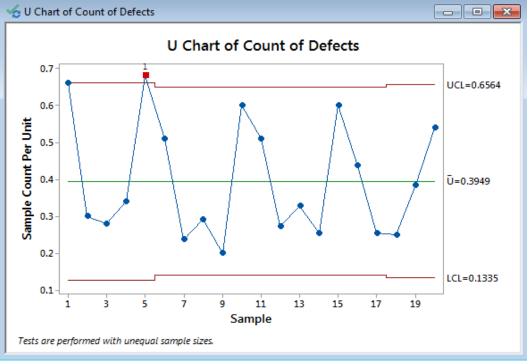


Fig. 5.12 U Chart output

U Chart Diagnosis

U Chart:

Since the sample sizes are not constant over time, the control limits are adjusted to different values accordingly. The data point circled above falls beyond the upper control limit. We conclude that the process is out of control. Further investigation is needed to determine the special causes that triggered the unnatural pattern of the process.

5.2.5 P CHART

P Chart

The *P* chart is a control chart monitoring the percentages of defectives. The P chart plots the percentage of defectives in one subgroup as a data point. It considers the situation when the subgroup size of inspected units is not constant. The underlying distribution of the P chart is binomial distribution.

P Chart Equations

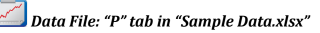
P chart

Data Point: $p_i = \frac{x_i}{n_i}$ Center Line: $\overline{p} = \frac{\sum_{i=1}^k x_i}{\sum_{i=1}^k n_i}$ Control Limits: $\overline{p} \pm 3\sqrt{\frac{\overline{p}(1-\overline{p})}{n_i}}$

Where:

- *n_i* is the subgroup size for the ith subgroup
- *k* is the number of subgroups
- x_i is the number of defectives in the ith subgroup.

Use Minitab to Plot a P Chart



Steps to plot a P chart in Minitab:

- 1. Click Stat \rightarrow Control Charts \rightarrow Attributes Charts \rightarrow P.
- 2. A new window named "P Chart" appears.
- 3. Select "Fail" as the "Variables."

4. Select "N" as the "Subgroup Sizes."

P Chart				×
C1 Day C2 Fail C3 Pass C4 N		Variables: Fail Subgroup sizes: N (enter a number or colu	mn containing the sizes)	~
		<u>S</u> cale Multiple Graphs	Labels Data Options	P Chart Options
s	elect			
Help			<u>0</u> K	Cancel

Fig. 5.13 P Chart dialog box with variable selections

- 5. Click the button "P Chart Options" to open a window named "P Chart Options".
- 6. Click the tab "Tests."
- 7. Select the item "Perform all tests for special causes" in the dropdown menu.

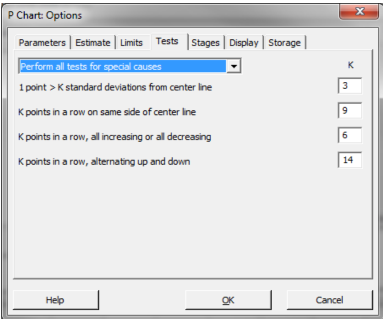


Fig. 5.14 P Chart: Options dialog box Test Tab with test selections

- 8. Click "OK" in the window "P Chart Options."
- 9. Click "OK."

10. The P chart appears in the newly-generated window.

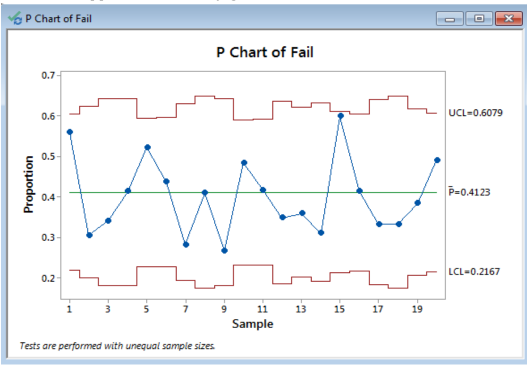


Fig. 5.15 P Chart output

P Chart Diagnosis

P Chart:

Since the sample sizes are not constant over time, the control limits are adjusted to different values accordingly. All the data points fall within the control limits and spread randomly around the mean. We conclude that the process is in control.

5.2.6 NP CHART

NP Chart

The *NP chart* is a control chart monitoring the count of defectives. It plots the number of defectives in one subgroup as a data point. The subgroup size of the NP chart is constant. The underlying distribution of the NP chart is binomial distribution.

NP Chart Equations

NP chart

Data Point: \mathcal{X}_i

Center Line: $n\overline{p} = \frac{\sum_{i=1}^{n} x_i}{k}$

Control Limits: $n\overline{p} \pm 3 \times \sqrt{n\overline{p}(1-\overline{p})}$

Where:

- *n* is the constant subgroup size
- *k* is the number of subgroups
- x_i is the number of defectives in the ith subgroup.

Use Minitab to Plot an NP Chart

Data File: "NP" tab in "Sample Data.xlsx"

Steps to plot a NP chart in Minitab:

- 1. Click Stat \rightarrow Control Charts \rightarrow Attributes Charts \rightarrow NP.
- 2. A new window named "NP Chart" appears.
- 3. Select "Fail" as the "Variables."
- 4. Select "N" as the "Subgroup Sizes."

ſ	NP Chart			— ×
	C1 Day	<u>V</u> ariables:		
	C2 Fail C3 Pass C4 N	Fail		~
	CT N			~
		Subgroup sizes: N		
		(enter a number or colu	mn containing the sizes)
		Scale	Labels	
		<u></u>	Labels	
I		Multiple Graphs	Data Options	NP Chart Options
	Select			
	Help		<u>O</u> K	Cancel

Fig. 5.16 NP Chart dialog box with variable selections

- 5. Click the button "NP Chart Options" to open a window named "NP Chart Options".
- 6. Click the tab "Tests."

7. Select the item "Perform all tests for special causes" in the dropdown menu.

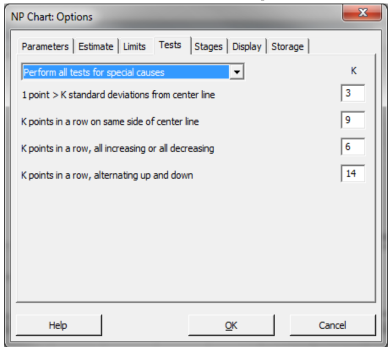


Fig. 5.17 NP Chart: Options dialog box Test tab with test selections

- 8. Click "OK" in the window "NP Chart Options."
- 9. Click "OK."
- 10. The NP chart appears in the newly-generated window.

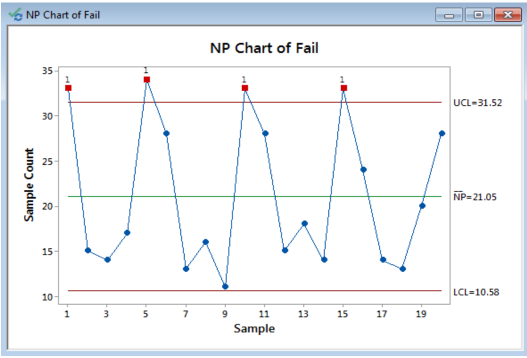


Fig. 5.18 NP Chart output

NP Chart Diagnosis

Four data points fall beyond the upper control limit. We conclude that the NP chart is out of control. Further investigation is needed to determine the special causes that triggered the unnatural pattern of the process.

5.2.7 X-S CHART

X-S Chart

The *X-S chart* (also called Xbar-S chart) is a control chart for continuous data with a constant subgroup size greater than ten. The Xbar chart plots the average of a subgroup as a data point. The S chart plots the standard deviation within a subgroup as a data point.

The Xbar chart monitors the process mean and the S chart monitors the variability within subgroups. The Xbar is valid only if the S chart is in control. The underlying distribution of the Xbar-S chart is normal distribution.

Xbar Chart Equations

Xbar Chart

Data Point: $\overline{X_i} = \frac{\sum_{j=1}^m x_{ij}}{m}$

Center Line:
$$\overline{\overline{X}} = \frac{\sum_{i=1}^{k} \overline{X}_{i}}{k}$$

Control Limits: $\overline{\overline{X}} \pm A_3 \overline{s}$

Where:

- *m* is the subgroup size
- *k* is the number of subgroups
- A₃ is a constant depending on the subgroup size.

S Chart Equations

S chart

Data Point:
$$s_i = \sqrt{\frac{\sum_{j=1}^{m} (x_{ij} - \bar{x}_i)^2}{m - 1}}$$

Center Line: $\bar{s} = \frac{\sum_{i=1}^{k} s_i}{k}$

Upper Control Limit: $B_4 \times s$

Lower Control Limit: $B_3 \times \overline{s}$

Where:

- *m* is the subgroup size
- *k* is the number of subgroups
- B₃ and B₄ are constants depending on the subgroup size.

Use Minitab to Plot Xbar-S Charts

Data File: "Xbar-S" tab in "Sample Data.xlsx"

Steps to plot Xbar-S charts in Minitab:

- 1. Click Stat \rightarrow Control Charts \rightarrow Variables Charts for Subgroups \rightarrow Xbar-S.
- 2. A new window named "Xbar-S Chart" appears.
- 3. Select "Measurement" by clicking the box below "All observations for a chart are in one column"
- 4. Select "Subgroup ID" as the "Subgroup Sizes."

3	Xbar-S	S Chart	×
	C1 C2	Measurement Subgroup ID	All observations for a chart are in one column:
			Measurement
			<u>S</u> cale <u>L</u> abels <u>M</u> ultiple Graphs <u>Da</u> ta Options
		Select	
		Help	<u>OK</u> Cancel

Fig. 5.19 Xbar-S Chart dialog box with variable selections

- 5. Click the button "Xbar-S Options" to open a window named "Xbar-S Chart Options".
- 6. Click the tab "Tests."

7. Select the item "Perform all tests for special causes" in the dropdown menu.

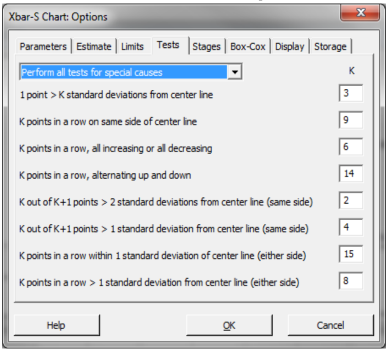


Fig. 5.20 Xbar-S Chart: Options dialog box Test tab with test selection

- 8. Click "OK" in the window "Xbar-S Chart Options."
- 9. Click "OK".
- 10. The Xbar-S charts appear in the newly-generated window.



Fig. 5.21 Xbar-S Chart output

Xbar-S Charts Diagnosis

Xbar-S Charts:

Since the S chart is in control, the Xbar chart is valid. In both charts, there are not any data points failing any tests for special causes (i.e., all the data points fall between the control limits and spread around the center line with a random pattern). We conclude that the process is in control.

5.2.8 CUMSUM CHART

CumSum Chart

The *CumSum chart* (also called cumulative sum control chart or CUSUM chart) is a control chart of monitoring the cumulative sum of the subgroup mean deviations from the process target. It detects the shift of the process mean from the process target over time. The underlying distribution of the CumSum chart is normal distribution.

There are two types of CumSum charts:

- 1. One two-sided CumSum charts
- 2. Two one-sided CumSum charts

Two-Sided CumSum

In two-sided CumSum charts, we use a V-mask to identify out-of-control data points. The Vmask is a transparent overlay shape of a horizontal "V" applied on the top of the CumSum chart. Its origin point is placed on the last data point of the CumSum chart and its center line is horizontal. If all of the data points stay inside the V-mask, we consider the process is in statistical control.

V-Mask

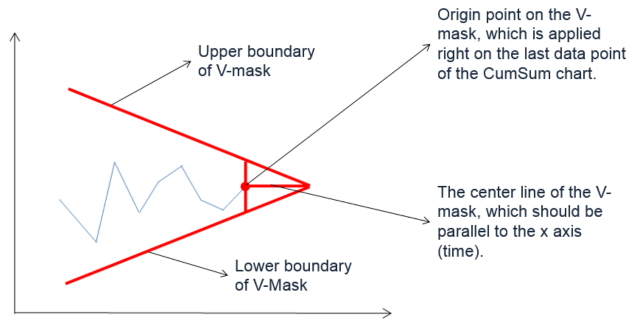


Fig. 5.22 V-Mask Explanation

The origin of the V-mask is the most recent data point, and the arms extend backwards. The process is out of control if any of the data points fall above or below the limits.

Two-Sided CumSum Equations

Two-Sided CumSum

Data Point:
$$\begin{cases} c_i = c_{i-1} + (\bar{x}_i - T) & i > 0\\ c_i = 0 & i = 0 \end{cases}$$

V-Mask Slope: $k \frac{\sigma}{\sqrt{m}}$

V-Mask Width at the Origin Point: $2h \frac{\sigma}{\sqrt{m}}$

Where:

- \overline{x}_i is the mean of the ith subgroup
- *T* is the process target
- σ is the estimation of process standard deviation
- *m* is the subgroup size.

One-Sided CumSum

We can also use two one-sided CumSum charts to detect the shift of the process mean from the process target. The upper one-sided CumSum detects the upward shifts of the process mean. The lower one-sided CumSum detects the downward shifts of the process mean.

One-Sided CumSum Equations

 $\begin{array}{ll} \textit{One-Sided CumSum} \\ \textit{Data Point:} \begin{cases} c_i = c_{i\text{-}1} + (\overline{x}_i\text{-}T) & i > 0 \\ c_i = 0 & i = 0 \end{cases} \end{array}$

Center Line: T

Upper Control Limit: $c_i^+ = \max\left(0, \overline{x}_i - (T+k) + c_{i-1}^+\right)$

Lower Control Limit: $c_i^- = \max\left(0, (T-k) - \bar{x}_i + c_{i-1}^-\right)$

Where:

- \overline{x}_i is the mean of the ith subgroup
- *T* is the process target
- *k* is the slope of the lower boundary of the V-mask.

Use Minitab to Plot a CumSum Chart

Data File: "CumSum" tab in "Sample Data.xlsx"

Steps in Minitab to plot CumSum charts:

- 1. Click Stat \rightarrow Control Charts \rightarrow Time Weighted Charts \rightarrow CUSUM.
- 2. A new window named "CUSUM Chart" appears.
- 3. Select "weight" in the box below "All observations for a chart are in one column."
- 4. Select "hour" as the "Subgroup sizes."

5. Enter "8.1" as "Target."

CUSUM Chart	
	All observations for a chart are in one column: weight Subgroup sizes: hour (enter a number or ID column) Target: 8.1
Select	Multiple Graphs Data Options CUSUM Options
Help	OK Cancel

Fig. 5.23 CUSUM Chart dialog box with variables selected

- 6. Click the button "CUSUM Options" to open a window named "CUSUM Chart Options".
- 7. Click the tab "Plan/Type"
- 8. Select the radio button "V-mask"

CUSUM Chart: Options	
Parameters Estimate Plan/Type Stages Box-C Type of CUSUM Iabular Iabular Image: Imag	CUSUM Plan
	Cancel

*Fig. 5.24 CUSUM Chart: Options dialog box with selections*9. Click "OK" in the window "CUSUM Chart Options."

10. Click "OK."

11. The CUSUM chart appears in the newly-generated window.

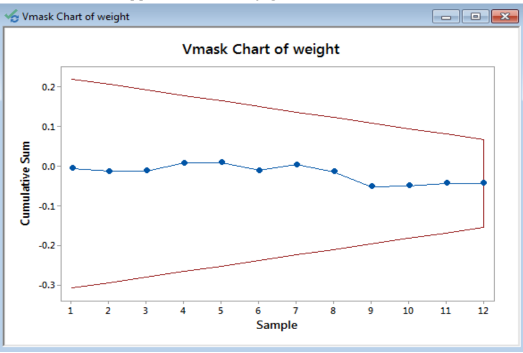


Fig. 5.25 CUSUM output

The process is in control since all of the data points fall between the two arms of the V-mask.

5.2.9 EWMA CHART

EWMA Chart

The *EWMA chart* (Exponentially-Weighted Moving Average Chart) is a control chart monitoring the exponentially-weighted average of previous and present subgroup means. The more recent data get more weight than older data. It detects the shift of the process mean from the process target over time. The underlying distribution of the EWMA chart is normal distribution.

EWMA Chart Equations

EWMA Chart Data Point: $z_i = \lambda x_i + (1 - \lambda) z_{i-1}$ where $0 < \lambda < 1$

Center Line: \overline{X}

Control Limits: $\overline{X} \pm k \cdot \frac{s}{\sqrt{n}} \sqrt{\left(\frac{\lambda}{2-\lambda}\right) \left[1 - (1-\lambda)^{2i}\right]}$

Where:

- \overline{x}_i is the mean of the ith subgroup
- λ and k are user-defined parameters to calculate the EWMA data points and the control limits.

Use Minitab to Plot an EWMA Chart

Data File: "EWMA" tab in "Sample Data.xlsx"

Steps in Minitab to plot EWMA charts:

- 1. Click Stat \rightarrow Control Charts \rightarrow Time Weighted Charts \rightarrow EWMA.
- 2. A new window named "EWMA Chart" appears.
- 3. Select "Gap" in the box below "All observations for a chart are in one column."
- 4. Select "Sample" as the "Subgroup sizes."
- 5. Enter "0.5" as "Weight of EWMA."

EWMA Chart	
	All observations for a chart are in one column:
	Subgroup sizes: Sample (enter a number or ID column) Weight of EWMA: 0.5
	Scale Labels Multiple Graphs Data Options
Select	
Help	<u>Q</u> K Cancel

Fig. 5.26 EWMA Chart dialog box with variable selections

6. Click "OK."

7. The EWMA chart appears in the newly-generated window.

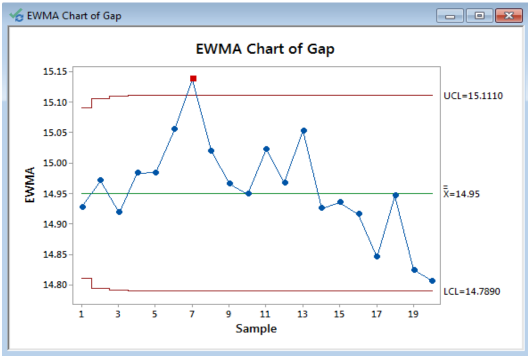


Fig. 5.27 EWMA Chart output

One data point falls beyond the upper control limit and we conclude that the process is out of control. Further investigation is needed to discover the root cause for the outlier.

5.2.10 CONTROL METHODS

Control Methods

The fundamental reason for the Control phase is to sustain improvements made by the project and ensure that the process remains stable with minimal variation. There are many control methods available to keep the process stable and minimize the variation. The most common control methods are:

- SPC (statistical process control)—using control charts to monitor process quality over time
- 5S method—to keep the workplace organized and efficient
- Kanban—for visual inventory management
- Poka-Yoke (mistake proofing)

SPC

SPC (Statistical Process Control) is a quantitative control method to monitor the stability of the process performance by identifying the special cause variation in the process. It uses control charts to detect the unanticipated changes in the process.

Which control chart to use depends on:

- Whether the data are continuous or discrete
- How large the subgroup size is
- Whether the subgroup size is constant
- Whether we are interested in measuring defects or defectives
- Whether we are interested in detecting the shifts in the process mean

5S

The *5S* methodology is a systematic approach of cleaning and organizing the workplace through:

- 1. Seiri (sorting)
- 2. Seiton (straightening)
- 3. Seiso (shining)
- 4. Seiketsu (standardizing)
- 5. Shisuke (sustaining)

In other words, 5S describes how to organize a workspace for efficiency and effectiveness by identifying and sorting tools and materials used for a process, and maintaining and sustaining the order of the area and items.

Kanban

A *Kanban* system is a demand-driven system. The customer demand is the signal to trigger or pull the production. Products are made only to meet the immediate demand. When there is no demand, there is no production. Kanban was designed to reduce the waste in inventory and increase the speed of responding to immediate demand.

Poka-Yoke

Poka-yoke means "mistake proofing" and is a mechanism to eliminate the defects as early as possible in the process.

Contact Method The contact method uses physical characteristics (e.g., shape, color, size) to identify defects.

Constant Number Method

The constant number method alerts an operator about a defect when a certain number of motions are not completed.

Sequence Method

The sequence method (or motion-step) identifies a defect when the steps of a process are not performed correctly or in the right order.

Control Chart Calculations Summary

Chart	Center Line	Control Limits	σ _x
I Chart	$\frac{\sum_{i=1}^{n} x_i}{n}$	$\frac{\sum_{i=1}^{n} x_i}{n} \pm 3 \times \frac{\overline{MR}}{d_2}$	$\frac{MR}{d_{1}}$
MR Chart	$\frac{n}{MR} = \frac{\sum_{i=1}^{n-1} \mathbf{x}_{i,1} - \mathbf{x}_i }{n-1}$	$UCL = D_4 \times \overline{MR}$ $LCL = D_3 \times \overline{MR}$	
Xbar Chart (Xbar-R)	$\overline{\overline{X}} = \frac{\sum_{i=1}^{k} \overline{X}_{i}}{k}$	$\overline{\overline{X}} \pm A_2 \overline{R}$	$\frac{R}{d_{z}}$
Xbar Chart (Xbar-S)	$\overline{\overline{X}} = \frac{\sum_{i=1}^{k} \overline{X}_{i}}{k}$	$\overline{\overline{X}} \pm A_3 \overline{s}$	<u></u>
R Chart	$\overline{R} = \frac{\sum_{i=1}^{k} R_i}{k}$	$UCL = D_4 \times \overline{R}$ $LCL = D_3 \times \overline{R}$	
S Chart	$\frac{1}{s} = \frac{\sum_{i=1}^{k} s_i}{k}$	$UCL = B_4 \times \overline{s}$ $LCL = B_3 \times \overline{s}$	
U Chart	$\frac{1}{u} = \frac{\sum_{i=1}^{k} u_i}{k}$	$\overline{u} \pm 3 \times \sqrt{\frac{\overline{u}}{n_i}}$	$\sqrt{\frac{u}{n_i}}$
P Chart	$\frac{\sum_{p=1}^{n} x_{1}}{\sum_{p=1}^{n} x_{p}}$	$\overline{p} \pm 3 \sqrt{\frac{\overline{p}(1-\overline{p})}{n_i}}$	$\sqrt{\frac{p(1-p)}{n_i}}$
NP Chart	$n\overline{p} = \frac{\sum_{i=1}^{k} x_i}{k}$	$n\overline{p}\pm 3 \times \sqrt{n\overline{p}(1-\overline{p})}$	$\sqrt{n\overline{p}(1-\overline{p})}$

Table 5.1 Control Chart Calculations

This is a helpful reference page to understand how the parameters of various control charts are calculated.

Subgroup Size	A2	A3	B3	B4	c4	d2	D3	D4
2	1.88	2.659	-	3.267	0.7979	1.128	-	3.267
3	1.023	1.954	-	2.568	0.8862	1.693	-	2.574
4	0.729	1.628	-	2.266	0.9213	2.059	-	2.282
5	0.577	1.427	-	2.089	0.94	2.326	-	2.114
6	0.483	1.287	0.03	1.97	0.9515	2.534	-	2.004
7	0.419	1.182	0.118	1.882	0.9594	2.704	0.076	1.924
8	0.373	1.099	0.185	1.815	0.965	2.847	0.136	1.864
9	0.337	1.032	0.239	1.761	0.9693	2.97	0.184	1.816
10	0.308	0.975	0.284	1.716	0.9727	3.078	0.223	1.777
15	0.223	0.789	0.428	1.572	0.9823	3.472	0.347	1.653
25	0.153	0.606	0.565	1.435	0.9896	3.931	0.459	1.541

Control Chart Constants

Table 5.2 Control Chart Constants

This is another helpful reference page. Many control charts use some of these constants in the calculation of parameters. They vary depending on the subgroup size.

Unnatural Patterns

If there are *unnatural patterns* in the control chart of a process, we consider the process out of statistical control. Control charts are designed to help us decipher special cause variation from common cause variation. Typical unnatural patterns in control charts are:

- Outliers (data points that fall outside of the upper or lower control limits)
- Trending
- Cycling
- Auto-correlative
- Mixture

A process is *in control* if all the data points on the control chart are randomly spread out within the control limits.

Western Electric Rules

Western Electric Rules are the most popular decision rules to detect unnatural patterns in the control charts. They are a group of tests for special causes in a process. The Western Electric Rules were developed by the manufacturing division of the Western Electric Company in 1956.

The area between the upper and lower control limits is separated into six subzones defined by the number of standard deviations from the center line.

- Zone A: between two and three standard deviations from the center line
- Zone B: between one and two standard deviations from the center line
- Zone C: within one standard deviation from the center line.

Western Electric Rules

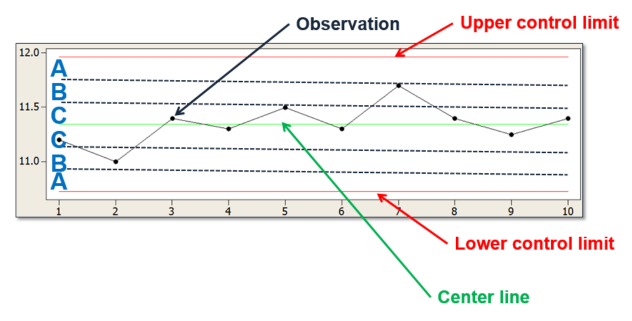


Fig. 5.28 Western Electric Rules

- Zone A: three sigma zone
- Zone B: two sigma zone
- Zone C: one sigma zone

If a data point falls onto the dividing line of two consecutive zones, the point belongs to the outer zone.

Test 1: One point more than three standard deviations from the center line (i.e., one point beyond zone A). Such a point is typically known as an outlier.

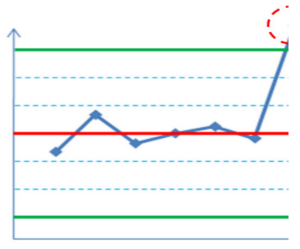


Fig. 5.29 Western Electric Test 1

Test 2: Nine points in a row on the same side of the center line.

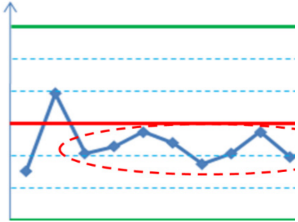


Fig. 5.30 Western Electric Test 2

Test 2 identifies situations where the process mean has temporarily shifts in the process.

Test 3: Six points in a row steadily increasing or steadily decreasing. Test 3 identifies significant trends in performance.

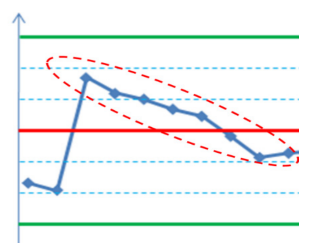


Fig. 5.31 Western Electric Test 3

Test 4: Fourteen points in a row alternating up and down. Test 4 indicates a cycle.

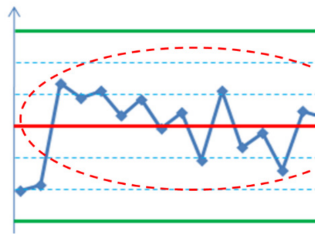


Fig. 5.32 Western Electric Test 4

Test 5: Two out of three points in a row at least two standard deviations from the center line (in zone A or beyond) on the same side of the center line

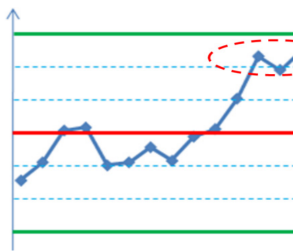


Fig. 5.33 Western Electric Test 5

Test 5 is similar to test 2, but represents a more acute shift in the process mean.

Test 6: Four out of five points in a row at least one standard deviation from the center line (in zone B or beyond) on the same side of the center line

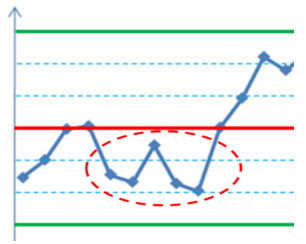


Fig. 5.34 Western Electric Test 6

Test 6 is similar to test 2, but represents a more acute shift in the process mean.

Test 7: Fifteen points in a row within one standard deviation from the center line (in zone C) on either side of the center line

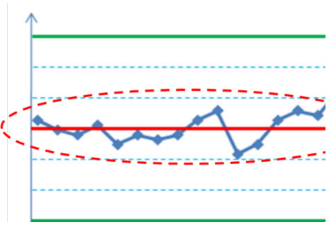
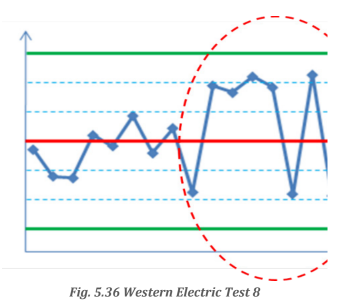


Fig. 5.35 Western Electric Test 7

Test 7 indicates a change in the level of variation around the process mean.

Test 8: Eight points in a row beyond one standard deviation from the center line (beyond zone C) on either side of the center line



While test 7 indicates a narrowing of variation, test 8 indicates an increase in variation.

Next Steps

If no data points fail any tests for special causes (Western Electric rules), the process is in *statistical control*. If any data point fails any tests for special causes, the process is *unstable* and we will need to investigate the observation thoroughly to discover and take actions on the special causes leading to the changes. Special cause variation must be addressed and the process must be stable before process capability analysis can be done. Process stability is the prerequisite of process capability analysis.

5.2.12 SUBGROUPS AND SAMPLING

Subgroups

Rational subgrouping is the basic sampling scheme in SPC (Statistical Process Control). When sampling, we randomly select a group (i.e., a subgroup) of items from the population of interest. The subgroup size is the count of samples in a subgroup. It can be constant or variable. Depending on the subgroup sizes, we select different control charts accordingly.

Impact of Variation

The *rational subgrouping* strategy is designed to minimize the opportunity of having special cause variation within subgroups, making it easier to identify special cause between subgroups. If there is only random variation (background noise) within subgroups, all the special cause variation would be reflected between subgroups. It is easier to detect the out-of-control situation. Random variation is inherent and indelible in the process. We are more interested in identifying and taking actions on special cause variation.

Frequency of Sampling

The *frequency of sampling* in SPC depends on whether we have sufficient data to signal the changes in a process with reasonable time and costs. The more frequently we sample, the higher the costs sampling may trigger. We need the subject matter experts' knowledge on the nature and characteristics of the process to make good decisions on sampling frequency.

5.3 SIX SIGMA CONTROL PLANS

5.3.1 Cost Benefit Analysis

What is Cost-Benefit Analysis?

For a project to be feasible, the benefits must outweigh the costs. The *cost-benefit analysis* is a systematic method to assess and compare the financial costs and benefits of multiple scenarios in order to make sound economic decisions.

A cost-benefit analysis is recommended to be done at the beginning of the project based on estimations of the experts from the finance team in order to determine whether the project is financially feasible. It is recommended to update the cost-benefit analysis at each DMAIC phase of the project.

Why Cost-Benefit Analysis?

In the Define phase of the project, the cost-benefit analysis helps us understand the financial feasibility of the project. In the middle phases of the project, updating and reviewing the costbenefit analysis helps us compare potential solutions and make robust data-driven decisions. In the Control phase of the project, the cost-benefit analysis helps us track the project's profitability.

Return on Investment

A common financial measure of a project's impact to profitability, *return on investment* (also called ROI, rate of return, or ROR) is the ratio of the net financial benefits (either gain or loss) of a project or investment to the financial costs.

$$ROI = \frac{TotalNetBenefits}{TotalCosts} \times 100\%$$

Where:

$$TotalNetBenefits = TotalBenefits - TotalCosts$$

Return on Investment (ROI)

The return on investment is used to evaluate the financial feasibility and profitability of a project or investment.

- If ROI < 0, the investment is not financially viable.
- If ROI = 0, the investment has neither gain nor loss.
- If ROI > 0, the investment has financial gains.

The higher the ROI, the more profitable the project. Any value over 100 percent means the project has a return greater than the cost, and the higher the ROI, the more profitable the project.

Net Present Value (NPV)

The *net present value* (also called NPV, net present worth, or NPW) is the total present value of the cash flows calculated using a discount rate. The NPV accounts for the time value of money.

$$NPV = \frac{NetCashFlow_t}{(1+r)^t}$$

Where:

- *NetCashFlow*_t is the net cash flow happening at time t
- *r* is the discount rate
- *t* is the time of the cash flow.

Often projects do not have immediate returns and it takes time to recover the costs incurred during the project.

Cost Estimation

There are many types of costs that can be incurred during a project. Some are one-time investments, such as equipment, consulting, or other assets, while others are incremental costs to be incurred going forward.

Examples of costs triggered by the project:

- Administration
- Asset
- Equipment
- Material
- Delivery
- Real estate
- Labor
- Training
- Consulting

Benefits Estimation

Benefits can take many forms: additional revenue, reduced wasted, operational resource costs, productivity improvements, or market share increases. Other types of benefits that are harder to qualify include customer satisfaction and associate satisfaction.

Examples of benefits generated by the project:

- Direct revenue increase
- Waste reduction
- Operation cost reduction
- Quality and productivity improvement
- Market share increase
- Cost avoidance
- Customer satisfaction improvement
- Associate satisfaction improvement

Challenges in Cost and Benefit Estimation

Different analysts might come up with different cost and benefit estimations due to their subjectivity in determining:

- The discount rate
- The time length of the project and its impact
- Potential costs of the project
- The tangible/intangible benefits of the project
- The specific contribution of the project to the relevant financial gains/loss

5.3.2 ELEMENTS OF CONTROL PLANS

Control Plans

A key component of a solid DMAIC project, the *control plan* ensures that the changes introduced by a Six Sigma project are sustained over time.

Benefits of the Control phase:

- Methodical roll-out of changes including standardization of processes and work procedures
- Ensure compliance with changes through methods like auditing and corrective actions
- Transfer solutions and learning across the enterprise
- Plan and communicate standardized work procedures
- Coordinate ongoing team and individual involvement
- Standardize data collection and procedures
- Measure process performance, stability, and capability
- Plan actions that mitigate possible out-of-control conditions

• Sustain changes over time

The Control phase ensures new processes and procedures are standardized and compliance is assured.

What is a Control Plan?

A *control plan* is a management planning tool to identify, describe, and monitor the process performance metrics in order to meet the customer specifications steadily. It proposes the plan of monitoring the stability and capability of inputs and outputs of critical process steps in the Control phase of a project. It covers the data collection plan of gathering the process performance measurements. Control plans are the most overlooked element of most projects. It is critical that a good solution be solidified with a great control plan!

Control Plan Elements

There are many possible elements within a control plan. All of these need to be agreed to and owned by the process owner—the person who is responsible for the process performance.

Control Plan—The clear and concise summary document that details key process steps, CTQs metrics, measurements, and corrective actions.

Standard Operating Procedures (SOPs) —Supporting documentation showing the "who does what, when, and how" in completing the tasks.

Communication Plan—Document outlining messages to be delivered and the target audience.

Training Plan—Document outlining the necessary training for employees to successfully perform new processes and procedures.

Audit Checklists—Document that provides auditors with the audit questions they need to ask.

Corrective Actions—Activities that need to be conducted when an audit fails.

Control Plan

The control plan identifies critical process steps that have significant impact on the products or services and the appropriate control mechanisms. It includes measurement systems that monitor and help manage key process performance measures. Specified limits and targets for these performance metrics should be clearly defined and communicated. Sampling plans should be used to collect the data for the measurements.

Lean Six Sigma Control Plan

Process: Custome Stakehol Business	der:					Preparer: Email: Phone: Owner:	Ξ				Reference No:	of
Process	Process Step	CTQ/Metric	CTQ / Metric Equation	Specifi Requir		Measurement Method	Sample Size	Measure Frequency		Link or Report Name	Corrective Action	Responsible for Action
				LSL	USL					Name		
3		8		8			8	2	<i>23</i>	2		

Fig. 5.37 Control Plan Key Process and Critical Measurements

Business:			
Process	Process Step	CTQ/Metric	CTQ / Metric Equation

Fig. 5.38 Process and Metrics portion of Control Plan

- Key processes and process steps are identified.
- Critical information regarding key measurements is documented and clarified.

			Owner:			
CTQ / Metric Equation		ication/ rement	Measurement Method	Sample Size	Measure Frequency	Responsible for Metric
	LSL	USL				
-		- Carl	and the second			Marel Land

Fig. 5.39 Measurements portion of Control Plan

- Measurements are clearly defined with equations.
- Customer specifications are declared.
- Other key measurement information is documented: sample size, measurement frequency, people responsible for the measurement, etc.



Fig. 5.40 Corrective Actions and Responsible Parties Portion of Control Plan

- Where will this measurement or report be found? Good control plans provide linking information or other report reference information.
- Control plans identify the mitigating action or corrective actions required in the event the measurement falls out of spec or control. Responsible parties are also declared.

Not only does a control plan spell out what is measured and what the limits are around the measurement, but it also details actions that must be taken when out-of-control or out-of-spec conditions occur.

Standard Operating Procedures (SOPs)

Standard Operating Procedures (SOPs) are documents that focus on process steps, activities, and specific tasks required to complete an operation. SOPs should not be much more than two to four pages and should be written to the user's level of required detail and information. The level of detail is dependent on the position's required skills and training. Good SOPs are auditable, easy to follow, and not difficult to find. Auditable characteristics are: observable actions and countable frequencies. Results should be evident to a third party (*compliance to the SOP must be measurable*). Standard operating procedures are an important element in a control plan when a specific process is being prescribed to achieve quality output.

SOP Elements

SOPs are intended to impart high value information in concise and well-documented manner. There are many components to maintain currency, ensure that employees understand why the SOP is necessary and important, and other supporting information to make scope very clear for the SOP.

- SOP Title and Version Number—Provide a title and unique identification number with version information.
- Date—List the original creation date; add all revision dates.
- Purpose—State the reason for the SOP and what it intends to accomplish.
- Scope—Identify all functions, jobs, positions, and/or processes governed or affected by the SOP.

- Responsibilities—Identify job functions and positions (not people) responsible for carrying out activities listed in the SOP.
- Materials—List all material inputs: parts, files, data, information, instruments, etc.
- Process Map—Show high level or level two to three process maps or other graphical representations of operating steps.
- Process Metrics—Declare all process metrics and targets or specifications.
- Procedures—List actual steps required to perform the function.
- References—List any documents that support the SOP.

SOP Template

S	tandard Operating	Procedures	
SOP Name/Title:			
Document Storage Loc	ation/Source:	Document No:	
SOP Originator:	Approving Position:	Effective Date:	
Name:	Name:	Last Edited Date:	
Signature:	Signature:	Other:	

1. Purpose

"Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut

2. Scope

"Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut

3. Responsibilities (RACI)

Responsible	Accountable	Consulted	Informed
John Doe	Jane Doe	Jack Doe	Jill Doe
Pam Doe	Paul Doe	Phil Doe	Peqqy Doe

4. Materials

"Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut

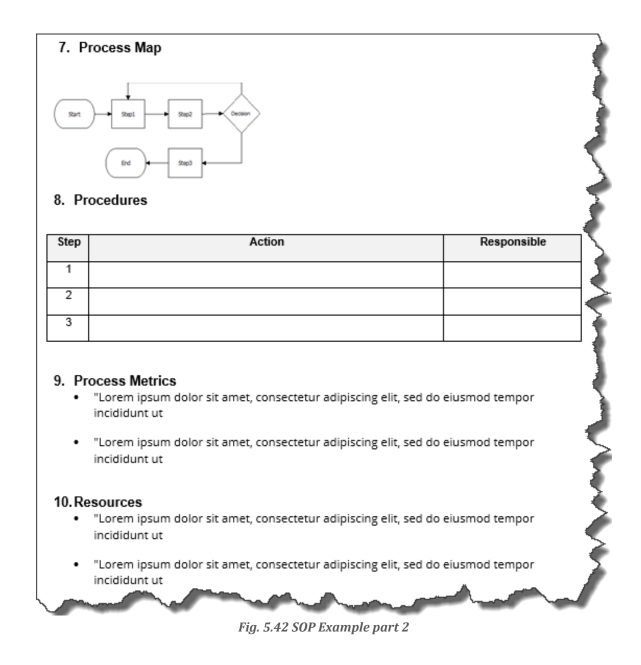
5. Related Documents

"Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut

6. Definitions

"Lorem ipsum dolor sit amet, consectetur adipiscing elit, sed do eiusmod tempor incididunt ut

Fig. 5.41 SOP Example part 1



Communication Plans

Communication plans are documents that focus on planning and preparing for the dissemination of information. They organize messages and ensure that the proper audiences receive the correct message at the right time. A good communication plan identifies:

- Audience
- Key points/message
- Medium (how the message is to be delivered)
- Delivery schedule
- Messenger
- Dependencies and escalation points

• Follow-up messages and delivery mediums

Communication plans help develop and execute strategies for delivering changes to an organization.

Communication Plan Template

					C	commur	nication	Plan T	emplate
Process/Funct	ion Name	Project/Prog	ram <u>Name</u>	Projec	t Lead		Project Spons	or/Champio	<u>n</u>
Communication Pu	rpose:								,
Target Audience	Key Message	Message Dependencies	Delivery Date	Location	Medium	Follow up Medium	Messenger	Escalation Path	Contact Information

Fig. 5.43 Communication Plan Template

Training Plans

Training plans are used to manage the delivery of training for new processes and procedures. Most GB or BB projects will require changes to processes and/or procedures that must be executed or followed by various employees. Training plans should:

- Incorporate all SOPs related to performing new or modified tasks
- Use and support existing SOPs and do not supersede them
- Include logistics
 - \circ One-on-one or classroom
 - Instruction time
 - Location of training materials
 - Master training reference materials
 - $\circ \quad \text{Instructors and intended audience}$
 - Trainee names

Different components of training plan include: training delivery method, training duration, location of materials and references, instructor names, and intended audience.

Training Plan Template

Spons	on	Business Divisio		ad	Project Le	ocess	Pr	ject	Pro
Trainer	Other Requirements	Technology Requirements	Supporting Docs	Training Medium	Key Change/Process	How Many	When	Where	Who
	Trainer		57						

Fig. 5.44 Training Plan Template

Audits

What is an *audit*?

ISO 9000 defines an audit as "a systematic and independent examination to determine whether quality activities and related results comply with planned arrangements and whether these arrangements are implemented effectively and are suitable to achieve objectives."

Audits are used to ensure actions, processes, procedures, and other tasks are performed as expected. "Say what you do and do what you say."—this is fundamentally what audits are checking for.

Audit Guidelines

Audits must be done carefully to achieve the desired result. Audits should be directed by managers, supervisors, and other accountable positions. An audit's purpose must be well-defined and executed by independent unbiased personnel.

Auditors must:

- Be qualified to perform their tasks
- Attend and successfully complete an internal auditing training session
- Be able to identify whether or not activities are being followed according to the defined SOP
- Base conclusions on facts and objective evidence
- Use a well-documented audit checklist
- Audits should confirm compliance or declare non-compliance

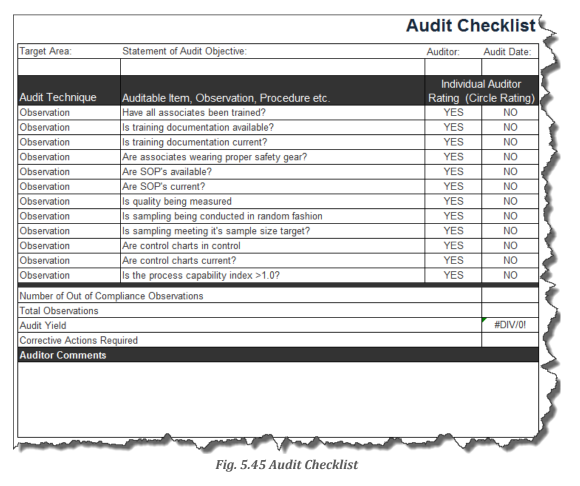
Audit Checklists

Auditors should review the SOPs before preparing checklists or ensure that existing checklists properly reference SOPs.

Audit checklists:

- Serve as guides for identifying items to be examined
- Are used in conjunction with understanding of the procedure
- Ensure a well-defined audit scope
- Identify needed facts during audits
- Provide places to record gathered facts.
- Checklists should include:
- A review of training records
- A review of maintenance records
- Questions or observations that focus on expected behaviors
- Questions should be open-ended where possible
- Definitive observations yes/no, true/false, present/absent, etc.

Audit Checklist Template



5.3.3 Response Plan Elements

What is a Response Plan?

A *response plan* should be a component of as many control plan elements as possible. Response plans are a management planning tool to describe corrective actions necessary in the event of

out-of-control situations. There is never any guarantee that processes will always perform as designed. Therefore, it is wise to prepare for occasions when special causes are present. Response plans help us mitigate risks and, as already mentioned, should be part of several control plan elements.

Response Plan Elements

- Action triggers
 - When do we need to take actions to correct a problem or issue?
- Action recommendation
 - What activities are required in order to solve the problem in the process? The action recommended can be short-term (quick fix) or long-term (true process improvement).
- Action respondent
 - Who is responsible for taking actions?
- Action date
 - When did the actions happen?
- Action results
 - What actions have been taken?
 - When were actions taken?
 - What are the outcomes of the actions taken?

A good response plan clearly describes when the response plan is acted upon, what the action is, who is responsible for taking action and when, and what the results are after the action is taken.



Fig. 5.46 Corrective Action and Responsible Party portion of Control Plan

A response plan denotes what the action is to be taken and who is responsible. Note the response plan element in this control plan template.

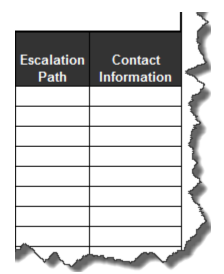


Fig. 5.47 Escalation contact information for communication plan

The communication plan has a section denoting how information is disseminated. Note the response element by the identification of escalations and contact information.

Target Area:	Statement of Audit Objective:	Auditor:	Audit Date:
Audit Technique	Auditable Item, Observation, Procedure etc.		al Auditor
Observation	Have all associates been trained?	YES	NO
Observation	Is training documentation available?	YES	NO
Observation	Is training documentation current?	YES	NO
Observation	Are associates wearing proper safety gear?	YES	NO
Observation	Are SOP's available?	YES	NO
Observation	Are SOP's current?	YES	NO
Observation	Is quality being measured	YES	NO
Observation	Is sampling being conducted in random fashion	YES	NO
Observation	Is sampling meeting it's sample size target?	YES	NO
Observation	Are control charts in control	YES	NO
Observation	Are control charts current?	YES	NO
Observation	Is the process capability index >1.0?	YES	NO
	Compliance Observations		
Total Observations Audit Yield			#DIV/0!
Aught rield	Required		#DIV/0!

Fig. 5.48 Audit Checklist

An audit checklist denotes corrective actions necessary as a result of the audit. The is a form of response plan.

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