Implementing ISO 9000 Quality Management System

Implementation of ISO 9000 affects the entire organization right from the start. If pursued with total dedication, it results in 'cultural transition' to an atmosphere of continuous improvement.

The process of implementing ISO 9000 depends on:

- The sophistication of your existing quality program,
- The size of your organization, and
- The complexity of your process.

The 14 essential steps, briefly described below, are to be followed through in order to implement ISO 9000 quality management system successfully.

Step 1: Top management commitment  
Step 2: Establish implementation team  
Step 3. Start ISO 9000 awareness programs  
Step 4: Provide Training  
Step 5. Conduct initial status survey  
Step 6: Create a documented implementation plan  
Step 7. Develop quality management system documentation  
Step 8: Document control  
Step 9. Implementation  
Step 10. Internal quality audit  
Step 11. Management review  
Step 12. Pre-assessment audit  
Step 13. Certification and registration  
Step 14: Continual Improvement

Step 1: Top Management Commitment

The top management (managing director or chief executive) should demonstrate a commitment and a determination to implement an ISO 9000 quality management system in the organization. Without top management commitment, no quality initiative can succeed. Top management must be convinced that registration and certification will enable the organization to demonstrate to its customers a visible commitment to quality. It should realize that a quality management system would improve overall business efficiency by elimination of wasteful duplication in management system.

The top management should provide evidence of its commitment to the development and implementation of the quality management system and continually improve its effectiveness by:

- Communicating to the organization the importance of meeting customer as well as statutory and regulatory requirements,
- Defining the organization's quality policy and make this known to every employee,
- Ensuring that quality objectives are established at all levels and functions,
Ensuring the availability of resources required for the development and implementation of the quality management system,
Appointing a management representative to coordinate quality management system activities, and
Conducting management review.

The top management should also consider actions such as:

- Leading the organization by example,
- Participating in improvement projects,
- Creating an environment that encourages the involvement of people.

This type of top management commitment may be driven by:

- **Direct marketplace pressure**: requirements of crucial customers or parent conglomerates.
- **Indirect marketplace pressure**: increased quality levels and visibility among competitors.
- **Growth ambitions**: desire to exploit market opportunities.
- **Personal belief in the value of quality** as a goal and quality management systems as a means of reaching that goal.

The top management should identify the goals to be achieved through the quality management system. Typical goals may be:

- Be more efficient and profitable
- Produce products and services that consistently meet customers’ needs and expectations
- Achieve customers satisfaction
- Increase market share
- Improve communications and morale in the organization
- Reduce costs and liabilities
- Increase confidence in the production system

**Step 2. Establish Implementation Team**

ISO 9000 is implemented by people. The first phase of implementation calls for the commitment of top management - the CEO and perhaps a handful of other key people. The next step is to establish implementation team and appoint a Management Representative (MR) as its coordinator to plan and oversee implementation. Its members should include representatives of all functions of the organization - Marketing, Design and development, Planning, Production, Quality control, etc.

In the context of the standard, the MR is the person within the Organization who acts as interface between organization management and the ISO 9000 registrar. His role is, in fact, much broader than that. The MR should also act as the organization’s "quality management system champion," and must be a person with:
- Total backing from the CEO,
- Genuine and passionate commitment to quality in general and the ISO 9000 quality management system in particular,
- The dignity - resulting from rank, seniority, or both - to influence managers and others of all levels and functions,
- Detailed knowledge of quality methods in general and ISO 9000 in particular.

The members of the implementation team should also be trained on ISO 9000 quality management systems by a professional training organization.

**Step 3. Start ISO 9000 Awareness Programs**

ISO 9000 awareness programs should be conducted to communicate to the employees the aim of the ISO 9000 quality management system; the advantage it offers to employees, customers and the organization; how it will work; and their roles and responsibilities within the system. Suppliers of materials and components should also participate in these programs.

The awareness program should emphasize the benefits that the organization expects to realize through its ISO 9000 quality management system. The program should also stress the higher levels of participation and self-direction that the quality management system renders to employees. Such a focus will go far to enlist employee support and commitment.

The programs could be run either by the implementation team or by experts hired to talk to different levels of employees.

**Step 4. Provide Training**

Since the ISO 9000 quality management system affects all the areas and all personnel in the organization, training programs should be structured for different categories of employees - senior managers, middle-level managers, supervisors and workers. The ISO 9000 implementation plan should make provision for this training. The training should cover the basic concepts of quality management systems and the standard and their overall impact on the strategic goals of the organization, the changed processes, and the likely work culture implications of the system. In addition, initial training may also be necessary on writing quality manuals, procedures and work instruction; auditing principles; techniques of laboratory management; calibration; testing procedures, etc.

When in-house capacity to carry out such training is not available, it may be necessary to participate in external training courses run by professional training organizations. Alternatively, an external training institution could be invited to conduct in-house training courses.
Step 5. Conduct Initial Status Survey

ISO 9000 does not require duplication of effort or redundant system. The goal of ISO 9000 is to create a quality management system that conforms to the standard. This does not preclude incorporating, adapting, and adding onto quality programs already in place. So the next step in the implementation process is to compare the organization's existing quality management system, if there is one – with the requirements of the standard (ISO 9001:2000).

For this purpose, an organization flow chart showing how information actually flows [not what should be done] from order placement by the customer to delivery to this customer should be drawn up. From this overall flow chart, a flow chart of activities in each department should be prepared.

With the aid of the flow charts, a record of existing quality management system should be established. A significant number of written procedures may already be in place. Unless they are very much out of date, these documents should not be discarded. Rather, they should be incorporated into the new quality management system. Documents requiring modification or elaboration should be identified and listed. This exercise is sometimes referred to as "gap analysis". During these review processes, wide consultation with executives and representatives of various unions and associations within the organization is required to enlist their active cooperation.

In the review process, documents should be collected, studied and registered for further use, possibly after they have been revised. Before developing new quality management system documentation, you need to consider with which quality requirements or department you should start. The best is to select an area where processes are fairly well organized, running effectively and functioning satisfactorily.

The basic approach is to determine and record how a process is currently carried out. We can do this by identifying the people involved and obtaining information from them during individual interviews. Unfortunately, it often happens that different people will give different, contradicting versions of a process. Each one may refer to oral instructions that are not accurate or clear. This is why the facts are often not described correctly the first time around, and have to be revised several times.

Once it has been agreed how to describe the current process, this process has to be adapted, supplemented and implemented according to the requirements of the quality standard (ISO 9001:2000). This requires organizational arrangements, the drawing up of additional documents and possible removal of existing documentation [e.g. procedures, inspection/test plans, inspection/test instructions] and records [e.g. inspection/test reports, inspection/test certificates].

In introducing a quality management system, the emphasis is on the improvement of the existing processes or the re-organization of processes.

In general, the steps to follow are the following:

- Ascertain and establish the following:
  What is the present operation/process? What already exists?
Analyze the relevant sections of the quality standard - ISO 9001:2000:
What is actually required?
If necessary, supplement and change operational arrangements in accordance with the standard, develop documents and records, and describe operations/processes:
What is the desired operation/process?

Figure 1: Steps in introducing a quality management system

The above gap analysis can be done internally, if the knowledge level is there. Or a formal pre-assessment can be obtained from any one of a large number of ISO 9000 consulting, implementing, and registration firms.

**Step 6. Create a Documented Implementation Plan**

Once the organization has obtained a clear picture of how its quality management system compares with the ISO 9001:2000 standard, all non-conformances must be addressed with a documented implementation plan. Usually, the plan calls for identifying and describing processes to make the organization’s quality management system fully in compliance with the standard.

The implementation plan should be thorough and specific, detailing:

- Quality documentation to be developed
- Objective of the system
- Pertinent ISO 9001:2000 section
- Person or team responsible
- Approval required
- Training required
- Resources required
- Estimated completion date

These elements should be organized into a detailed chart, to be reviewed and approved. The plan should define the responsibilities of different departments and personnel and set target dates for the completion of activities. Once approved, the Management Representative should control, review and update the plan as the implementation process proceeds.

Typical implementation action plan is shown in Figure 2. Use ISO 10005:1995 for guidance in quality planning.
Figure 2. Typical action plan

- Month 1: Appoint MR + establish implementation team
- Month 2: ISO 9000 awareness campaign
- Month 3: Initial status survey + planning
- Month 4: Develop Quality manual-Level A
- Month 5: Write Level B documents
- Month 6: Write Level C documents
- Month 7: Monitor implementation process
- Month 8: First internal audit
- Month 9: Clear nonconformities
- Month 10: Pre-registration audit
- Month 11: Quality training
- Month 12: Compliance audit
- Month 13: Compliance discrepancies
- Registration
Step 7. Develop Quality Management System Documentation

Documentation is the most common area of non-conformance among organizations wishing to implement ISO 9000 quality management systems. As one company pointed out: "When we started our implementation, we found that documentation was inadequate. Even absent, in some areas. Take calibration. Obviously it's necessary, and obviously we do it, but it wasn't being documented. Another area was inspection and testing. We inspect and test practically every item that leaves here, but our documentation was inadequate".

Documentation of the quality management system should include:

- Documented statements of a quality policy and quality objectives,
- A quality manual,
- Documented procedures and records required by the standard ISO 9001:2000, and
- Documents needed by the organization to ensure the effective planning, operation and control of its processes.

Quality documentation is generally prepared in the three levels indicated in the box that follows. Use ISO 10013:1995 for guidance in quality documentation.

<table>
<thead>
<tr>
<th>Level A: Quality manual</th>
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<tr>
<td>States the scope of the quality management system, including exclusions and details of their justification; and describes the processes of the quality management system and their interaction. Generally gives an organization profile; presents the organizational relationships and responsibilities of persons whose work affects quality and outlines the main procedures. It may also describe organization's quality policy and quality objectives.</td>
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<th>Level B: Quality management system procedures</th>
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<td>Describes the activities of individual departments, how quality is controlled in each department and the checks that are carried out.</td>
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<tr>
<th>Level C: Quality documents (forms, reports, work instructions, etc.)</th>
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<tr>
<td>Work instructions describe in detail how specific tasks are performed; include drawing standards, methods of tests, customer's specifications, etc.</td>
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<td>Presents forms to be used for recording observations, etc.</td>
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In small companies, the above levels of documentation could be presented in one manual; otherwise, separate manuals should be prepared.
A list of the documents to be prepared should be drawn up and the responsibility for writing the documents should be assigned to the persons concerned in various functional departments. They should be advised to prepare the drafts within a specific time frame.

**Step 8: Document Control**

Once the necessary quality management system documentation has been generated, a documented system must be created to control it. Control is simply a means of managing the creation, approval, distribution, revision, storage, and disposal of the various types of documentation. Document control systems should be as simple and as easy to operate as possible - sufficient to meet ISO 9001:2000 requirements and that is all.

Document control should include:

- Approval for adequacy by authorized person(s) before issue,
- Review, updating and re-approval of documents by authorized person(s),
- Identification of changes and of the revision status of documents,
- Availability of relevant versions of documents at points of use,
- Identification and control of documents of external origin,
- Assurance of legibility and identifiability of documents, and
- Prevention of unintended use of obsolete documents.

The principle of ISO 9000 document control is that employees should have access to the documentation and records needed to fulfil their responsibilities.

**Step 9. Implementation**

It is good practice to implement the quality management system being documented as the documentation is developed, although this may be more effective in larger firms. In smaller companies, the quality management system is often implemented all at once throughout the organization. Where phased implementation takes place, the effectiveness of the system in selected areas can be evaluated.

It would be a good idea initially to evaluate areas where the chances of a positive evaluation are high, to maintain the confidence of both management and staff in the merits of implementing the quality management system.

The implementation progress should be monitored to ensure that the quality management system is effective and conforms to the standard. These activities include internal quality audit, formal corrective action and management review.

**Step 10. Internal Quality Audit**

As the system is being installed, its effectiveness should be checked by regular internal quality audits. Internal quality audits are conducted to verify that the installed quality management system:
Conform to the planned arrangements, to the requirements of the standard [ISO 9001:2000] and to the quality management system requirements established by your organization, and

Is effectively implemented and maintained.

Even after the system stabilizes and starts functioning, internal audits should be planned and performed as part of an ongoing strategy.

A few staff members should be trained to carry out internal auditing. Use ISO 19011 for guidance in auditing, auditor qualification and programmes.

**Step 11. Management Review**

When the installed quality management system has been operating for three to six months, an internal audit and management review should be conducted and corrective actions implemented. The management reviews are conducted to ensure the continuing suitability, adequacy and effectiveness of the quality management system. The review should include assessing opportunities for improvement and the need for changes to the quality management system, including the quality policy and quality objectives.

The input to management review should include information on:

- Results of audits,
- Customer feedback,
- Process performance and product conformity,
- Status of preventive and corrective actions,
- Follow-up actions from previous management reviews,
- Changes that could affect the quality management system, and
- Recommendations for improvements.

Management reviews should also address the *pitfalls* to effective implementation, including *lack of CEO commitment, failure to involve everyone in the process, and failure to monitor progress and enforce deadlines.*

**Step 12. Pre-assessment Audit**

When system deficiencies are no longer visible, it is normally time to apply for certification. However, before doing so, a pre-assessment audit should be arranged with an independent and qualified auditor. Sometimes certification bodies provide this service for a nominal charge. The pre-assessment audit would provide a degree of confidence for formally going ahead with an application for certification.

**Step 13. Certification and Registration**

Once the quality management system has been in operation for a few months and has stabilized, a formal application for certification could be made to a selected certification agency. The certification agency first carries out an audit of the documents (referred to as an "adequacy audit"). If the documents conform to the requirements of the
quality standard, then on-site audit is carried out. If the certification body finds the
system to be working satisfactorily, it awards the organization a certificate, generally
for a period of three years. During this three-year period, it will carry out periodic
surveillance audits to ensure that the system is continuing to operate satisfactorily.

**Step 14: Continual Improvement**

Certification to ISO 9000 should not be an end. You should continually seek to improve
the effectiveness and suitability of the quality management system through the use of:

- Quality policy
- Quality objectives
- Audit results
- Analysis of data
- Corrective and preventive actions
- Management review

**ISO 9004:2000** provides a methodology for continual improvement.