Sensing and Control

Quality Manual

ISO/TS 16949:2009
Quality Management System

CORPORATE HEADQUARTERS:
OPTEK Technology
1645 Wallace Drive
Carrollton, TX  75006

MANUFACTURING:
Optron de Mexico
Avenue Rio Bravo #1551
Parque Industrial Rio Bravo
Cd. Juarez, Chihuahua 32700
# Table of Contents

ABOUT THIS MANUAL ............................................................................................................. 6

APPROVALS: ............................................................................................................................ 6

OVERVIEW OF TT ELECTRONICS / OPTEK .......................................................................... 6

SCOPE ...................................................................................................................................... 7

APPLICABLE STANDARDS ...................................................................................................... 7

MANAGEMENT STRUCTURE .................................................................................................. 8

1 MANAGEMENT PROCESSES .......................................................................................... 8

1.1 Quality Management System Implementation & Maintenance ....................................... 8
   1.1.1 QMS General Requirements .................................................................................... 8
   1.1.2 Management Commitment .................................................................................... 10
   1.1.3 Customer Focus .................................................................................................... 10
   1.1.4 Quality Policy ......................................................................................................... 10
   1.1.5 Quality Management System Planning .................................................................. 11
   1.1.6 Responsibility and Authority ................................................................................... 12
       1.1.6.1 Responsibility for quality ................................................................................. 12
       1.1.6.2 Departmental responsibility for quality ............................................................ 12
       1.1.6.3 Individual responsibility ................................................................................... 12
   1.1.7 Management Representative ................................................................................. 13
       1.1.7.1 Customer representative ................................................................................ 13
   1.1.8 Provision of Resources .......................................................................................... 14
   1.1.9 Human resources - General ................................................................................... 14
       1.1.9.1 Competence, awareness and training ............................................................ 14
       1.1.9.2 Product design skills ....................................................................................... 15
       1.1.9.3 Training ........................................................................................................... 15
       1.1.9.4 Training on the job .......................................................................................... 15
       1.1.9.5 Employee motivation and empowerment ........................................................ 15
   1.1.10 Planning of Product Realization ............................................................................ 15
       1.1.10.1 Planning of product realization - Supplemental .............................................. 16
       1.1.10.2 Acceptance criteria ......................................................................................... 16
       1.1.10.3 Confidentiality ............................................................................................... 16
       1.1.10.4 Change control ............................................................................................... 16

1.2 Management Review ................................................................................................... 16
   1.2.1 General - Quality Management System Performance (Mgmt. Review) ............ 16
1.2.1.1 Review Input ................................................................. 17
1.2.1.2 Review Output .............................................................. 18
1.2.2 Quality Objectives .......................................................... 18
1.2.3 Measurement Analysis & Improvement - General .................. 21
  1.2.3.1 Identification of Statistical Tools .................................. 21
  1.2.3.2 Knowledge of Basic Statistical Concepts ..................... 21
1.2.4 Customer Satisfaction ..................................................... 21
1.2.5 Analysis and Use of Data .................................................. 22
1.2.6 Continual Improvement .................................................... 23

1.3 Communication ........................................................................ 23
  1.3.1 Internal Communication ................................................ 23
  1.3.2 Customer Communication .............................................. 23

1.4 Internal Auditing ...................................................................... 24
  1.4.1 Internal Audit of the Quality Management System ............ 24
  1.4.2 Internal Audit Plans ....................................................... 24
  1.4.3 Internal Auditor Qualification ......................................... 25
  1.4.4 Manufacturing Process Audit ......................................... 25
  1.4.5 Product Audit ............................................................... 25

1.5 Corrective & Preventive Action ............................................... 25
  1.5.1 Corrective Action ........................................................... 25
    1.5.1.1 Problem Solving ..................................................... 26
    1.5.1.2 Error-Proofing ....................................................... 26
    1.5.1.3 Corrective Action Impact ....................................... 26
    1.5.1.4 Rejected Product Test / Analysis ............................... 26
  1.5.2 Preventive Action .......................................................... 26

2 REALIZATION PROCESSES: .................................................... 27

2.1 Requirements Determination & Review .................................... 27
  2.1.1 Determination of requirements related to product ................ 27
  2.1.2 Review of Requirements Related to Product, Manufacturing Feasibility .......... 28

2.2 Design & Development ......................................................... 28
  2.2.1 Design and Development ............................................... 28
  2.2.2 Design and Development Planning ................................... 28
  2.2.3 Multidisciplinary Approach ............................................ 29
  2.2.4 Design and Development Input ....................................... 29
  2.2.5 Special Characteristics .................................................. 29
  2.2.6 Design and Development Output ...................................... 29
  2.2.7 Design and Development Review, Monitoring ................. 30
  2.2.8 Design and Development Verification ............................. 30
  2.2.9 Design and Development Validation ............................... 30
  2.2.10 Prototype Program ..................................................... 30
2.2.11 Product Approval Process ................................................................. 30
2.2.12 Control of Design and Development Changes ............................... 31

2.3 Purchasing .......................................................................................... 31
2.3.1 Purchasing Process ........................................................................... 31
2.3.2 Regulatory Conformity ..................................................................... 31
2.3.3 Supplier Quality Management System Development .................... 31
2.3.4 Customer-Approved Sources .......................................................... 32
2.3.5 Purchasing Information .................................................................... 32
2.3.6 Verification of Purchased Product and Incoming Product Quality .... 32
2.3.7 Supplier Monitoring ........................................................................... 33

2.4 Facilities & Equipment Management ................................................. 33
2.4.1 Infrastructure ................................................................................... 33
2.4.1.1 Plant, facility and equipment planning ......................................... 33
2.4.1.2 Contingency plans ....................................................................... 33
2.4.2 Work Environment ........................................................................... 34
2.4.2.1 Work Environment ...................................................................... 34
2.4.2.2 Personnel Safety to Achieve Product Quality ............................... 34
2.4.2.3 Cleanliness of Premises .............................................................. 34
2.4.3 Customer Property .......................................................................... 34
2.4.3.1 Customer Property ...................................................................... 34
2.4.3.2 Customer Owned Production Tooling ......................................... 35
2.4.4 Control of Monitoring & Measurement Devices .............................. 35
2.4.4.1 Control of Monitoring and Measuring Devices ............................. 35
2.4.4.2 Measurement System Analysis .................................................... 36
2.4.4.3 Calibration / Verification Records ............................................... 36
2.4.4.4 Laboratory Requirements ............................................................ 36
2.4.4.4.1 Internal Laboratory ................................................................. 36
2.4.4.4.2 External Laboratory ............................................................... 36

2.5 Company Control Specifics ................................................................. 37
2.5.1 Document Control .......................................................................... 37
2.5.1.1 Control of Documents and Engineering Specifications .............. 37
2.5.2 Records Control .............................................................................. 37
2.5.2.1 Control and Retention of Records ................................................ 37
2.5.3 Control of Production Provision ...................................................... 38
2.5.3.1 Control of Production ................................................................. 38
2.5.3.2 Control Plan ................................................................................. 38
2.5.3.3 Work Instructions ...................................................................... 39
2.5.3.4 Verification of job set-ups ......................................................... 39
2.5.3.5 Preventive and Predictive Maintenance ...................................... 39
2.5.3.6 Management of Production Tooling ............................................ 39
2.5.3.7 Production Scheduling ............................................................... 40
2.5.3.8 Feedback of Information ............................................................. 40
2.5.4 Validation of Processes ................................................................... 40
About this Manual

This manual was developed for a quality management system aimed primarily at achieving customer satisfaction by meeting customer requirements through the application of the system, the continual improvement of the system and the prevention of nonconformity. The quality management system developed by this manual was based on & demonstrates throughout its wording good quality principles such as leadership, customer focus, continual improvement, involvement of the employees, mutually beneficial supplier relationships, process focus, management by a system of processes and ultimately decisions that are based on good sound data.

The Management Representative maintains this manual for the Quality Management System and for Optek employees’ use. The online version incorporates links (in blue) to other pertinent procedures, process maps, and documents making it use easier than a printed version.

Requests for changes should be submitted to the Management Representative. Updates of the manual are issued as required. The online version of this quality manual is available to all employees via computers. The online version is to be considered the most current and takes precedence over any printed copy.

It is the responsibility of the Department Heads to ensure that employees are familiar with the manual's content related to their work and responsibilities, and that they are kept informed of any changes and updates.

Effective date of this quality manual: May 3, 2004

Approvals:

<table>
<thead>
<tr>
<th></th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jeff Beatty</td>
<td>3-31-14</td>
</tr>
<tr>
<td>Vice President / General Manager</td>
<td>3-31-14</td>
</tr>
</tbody>
</table>

Overview of TT electronics / OPTEK

**TT electronics / OPTEK Technology**, hereafter called **OPTEK**, is headquartered at 1645 Wallace Drive, Carrollton, Texas 75006 with manufacturing facilities at Optron de Mexico, Avenue Rio Bravo #1551 Park Industrial Rio Bravo, Cd. Juarez, Chihuahua Mexico 32700. OPTEK’s business activities are the design, manufacturing, and sale of electronic sensors. OPTEK provides sensor technologies such as optical (IR, visible, UV & VCSEL), magnetic (Hall-effect) and fiber optic (LED & VCSEL to 2.5G). OPTEK sensors are found in office
equipment, industrial applications, encoders, military & Hi-Rel applications, medical diagnostic equipment, automotive engine and door controls and ignition security and fiber optic data communication applications. Sensors can be standard product or custom made to customer’s specification.

This quality manual describes the Quality Management System of OPTEK. Its purpose is:

- **for internal use**, to communicate to employees the company’s Quality Policy and quality objectives, to make the employees familiar with the method of compliance with ISO/TS 16949 requirements, to facilitate the implementation and maintenance of the Quality Management System and to ensure its continuity and required updates during changing circumstances, to provide effective communication and control of quality related activities and a documented base for quality system audits.

- **for external use**, to inform OPTEK’s customers and other interested external partners about OPTEK’s Quality Policy, OPTEK’s implemented Quality Management System, and measures of compliance with the requirements of ISO/TS 16949.

Scope

This Quality Management System described hereafter complies with all the requirements of ISO/TS 16949 Revision 2009, is focused on the enhancement of customer satisfaction through continual improvement of processes and products, and demonstrates compliance with customer and regulatory requirements.

The scope of the Quality Management System covers the design and manufacturing of sensor systems, the manufacture of custom made piece parts and final products as well as the construction of dies and fixtures at either OPTEK’s location at 1645 Wallace Drive, Carrollton, Texas 75006 or Optron de Mexico, Juarez manufacturing site.

OPTEK’s Quality Management System meets all requirements of ISO/TS 16949:2009 and no exclusions are applied.

Applicable Standards

The content and application of this quality manual makes reference to the following publications and documents:

- **ISO 9001: 2008**, Quality Management Systems - requirements
- **ISO/IEC 80079-34**
For this quality manual, the terms and definitions for the automotive industry specified in clause 3.1 of ISO/TS 16949 are applied.

The quality manual includes the scope of the Quality Management System with no exclusions from the requirements of ISO/TS 16949 and their justification, makes reference to applicable operating procedures and other pertinent documents. Process Map #001 is a description of the Quality Management System interaction of processes.

Management Structure
The management of OPTEK consists of the President/CEO. Reporting to the President/CEO and his Executive Staff are the department managers of Quality Assurance, Business Unit Managers, Manufacturing, Engineering and Administration.

1 Management Processes

1.1 Quality Management System Implementation & Maintenance
It is the responsibility of the Management Representative

- to ensure that the Quality Management System of OPTEK is established, documented as required, implemented, managed and maintained according to the requirements of ISO/TS 16949
- to continually improve the effectiveness of the Quality Management System

1.1.1 QMS General Requirements

a) Operational and administrative activities affecting quality of the departments Engineering, Manufacturing, Quality Assurance, Purchasing, Warehouse, Toolroom, Sales and Customer Service and Quality Management System Administration are in compliance with ISO/TS 16949. It is the responsibility of the Management Representative and Department Heads to ensure that the activities/processes included in the scope of this Quality Management System are identified and are performed in compliance with ISO/TS 16949.

b) It is the responsibility of the Management Representative and Department Heads to ensure that the sequence and interaction of processes or activities of this Quality Management System are determined in a suitable manner, utilizing tools such as process maps, quality plans, flow charts, operating procedures, etc.

c) It is the responsibility of the Management Representative and Department Heads to apply the necessary techniques and criteria in order to verify that established processes/activities and their implemented controls are effective.

d) It is the responsibility of the President/CEO, Management Representative and the Department Heads to ensure that the necessary human and material resources, as well as the necessary information, are available to ensure the effective operation and control of the processes of the Quality Management System.
e) Compliance of product with IECE x Certificate of Conformity # IECE x BAS 11.0123u

f) It is the responsibility of the Management Representative and Department Heads to ensure that the processes / activities which are part of the Quality Management System are monitored, measured and analyzed regarding their achievement of planned results.

g) As required, the Management Representative and Department Heads ensure that action is taken to obtain expected results of processes/activities, as well as the continual improvement of these processes/activities.

In the event that processes, which affect product conformity, are outsourced, Quality Assurance via Supplier Quality establishes and implements the necessary controls for approval processes to ensure conformance to specified requirements. These implemented controls however do not absolve OPTEK from the responsibility of supplying products that meet customer requirements.

h) QMS Documentation requirements
As a minimum, the documentation of OPTEK’s Quality Management System includes
- a Quality Policy and objectives
- this quality manual
- required documented procedures or maps for ISO/TS 16949 clauses: 4.2.3 Control of Documents; 4.2.4 Control of Records; 6.2.2.2 Training, 8.2.2 Internal Audit; 8.3 Control of Nonconforming Product; 8.5.2 Corrective Action and 8.5.3 Preventive Action
- other documents needed by OPTEK for the effective planning, operation and control of processes of the Quality Management System
- records required by ISO/TS 16949 and by OPTEK to ensure appropriate control and evidence of compliance with requirements.

The document structure of OPTEK’s Quality Management System consists of two levels:


2) Operating procedures, work instructions, forms, master lists, operating instructions, quality plans, control plans and other documents needed by OPTEK for the effective and efficient operation of the Quality Management System.

In addition, records are created as required by ISO/TS 16949, as well as records necessary to meet other internal and external requirements.
It is the responsibility of the Management Representative to ensure the availability of corporate documents of the Quality Management System and documents required by ISO/TS 16949.

It is the responsibility of the Department Heads to ensure the development and availability of documented procedures, work instructions, operating instructions and any other documents related to their departments, that are necessary to ensure the effective implementation, control and functioning of the Quality Management System and its processes.

1.1.2 Management Commitment

The management of OPTEK is committed to the development, implementation and optimum functioning of the Quality Management System and the continual improvement of its effectiveness. In order to provide this evidence, the President/CEO ensures that:

- a corporate Quality Policy is established
- quality objectives are established by selected departments based on the company business plan
- the importance of meeting customer requirements and statutory and regulatory requirements is part of the training of each employee (QS02, MF09)
- resources for the implementation and maintenance of the Quality Management System and its processes are provided in a timely manner
- at a minimum, yearly management reviews are conducted to verify the effectiveness, efficiency and proper functioning of the Quality Management System, including product realization processes and support processes.

1.1.3 Customer Focus

The President/CEO ensures that procedures for determining and meeting customer requirements are established and implemented. The effectiveness of these procedures is measured through customer satisfaction surveys, which are part of Management Reviews (see 2.1.1 & 1.2.4).

1.1.4 Quality Policy

The management of OPTEK has developed a Corporate Quality Policy, which meets the needs of OPTEK and its customers.

Corporate Quality Policy

OPTEK is dedicated to delivering Reliable World Class competitive products on time to meet the requirements of our customers. OPTEK is committed to its Quality Management System, Quality Objectives and will continually improve their effectiveness.
It is the responsibility of OPTEK’s management to implement and maintain this Quality Policy. The Quality Policy includes OPTEK’s commitment for continual improvement, for meeting internal requirements and customer requirements, and provides a basis for the establishment and review of quality objectives. The Quality Policy is made known within the organization and understood and adhered to by employees. During management reviews, the Quality Policy is reviewed for its continuing suitability.

1.1.5 Quality Management System Planning

In the first and the third quarter of the year, the Management Representative calls for a meeting of the Department Heads with the purpose to review, coordinate and plan the efficiency and effectiveness of the Quality Management System and the realization of established quality objectives of the departments, as well the coordination of improvement opportunities. The General Requirements of clause 4.1 of ISO/TS 16949 are included in this planning process.

The output of these planning activities includes the identification of required resources. As appropriate, results from audits of the Quality Management System as well as permissible exclusions according to ISO/TS 16949 are considered. Planning activities are documented and are consistent with other requirements of the Quality Management System.

It is the responsibility of the Management Representative to ensure that resulting organizational changes and their consequences are identified and defined, that changes resulting from planning activities are coordinated and implemented in a controlled manner, that changes to the Quality Management System are documented, implemented and approved, and that the Quality Management System is properly maintained during these changes.

Note: the Quality Planning Team under the responsibility of the Manufacturing Department performs Quality planning for manufacturing process activities.

Each Department Head develops and maintains a process map, flow chart, quality plan, operating procedure, etc for his/her department, showing the workflow of the department as well as evidence of compliance with the requirements of the quality system.
1.1.6 Responsibility and Authority

It is the responsibility of the Management Representative to develop and maintain an organization chart of OPTEK. Updated charts are distributed to Department Heads and are available to employees on a need to know basis.

1.1.6.1 Responsibility for quality

It is the responsibility of personnel in Quality Assurance, or personnel in Production and Warehouse, to inform the responsible Department Head of any nonconformity of products or processes. Corrective action is taken as appropriate, including action for the review and improvement of processes. Documents are updated as required.

If necessary, the Department Head can transfer the nonconformity to the Quality Planning Team for review and action.

In the event of nonconformity in production, all personnel are responsible for product quality and have the authority to stop production in order to correct any quality problems. It is ensured that an employee responsible for quality is present at all times during production.

1.1.6.2 Departmental responsibility for quality

The responsibility for quality in each department rests with the Department Head. The Department Head is responsible for the development and approval of the department’s procedures and work instructions. The Department Heads ensure that the department staff understands and follows the applicable policies and guidelines outlined in the Quality Manual, that the department’s personnel adheres to all applicable procedures and work instructions and participates, as appropriate, in the quality improvement process.

In addition, employees are made aware of the importance to meet customer requirements and expectations. It is the responsibility of the Department Heads to ensure that customer requirements and customer expectations, which relate to activities under the department’s responsibility, are identified, defined, documented and met.

Department Heads ensure that the responsibility of employees or functions whose activities affect quality, are defined in procedures and job descriptions. In yearly performance reviews, or when required, these responsibilities for quality, together with other responsibilities of the function, are reviewed and discussed between the Department Head and the employee, and are assessed and re-defined as necessary.

1.1.6.3 Individual responsibility

All employees follow the policies and guidelines outlined in the quality manual and in established procedures. It is the duty of each employee to inform the Department Head or the Management Representative when performed activities do not match
the established procedures, or when established procedures and work instructions are unclear or ambiguous. The Department Head is notified of any identified nonconformity or deficiency where the correction or prevention of such nonconformity or deficiency is out of the employee’s scope of responsibility.

1.1.7 Management Representative

The management of OPTEK appoints a Management Representative who, irrespective of other duties and responsibilities, has the defined authority and responsibility for

- ensuring that a Quality Management System is established, implemented and maintained in accordance with the requirements of the international standard ISO/TS 16949
- reporting on the performance of the Quality Management System to management for review and corrective action, including needs for improvements
- the proper coordination - where required - of quality related issues and activities within the company, with external customers and suppliers and other external partners and authorities
- promoting the awareness of customer requirements and expectations throughout the company

The Management Representative is responsible for the overall coordination, implementation and administration of the Quality Management System.

1.1.7.1 Customer representative

The management of OPTEK has assigned the inter-departmental coordination of customer requirements to the manager of the Sales Department, and the direct responsibility of addressing customer requirements to the Department Heads of Sales, Engineering and Manufacturing.

Responsibilities related to customer requirements include:

**Sales:**
- coordination of issues related to customer requirements
- recommendations for corporate quality objectives, including quality objectives for other departments
- conveyance of temporary deviations to customers
- analysis of feedback from customers regarding nonconformities
- follow-up on corrective actions

**Engineering:**
- recommendations for corporate quality objectives, including quality objectives for other departments
- quality planning activities
- product design and development
- communication with customers on technical issues
• customer prototype support

Manufacturing:
• recommendations for corporate quality objectives, including quality objectives for other departments
• special training requirements for production and warehouse
• production planning
• issues related to customer supplied products (in coordination with Purchasing and/or Supplier Quality Engineering)

• delivery requirements - shipping inspection

The Sales Department is informed of corrective actions taken by the responsible department regarding the compliance with customer requirements.

1.1.8 Provision of Resources
Management ensures that approved material and human resources, which have been identified by the Department Heads during budget planning and quality planning, are available in a timely manner. This refers to resources required for the implementation, maintenance and continual improvement of the processes of the Quality Management System, for meeting customer requirements and achieving customer satisfaction. Also included are resource requirements for new projects and other quality related activities. Related expenses are included in the company’s financial budget.

1.1.9 Human resources - General
It is the responsibility of the Department Heads to identify qualification requirements of functions or personnel assigned to defined activities that affect quality of product. Qualification requirements include education, training, skills and experience as appropriate. In the department’s budget are provisions for the employment and assignment of qualified and trained personnel.

1.1.9.1 Competence, awareness and training
Department Heads ensure that the qualification requirements (such as education, skills, training, experience) for each job are identified, determined and documented in job descriptions. Training is provided to employees (QS02, MF01) or other actions are taken in order to meet defined qualification requirements. The effectiveness of provided training or of related actions is evaluated.

Employees are made aware of the importance and the impact of their work in relation to product quality, to the achievement of quality objectives and customer satisfaction

Records of employees’ education, experience and other qualifications are maintained.
1.1.9.2 Product design skills

It is the responsibility of Engineering to ensure that personnel with responsibility of product design are qualified for the job and have the necessary skills and experience as specified in the applicable job description.

1.1.9.3 Training

The Management Representative establishes and maintains the documented procedure (QS02 Training) for identifying training needs and for providing required training to employees who are performing activities affecting product quality. The procedure includes training for the fulfillment of specific customer requirements. The HR Representative (MF09) provides training for safety and the handling of hazardous materials.

1.1.9.4 Training on the job

Department Heads ensure that personnel, assigned to new or modified responsibilities affecting product quality, are trained on-the-job. This applies only to on-site contracted personnel. Personnel performing activities that can affect quality are informed of potential consequences to the customer in the event that defined requirements will not be met (QS02).

1.1.9.5 Employee motivation and empowerment

To promote innovation, to motivate employees in accomplishing quality objectives and to participate in the continual improvement process, the Management Representative publishes quarterly at least one article in the company’s bulletin regarding the importance, advantages, challenges and past achievements of these activities. Promoting the awareness for quality and technology to all employees is part of this process. An alternative would be to post these articles on the company’s bulletin boards/electronic bulletin boards.

Performance reviews conducted by the Department Heads include and document the degree of employee’s awareness regarding the importance of their work and their contribution in achieving quality objectives.

1.1.10 Planning of Product Realization

The Engineering Department is responsible for the quality planning (EG01) of the production processes of new products and for changes of existing products. Planning activities are consistent with other requirements of the Quality Management System. Prior and during the planning process, quality objectives and quality requirements for product related to the planning project are established by the quality planning team.

As appropriate, the planning process covers provision of resources necessary, manufacturing processes and documents, required verification, validation, monitoring, inspection and test activities, and criteria for product acceptance.
Records for providing evidence that manufacturing processes, and manufactured product meet requirements are defined and specified.

1.1.10.1 Planning of product realization - Supplemental

Customer requirements and references to technical specifications are included in the quality plan.

1.1.10.2 Acceptance criteria

Acceptance criteria are defined in the planning process and, as required, approved by the customer. Acceptance for attribute data sampling is zero defects (EG01).

1.1.10.3 Confidentiality

Confidentiality of information and data about customer-contracted products/projects is ensured (EG01).

1.1.10.4 Change control

Changes to production processes (EG01), including changes to products/materials from suppliers, are assessed, validated and approved by Engineering prior to use and implementation. For proprietary designs, the impact of changes is reviewed with the customer. If requested by the customer, additional verification/identification requirements are met.

1.2 Management Review

1.2.1 General - Quality Management System Performance (Mgmt. Review)

At planned intervals, management and the Management Representative have a formal meeting to discuss and review (Process Method #002) the continuing effectiveness and adequacy of the Quality Management System. The review includes the evaluation of the need for changes to the Quality Management System, the Quality Policy and quality objectives, as well as the assessment of improvement opportunities based on the review and analysis of performance trends, achievement of quality objectives and customer satisfaction, and the cost of poor quality according to clauses 8.4.1 and 8.5.1 of ISO/TS16949. The maximum interval between Management Reviews should be twelve (12) months, not to exceed fourteen (14) months.

This review covers all clauses of the Quality Management System. As required, Department Heads and employees are invited to attend the meeting when issues of his/her area of responsibility are discussed.

The Management Representative, with the input from management, prepares the agenda of upcoming meetings, ensures that the required data and documents are available for management review, writes the minutes of the meeting, informs results to the Department Heads and individuals concerned and follows-up on required actions
resulting from these meetings. Management is kept informed on the status of follow-up activities. Records of management reviews are maintained.

### 1.2.1.1 Review Input

As a minimum, the following inputs are to be considered for at least one meeting during the year:

- performance of production processes, product conformity, including performance trends
- status and effectiveness of corrective and preventive actions
- follow-up actions from previous management reviews
- planned changes that could affect the Quality Management System
- assessment of improvement opportunities and recommendations for improvements
- employees’ knowledge, understanding and adherence to quality policies, directives and procedures, and their involvement in the quality improvement process
- availability and effectiveness of internal and external information within the company, at all levels
- human resources, training and staffing requirements
- suitability of working environment
- availability of material resources
- effectiveness of quality planning (quality system, design/development of products and processes)
- achievement of corporate quality objectives, including those specified in the business plan
- achievement of departmental quality objectives
- effectiveness of continual improvement activities of products, processes, and Quality Management System
- results of internal and external audits of the quality system, incl. audits of manufacturing processes
- statistical results of operational performance, based on the analysis of collected data, which includes:
  - quality system audits conducted by the Registrar
  - internal quality audits
  - customer satisfaction surveys and other customer feedback regarding customer satisfaction
  - customer complaints
  - suppliers performance
  - product and service quality and nonconformities
  - direct and indirect costs and benefits of the quality system (relation cost - benefit)
  - cost of poor quality - scrap, rework, returns, warranty repairs, excessive freight charges, etc.
  - analysis of field-failures (returns, warranty repairs), their impact on quality, safety, environment
the impact or potential impact of changes that could affect the Quality Management System
opportunities for improvement resulting from additions or changes
• ISO/IEC 80079-34 standards for “Equipment for use in explosive atmospheres”
• Review of internal audit, and overall processes & external (authorized product line engineer must attend meeting)

1.2.1.2 Review Output
Results of the assessment and conclusions of management reviews include the following output:
• effectiveness of corrective and/or preventive actions
• the suitability and effectiveness of the Quality Management System
• improvement of the effectiveness of the Quality Management System and its processes
• improvement of product with focus on customer requirements
• availability of human and material resources
• suitability of the corporate Quality Policy
• frequency of Management Reviews
• required corrective and or preventive actions regarding items reviewed

1.2.2 Quality Objectives
Each year, management defines Corporate Quality Initiatives in the Business Plan. Yearly quality objectives and measurements are established by Department Heads for their departments based on these Quality Initiatives. Management approves these departmental quality objectives.

Established quality objectives are consistent with the Quality Policy, include - as appropriate - objectives to meet product requirements (see 1.1.10), and are defined in such a way that their degree of achievement and results can be measured.

At least two quality objectives of the departments Manufacturing and Quality Assurance are related to the performance of product.

The completion and achievement of yearly quality objectives included in the business plan and departmental quality objectives are reviewed during management review regarding their level of achievement.

Corporate quality initiatives
Based on the corporate Quality Policy, the management of OPTEK has established the following corporate quality initiatives based on one premise – that of **Exceeding the Customer’s Expectations**. These quality initiatives will be reviewed each year to ensure alignment with the business plan.
• Provide Exemplary Customer Satisfaction
• Be a leader in Advanced Technology Development
• Eliminate waste wherever it exists
• Be the industry benchmark for manufacturing capability
• Provide a work environment that promotes competency, teamwork, empowerment and accountability
• Deliver exceptional financial performance over time
The translation from a Corporate Quality Initiative to a measurable index for demonstrating results and continual improvement is demonstrated in the following example:

The end result of this process is to clearly tie the business plan into operational terms that have meaning to front line personnel at all levels within the organization. Our business objectives are our Quality Objectives.
Based on internal and external audit results and statistics issued by the Management Representative (QS04), these quality initiatives are reviewed during management reviews regarding their continuing suitability.

**Departmental quality objectives**
Each year, the Department Heads of Administration, Sales, Engineering, Manufacturing, Warehouse, Toolroom and Quality Assurance establish quality objectives for his/her department. These quality objectives are in accordance with the corporate Quality Policy and are focused on the improvement of departmental processes/activities (see process maps). The departmental quality objectives for the coming year are submitted to the President/CEO for review and approval. A summary of the achievement of the department’s quality objectives of the past year is documented by the applicable Department Head and submitted to the President/CEO.

### 1.2.3 Measurement Analysis & Improvement - General

In order to demonstrate the conformity of manufactured product, the conformity of the Quality Management System and its continual improvement, the Management Representative develops and distributes quarterly statistics (QS04). These statistics are analyzed by Department Heads and corrective and preventive action for the continual improvement of the Quality Management System is taken as appropriate.

#### 1.2.3.1 Identification of Statistical Tools

During product quality planning (EG01), appropriate statistical tools are determined for each process and are included in the control plan. This includes statistical methods for product development (variation analysis, dependability analysis, etc.), for product verification (process capability, variation analysis, control charts, etc.), and other processes.

#### 1.2.3.2 Knowledge of Basic Statistical Concepts

Department Heads, with the assistance of the Management Representative ensure that personnel are trained in the use and application of basic statistical concepts defined by quality planning and used in their respective departments. Training records are maintained (QS02).

### 1.2.4 Customer Satisfaction

Periodic customer satisfaction surveys (QS07) are conducted by management to verify if customer satisfaction has been achieved. Survey results, which include customer complaints and feedback, customers' business disruptions (QS06), customer returns of nonconforming product (217-0027-001) and delivery performance (QS04) are analyzed and evaluated (QS05). As required, management takes corrective or preventive action. The Management Representative monitors the effectiveness of these corrective or preventive actions.
1.2.5 Analysis and Use of Data

The Management Representative issues statistics regarding the performance of the Quality Management System (QS04) thru the analysis and use of data. Ratings on supplier performance are issued by Purchasing and/or Quality Assurance (PU04). The statistics are analyzed by the Management Representative regarding the effectiveness, suitability and opportunities for improvement of the processes of the Quality Management System, and by Department Heads regarding the performance and suitability of activities and processes under their responsibility. This includes the analysis of customer complaints (QS06) and customer returns (QA08).

A summary report is issued by the Management Representative, providing information on: customer satisfaction or dissatisfaction, product quality, characteristics and trends of processes and products including opportunities for preventive action, and supplier performance.

The Management Representative controls and coordinates the implementation of required corrective or preventive actions. Analysis results of statistics and actions are reported by the Department Heads to the Management Representative who monitors the progress and results of these actions.

In addition, trends in quality and operational performance are compared with progress toward objectives and lead to action to support: the development of priorities to resolve customer-related problems, to determine customer related trends and correlation for status review, decision making and longer term planning, and an information system for reporting of product information related to usage.

Improvement

It is the responsibility of the Management Representative to form and implement a Quality Planning Team for the handling of assigned activities related to the Quality Management System. Members should be from Quality Assurance, Manufacturing, Engineering, Sales, Purchasing and any other additional members deemed necessary.

The purpose of the Quality Planning Team is to review, analyze and make final decisions on Corrective Action Requests (QSW01) and Quality Improvement Proposals (QSW03), to make recommendations for preventive actions and quality improvements, to coordinate and implement preventive actions and quality improvement projects, monitor results, and to provide a forum for any quality issue which requires a cross-functional approach. Nonconformities and deficiencies are analyzed, root causes are determined and required action is taken or recommended as appropriate.

As required and/or decided by management, selected Quality Improvement Proposals are referred to the Quality Planning Team for review regarding their feasibility and benefits.
1.2.6  Continual Improvement

The planning, coordination and control of activities for continual improvement (QS11) is the responsibility of the Management Representative and the Quality Team. Continual improvement activities include - but are not be limited to - the following:

- activities of the Quality Team under the responsibility of the Management Representative
- actions on results from analysis of data (QS04)
- evaluation of suppliers (PU04)
- achievement of departmental quality objectives
- results from internal quality audits
- quality improvement proposals (QIP)
- corrective actions and preventive actions (CAR)
- periodic review of controlled documents (QS01)

The objectives of the corporate Quality Policy are taken into consideration for planning of improvement. During Management Reviews, the effectiveness of continual improvement is reviewed and opportunities for improvement are identified.

Manufacturing process improvement

It is the responsibility of personnel in Manufacturing to continually monitor the performance of manufacturing processes regarding conformity with product characteristics and process parameters. In monthly meetings with the production staff, process performances of production areas are analyzed, and opportunities for improvement are identified and implemented (QS11).

1.3 Communication

1.3.1  Internal Communication

Effective internal communication is essential for the proper functioning of the Quality Management System. The Management Representative, with the assistance of the Department Heads, ensures that required communication and information between departments and functions is defined in documented procedures, memos, forms and/or documents, and staff meetings.

Any communication problem regarding the Quality Management System is reported to the Management Representative for corrective action.

1.3.2  Customer Communication

In order to meet customer requirements and to ensure the proper and effective communication between the various departments within OPTEK and the customer, Sales establishes a list with some main contacts within OPTEK regarding customer inquiries. This list is updated as required, is distributed to functions concerned and is attached to the main directory available at the front lobby.
Internal and external communication related to the planning of products and processes (EG01) is defined by Engineering and/or Sales and/or the quality planning team, as applicable.

It is the responsibility of Engineering, with the assistance of the IS-department, to install and use electronic communication and design systems (such as CAD) that are compatible with the customers' systems, in order to effectively communicate and interchange information with the customers.

It is the responsibility of Sales, with the assistance of the IS-department, to develop, implement and maintain a computerized system (such as EDI) for the receipt of planning information of customer orders, shipping schedules and shipping information.

### 1.4 Internal Auditing

#### 1.4.1 Internal Audit of the Quality Management System

Following the established documented Process Map #002 Internal Quality Audits, the Management Representative is responsible for internal audits. Internal audits are planned and scheduled in such a way that all applicable clauses of ISO/TS 16949 and other additional quality system requirements are audited regarding compliance with this implemented Quality Management System and ISO/TS 16949 and ISO/IEC 80079-34. Audits do also verify if the Quality Management System is effectively implemented and maintained, and that it meets the requirements of OPTEK, including planned actions, objectives and results.

The Management Representative selects the auditors and ensures that they have required experience and knowledge to perform auditing activities.

Audits are scheduled on the basis of the importance of the activity to be audited. Audit activities are assigned to personnel not responsible for the area or activity to be audited.

Audit results are recorded and corrective action is taken as required. Where applicable, follow-up audits are conducted to ensure that corrective action was implemented and is effective.

Records of internal audits are maintained. As appropriate, management is informed of the results of audits and follow-up audits and takes additional corrective action. The Management Representative ensures that audit results are part of Management Review.

#### 1.4.2 Internal Audit Plans

Internal audits cover the entire Quality Management System and its processes, including all shifts of these processes, and are scheduled according to a yearly auditing plan and schedule prepared by the Management Representative (QS08). Due to special
circumstances, such as nonconformities and customer complaints, the auditing frequency is increased as appropriate.

1.4.3 Internal Auditor Qualification

It is the responsibility of the Management Representative to ensure that internal auditors of the Quality Management System have the necessary experience and qualification for performing internal quality audits (QS08). Training needs are identified and training is provided as required (QS02).

1.4.4 Manufacturing Process Audit

In addition to the normal internal audits of the Quality Management System, the Management Representative coordinates with the Manufacturing department the auditing of all manufacturing processes. Audits of manufacturing processes are performed at least once per year. Responsible personnel in Manufacturing take required corrective actions. A summary of audit results of the manufacturing processes is prepared and included in Management Reviews.

1.4.5 Product Audit

During the auditing of process in Manufacturing and the Warehouse, inspection and test results of product in process and finished product are audited to verify conformity to specified requirements. Incoming product, product in inventory and product ready for shipping is audited regarding compliance with packaging and labeling requirements. As appropriate, physical product can be inspected and tested by the auditor to confirm the product’s conformance to requirements and proper functionality.

1.5 Corrective & Preventive Action

1.5.1 Corrective Action

It is the responsibility of the Management Representative to implement and maintain the documented procedure QS10 Corrective Action that defines a corporate approach for corrective action.

Following the established procedure for corrective action (QS10), nonconformities are identified, root causes are determined, corrective action is evaluated and defined, recurrence of the nonconformity is prevented, corrective actions and their results are recorded, and the effectiveness of corrective action taken is reviewed. Corrective actions are appropriate to the importance and impact of the addressed nonconformity.

It is the responsibility of the Department Heads to inform the Sales/Service department of all customer complaints (QS06) and related corrective actions.

It is the responsibility of the Department Heads to establish and maintain records of corrective actions and their results.
1.5.1.1 Problem Solving

To determine the root cause of a problem or deficiency, and to establish required corrective action, a disciplined problem solving method as outlined in the work instruction QSW02, or any other suitable method, is used as appropriate.

1.5.1.2 Error-Proofing

As appropriate, the Quality Team applies error-proofing methods in the corrective action process to prevent recurrence of the problem.

1.5.1.3 Corrective Action Impact

As applicable, the Quality Team applies implemented corrective action to other similar processes or products in order to correct nonconformity.

1.5.1.4 Rejected Product Test / Analysis

Product returned from customers (QA08) is analyzed by Quality Assurance in order to initiate appropriate corrective action and to prevent recurrence (Process Map 010).

1.5.2 Preventive Action

It is the responsibility of the Management Representative to implement and maintain the documented procedure (QS09) Preventive Action that defines a corporate approach for preventive action to prevent the occurrence of potential nonconformities, deficiencies or problems. Any employee can suggest a preventive action to the responsible Department Head by initiating a CAR (QSW01).

The process of preventive action includes the following steps:
- identify potential nonconformities, deficiencies or problems
- determine the root causes
- determine the necessary preventive action
- implement the action
- follow-up on status and results
- review the effectiveness of preventive action.
- update all related documents with preventive action taken

Department Heads analyze and evaluate data of statistics (QS04) and perform periodic reviews of procedures (QS01) in order to detect deficiencies and problems and to take preventive action as required.

It is the responsibility of the Department Heads to establish and maintain records of preventive actions and their results. The Management Representative ensures that relevant information on preventive action is on the agenda of management reviews.
2 Realization Processes:

2.1 Requirements Determination & Review

2.1.1 Determination of requirements related to product

It is the responsibility of the Sales department to ensure that customer requirements related to product are identified and defined.

Potential new or modification to standard products will be reviewed per the New or Modified Standard Product Selection Procedure (99-00726-086) prior to the design and implementation into manufacturing.

Potential new or modification to existing ISO/IEC 80079-34 products will be reviewed per specification 99-00726-087 prior to the design and implementation into manufacturing.

It is the responsibility of Engineering, represented by the Quality Planning Team, to identify and determine requirements not specified by the customer but necessary for the proper and intended use of the product or service, as well as other requirements identified during product development and quality planning (EG01), including regulatory and statutory requirements Process Map #013 – Product Realization. Once these requirements are determined, they are used as input for product development and quality planning, and other functions concerned are informed as appropriate.

In addition to customer requirements included in design, development and quality planning (EG01), Department Heads ensure that other requirements specified by customers, as well as customer needs and expectations are identified, determined and documented by the responsible department, and that these requirements are met as appropriate. The Sales department or responsible Department Head also ensures that during set-up and maintenance of new customer files (217-0020-001), order taking and processing (217-0014-001, 217-0011-001), customer returns (217-0027-001) and shipping of products, customer requirements are identified and documented, and understood by all functions concerned.

Based on sales forecast and/or other special requirements documented by the Sales department, Manufacturing prepares production schedules and material requirement reports to ensure availability of product for the fulfillment of customer orders.

Optek has established an International Traffic in Arms Regulations (ITAR) compliance Manual (89-00015-179) that addresses legal and ethical compliance and more specifically, compliance with International Trade Regulations.

Customer-designated special characteristics

It is the responsibility of Engineering, the quality planning team and Manufacturing to apply, document and control special characteristics designated by the customer, with
focus on processes affecting safety, compliance with regulatory requirements, the fit or function of a product, or any other requirement of importance. Symbols to be used for these special characteristics are those designated by the customer or other commonly used symbols used in the industry.

2.1.2 Review of Requirements Related to Product, Manufacturing Feasibility
The Sales department is responsible for the review of product specifications and customer requirements (Process Map # 003, Process Map # 004, 217-0020-001) SDQA. Prior to the submission of a quotation (Process Map #011) to the customer, or the acceptance or confirmation of an order (Process Map 012) from a customer, the order or quotation is reviewed to ensure that
- the product and customer requirements are clearly defined and documented
- OPTEK has the capability to meet the requirements of the quotation or order
- requirements of verbal orders are recorded and confirmed prior to acceptance
- any differences between the customer’s order and OPTEK’s quotation are clarified and resolved.

Waiving the requirement for a formal review requires customer approval. Manufacturing feasibility is analyzed, and a risk analysis is performed, confirmed and documented (217-0020-001). The results of reviews and required actions are documented.

In the event of changes to product requirements, or other changes to a quotation or order, it is ensured that relevant documents and data are updated and that other functions concerned are notified. Records of contract reviews are maintained.

2.2 Design & Development

2.2.1 Design and Development
If there is a need for the design and/or development of a new custom product or for a change of the design or manufacturing process of a custom product, the originating department submits a Redfolder (217-0020-001) request to Engineering.

If there is a need for a design and/or development of a new standard product or for a change of the design or manufacturing process of a new standard product, the New or Modified Standard Product Selection Procedure (99-00726-086) will be followed.

2.2.2 Design and Development Planning
The planning and control of design and development of product is the responsibility of Engineering (EG01). During the planning process (Design Control Handbook), the project team determines the stages of the design and development project, defines the
review, verification and validation of each design and development stage as appropriate, and assigns responsibilities of required tasks and actions.

It is the responsibility of the Project Team Leader to ensure the effective communication and distribution of information between the team members and other functions participating in the planning process.

As the planning process develops, planning output is updated as appropriate.

2.2.3 Multidisciplinary Approach
Organizational and technical interfaces, including customer communication, are defined in the Project Plan and Schedule (Design Control Handbook) and are reviewed during each meeting of the team. As required, other functions are consulted within their areas of expertise. A multidisciplinary approach is also used for the development and monitoring of special characteristics, and the development and review of FMEAs and control plans.

2.2.4 Design and Development Input
The originator of the Greenfolder (217-0010-001) identifies and documents the input requirements, which are reviewed by Engineering. Input for product design and development includes functional and performance requirements, statutory and regulatory requirements, customer requirements, product quality and performance objectives, and any other identified requirements. Input for the development of manufacturing processes includes product design output data, targets for productivity, capability, cost, and customer requirements, as applicable (Design Control Handbook).

Any past experience or information from similar projects is applied as appropriate. Ambiguous, missing or conflicting information is clarified and resolved with the originator of the request before proceeding with the project. Records of design input are maintained.

2.2.5 Special Characteristics
Special characteristics for product and processes, and which are specified by the customer or by OPTEK, are identified and included in control plans, FMEAs and applicable documents in order to ensure proper identification of special requirements of product and processes (EG01).

2.2.6 Design and Development Output
The Project Team produces design output which is documented, is expressed in terms that can be verified and validated against design input requirements, meets design input requirements, contains or makes reference to acceptance criteria and includes critical and crucial characteristics for safety and functionality of the product or process. As applicable, design and development output provides data and information for product
design, manufacturing process design. Additional outputs for product design include FMEAs, special characteristics, product definitions, design reviews and other defined output results. Additional outputs for process design include drawings, FMEAs, control plans, process performance and other information and data to ensure that manufacturing processes meet requirements (Design Control Handbook).

Design outputs are reviewed prior to release.

### 2.2.7 Design and Development Review, Monitoring
The Project Team performs formal design reviews to identify any potential problems in meeting requirements and design goals. Problems are identified and appropriate action is taken.

Progress and end results of design and development planning (such as effectiveness, costs, lead time) are recorded at defined stages of the planning, and are submitted to the Management Representative for input to management reviews.

Records of design reviews and resulting actions are maintained.

### 2.2.8 Design and Development Verification
The Project Team performs periodic design verifications to verify that design and development outputs meet the design and development input requirements. Results of design verifications and resulting actions are recorded and maintained.

### 2.2.9 Design and Development Validation
The Project Team performs design validation to ensure that the designed product meets defined customer/user needs and requirements. Validation is according to customer requirements and includes program timing. If possible, this validation should be performed prior to production. However, if required, partial validation is acceptable. Results of validations and necessary actions are recorded and maintained.

### 2.2.10 Prototype Program
If required by the customer, the product development includes the development of a control plan and a prototype (EG01). Processes, equipment and materials used for the prototype should be the same as those used for final production runs. Testing activities are monitored regarding timely completion and compliance with requirements. In the event that services for prototype development are outsourced, it is understood that OPTEK is still responsible for the quality and performance of the prototype. As required, OPTEK provides technical assistance and support to contractors/suppliers.

### 2.2.11 Product Approval Process
Sample submission of production parts (QA10) for consequent customer approval is the responsibility of the Quality Assurance. Methods and guidelines specified by the customer are followed. Production part approval is requested for production parts, engineering change of production part, manufacturing location, material suppliers and
production process environment. Any change to these conditions requires customer notification and possible re-submission of production parts for approval. OPTEK is responsible for contracted materials and services.

As appropriate, production part approval is extended for engineering approval of purchased products.

The Quality Planning Team properly validates engineering changes.

### 2.2.12 Control of Design and Development Changes

Requests for design and development changes, including proposed changes from suppliers, are documented ([217-0010-001, Design Control Handbook](#)). Requests are reviewed and approved by Engineering. Results and necessary actions are documented and records are maintained.

### 2.3 Purchasing

#### 2.3.1 Purchasing Process

The Purchasing department is responsible for the effective and efficient operation of purchasing functions and activities ([PU01, PU02, PU03](#)).

Depending on the effect of the purchased product on the final product or production processes, the type of control applied to the supplier and the method used for verification of purchased product are identified and established by Engineering and/or Quality Assurance.

Production materials, products and services are only purchased from approved suppliers ([PU04](#)). Suppliers are evaluated and selected according to defined selection criteria and their ability to supply product that meets specified requirements. Records of evaluation and selection of suppliers as well as related actions are maintained by Supplier Quality Engineering (SQE) and/or Supplier Quality Development (SQD).

Supplier performance is monitored per the [Supplier Quality Requirements](#) paragraph 4 page 12 through the evaluation of product quality, problems reported by the manufacturing involving supplied product, delivery performance to acknowledged date and continual improvement.

#### 2.3.2 Regulatory Conformity

Quality Assurance verifies that incoming purchased products and materials are in compliance with applicable regulatory requirements ([QA02](#)).

#### 2.3.3 Supplier Quality Management System Development

The Purchasing staff encourages suppliers to prepare for and/or implement the necessary procedures in order to meet the requirements of ISO 9001:2000 and to become certified. All suppliers receive the manual TT Electronics/OPTEK [Supplier](#).
Quality Requirements as an initial step for getting familiar with quality system requirements. As required, the assistance of the Management Representative is requested. Purchasing &/or Supplier Development follows up on the suppliers' progress with the implementation of ISO 9001:2001 or an acceptable quality management system (Process Map: Supplier Development).

2.3.4 Customer-Approved Sources

Where according to customer contract, a product or material is to be purchased from a customer-designated supplier, it is the responsibility of the Purchasing department to ensure that these materials, products or services are only purchased from the customer-designated supplier. The customer must approve alternate suppliers for this product, material or service. Materials supplied by customer-designated suppliers are subject to receiving inspection by Quality Assurance (QA02). For the supply of materials for other applications, the customer-approved supplier must be approved according to OPTEK's approval criteria (PU04).

2.3.5 Purchasing Information

For products and services purchased, including customer supplied product, Purchasing, SQE and/or SQD ensures that required records are set up and maintained (PU02, PU03). The data describe and identify clearly the product to be ordered, requirements for the approval of product, procedures, processes and equipment, statutory and regulatory requirements, requirements for qualification of personnel, and Quality Management System requirements, as applicable. As appropriate, standards or other documents are referenced. The adequacy of specified purchase requirements is ensured prior to submission to the supplier.

2.3.6 Verification of Purchased Product and Incoming Product Quality

The extent of quality control exercised over a supplier or over the supplied product is determined by Quality Assurance and depends on the importance of the product or product class (PU03), the initial evaluation of the supplier, and/or type and extent of inspection performed by the supplier, and/or the results of ongoing performance ratings of the supplier. Incoming purchased product is submitted to an incoming inspection (QA02) performed and recorded by Quality Assurance when supplier product is non-certified.

In the event that OPTEK or one of OPTEK’s customers wants to verify purchased product at the supplier’s premises, these verification requirements and/or the method of product release are requested and defined by either Engineering, Quality Assurance or Sales, and are specified either in the purchase order and/or part specification. Verification activities at the supplier’s premises are coordinated through the Purchasing and/or Quality department. Verification at supplier’s site does not exclude subsequent rejection after receipt at OPTEK or OPTEK’s customer.
2.3.7 Supplier Monitoring

Supplier performance is monitored through evaluation of product quality, cost competitiveness, continuous improvement efforts, problems reported by manufacturing involving supplied product, delivery performance, and customer feedback.

Purchasing/Supplier Quality develops periodic performance ratings (PU04, Supplier Quality Requirements) of approved active suppliers and distributes on a regular basis to the top suppliers, others on request or at OPTEK’s discretion. Suppliers are informed of their rating and corrective action is taken as required. Records of supplier performance ratings are maintained.

2.4 Facilities & Equipment Management

2.4.1 Infrastructure

The required infrastructure and resources for manufacturing activities are identified during quality planning (EG01). As applicable, this includes building facilities, necessary workspace and utilities as well as needed equipment and services such as maintenance, warehousing and transportation.

Management ensures the timely availability of identified and approved resources.

2.4.1.1 Plant, facility and equipment planning

Plant, facility and equipment planning of the effectiveness of existing equipment and facilities are the responsibility of Manufacturing and involves departments and functions concerned. The productivity and effectiveness of existing operations is reviewed, monitored and evaluated considering

- human factors
- operator and line balance
- availability of supplies
- use of automation
- work plans

Records of planning activities are maintained as per applicable master list of records.

2.4.1.2 Contingency plans

The Management Representative, with participation from Manufacturing, Engineering and Sales, develops contingency plans to meet customer requirements in the event of a production halt or labor shortage. Contingency plans are reviewed in the first quarter of each year regarding their validity. New plans are developed as required.
2.4.2 Work Environment

2.4.2.1 Work Environment
The quality planning team defines special conditions of the work environment that are necessary for the processes to meet defined requirements of product quality and regulatory requirements (EG01). These special conditions are included in the quality plan, manufacturing plan, process sheet or other documents. It is the responsibility of the Department Head to implement these requirements.

2.4.2.2 Personnel Safety to Achieve Product Quality
It is the responsibility of the Department Heads to ensure the safety of employees and to minimize risk of injuries when performing their duties. Accidents at the workplace are recorded (First Report of Injury & Supervisors Investigation Report) with copy to the HR Representative who keeps a master list of accidents for corrective or preventive actions.

The HR Representative forms the Health and Safety Committee that includes representatives of applicable areas. Any issues or concerns regarding health and safety of processes are reported to the departmental representative.

Product safety is addressed during the design and development process under the responsibility of Engineering.

2.4.2.3 Cleanliness of Premises
It is the responsibility of management to ensure that the premises of OPTEK are kept clean and in a good state of order. It is the responsibility of Manufacturing and Warehouse to ensure that production facilities and the warehouse are kept clean and in good order. As required, housekeeping procedures are developed and implemented by individual Department Heads.

2.4.3 Customer Property

2.4.3.1 Customer Property
Customer owned product supplied for production is inspected by Quality Assurance according to defined inspection requirements (QA02).

The responsible Department Head ensures that customer owned product is identified, stored, used, handled and shipped in an appropriate manner in order to ensure its suitable condition for use.

During periodic cycle counts conducted by designated personnel in Manufacturing or Warehousing, a visual inspection of products, including customer owned product, is performed to verify the product’s condition and proper identification. Any loss, damage or deterioration of customer-supplied product is recorded and the customer is notified.
2.4.3.2 Customer Owned Production Tooling

It is the responsibility of the Toolroom & Manufacturing to ensure that customer owned tooling and fixtures are clearly identified with a metal plate or permanent marking method showing the ownership of the equipment.

2.4.4 Control of Monitoring & Measurement Devices

2.4.4.1 Control of Monitoring and Measuring Devices

To ensure accurate and reliable monitoring and inspection results, the Quality Assurance, Manufacturing, Toolroom and Engineering ensure that monitoring and measuring equipment and devices are controlled, calibrated and maintained (QA05).

The type of monitoring and measuring equipment/device/software to be used in Manufacturing, by Quality Assurance and Toolroom, and the required accuracy of these monitoring and measurement activities are defined during quality planning and specified in the manufacturing plan (EG01), process traveler and/or inspection reports (QA02, QA03, QA04).

It is the responsibility of the applicable department to ensure that monitoring and measuring processes are capable for their intended purpose and are performed in a manner that is consistent with requirements.

To ensure valid results, measuring equipment is

- calibrated and/or checked in defined intervals or prior to use, and according to a recognized standard; where no recognized standard is used, the basis applied for the calibration is documented.
- adjusted and re-adjusted as necessary to ensure required accuracy
- identified with a unique identification number and the current calibration status.
- kept in a secure and restricted location to prevent misuse and improper adjustments that could invalidate calibration settings.
- protected from damage and deterioration during handling, maintenance and storage

In the event that monitoring and measuring devices are found out of calibration, previous measuring results are reviewed regarding their validity (QA06). Corrective action on the measuring device or product affected is taken, including recall of nonconforming product, if required (MF03, QA01).

Prior to the use of computer software for monitoring and measuring activities, it is verified and confirmed that the software produces defined results. Records of these verifications are maintained.
2.4.4.2 Measurement System Analysis

It is the responsibility of Quality Assurance to ensure that a Repeatability and Reproducibility study is conducted (QA07) for each measuring device referenced in control plans. Records of these studies provide evidence of the variations present in the results of each type of measuring device and are taken into consideration when inspection reports are developed.

2.4.4.3 Calibration / Verification Records

The department performing the calibration of monitoring and measuring devices is responsible for the record keeping of calibration activities. These records include the identification of the equipment and the calibration standard, revisions due to engineering changes, and calibration results such as out-of-specification/conformity to specifications.

When monitoring and inspection equipment is found out-of-specifications, the impact on products previously measured with this equipment is reviewed and validated (QA06) and an Out-Of-Calibration Report is initiated as appropriate. If suspect product/material has been shipped, the customers are informed and the product is recalled as required (QA06).

2.4.4.4 Laboratory Requirements

2.4.4.4.1 Internal Laboratory

It is the responsibility of Quality Assurance to define and document the scope of the capability of tests and inspection activities, which can be performed by the in-house laboratory facility of the Quality Assurance department. Quality Assurance has specified and implemented technical requirements for the

- suitability of implemented procedures
- competency of personnel
- testing of product
- capability to perform these services correctly and according to pertinent process standards,
- the review of related records

2.4.4.4.2 External Laboratory

As required, qualified laboratories are used for inspection, test and calibration, and other contract services. Only laboratories are used which include in their defined laboratory scope the required service to be performed, and which are either capable based on business history and/or previous services provided, or laboratories that are accredited to ISO/IEC 17025 or other equivalent national standard. If a specific calibration service cannot be performed by an external laboratory, or in the absence of such a laboratory, the original equipment manufacturer or their accredited representative can provide this service, as long as the requirements of clause 7.6.3.1 of ISO/TS 16949 are met.
2.5 Company Control Specifics

2.5.1 Document Control

2.5.1.1 Control of Documents and Engineering Specifications

Documents required by the Quality Management System are controlled documents.

It is the responsibility of the Management Representative to implement and maintain the documented procedure (Process Maps #005, #006, #007, #015, #016) Control of Documents, which defines the responsibilities for the development of controlled documents, their approval for adequacy, changes and re-approval, revision status, document formats, identification and distribution.

Following the documented procedure EG02 Document and Data Control, the Engineering Department is responsible for the identification, control and distribution of technical engineering documents, including documents and data of external origin such as standards and customer drawings. Engineering documents developed by Engineering or engineering documents from the customer, including the distribution of these documents, are recorded. Incoming customer engineering standards and specifications, including changes, are reviewed as soon as possible, not to exceed two weeks, by Engineering, and are then distributed and implemented as required. Records of implementation dates in production are maintained.

It is the responsibility of the applicable Department Head to ensure that current revisions of controlled documents are legible, readily available where needed, that obsolete copies are replaced and destroyed or invalidated, and that obsolete documents retained for any purpose are clearly identified.

As required, it is the responsibility of the IS Department to establish a schedule for producing back-ups of defined computer data. These back-ups on tape or CD-ROM are kept in a secure place outside of the company’s premises. Schedules for back-up responsibilities and compliance are documented.

2.5.2 Records Control

2.5.2.1 Control and Retention of Records

Records are maintained to provide evidence of activities and their results, of conformance to requirements and of the effective operation of the Quality Management System. Department Heads are responsible for the proper identification, storage, retrieval, protection, retention time and disposition of records according to the established documented procedure QS12.
2.5.3 Control of Production Provision

2.5.3.1 Control of Production

Manufacturing processes activities are performed under controlled conditions. Based on the output from quality planning (EG01), Manufacturing and Document Control ensure that the necessary documents, data and operating instructions for the performance of manufacturing processes are developed and available to personnel. These documents or data describe in sufficient detail the product characteristics, production processes, the equipment to be used, as well as the activities for monitoring and measuring of these processes. Included are procedures for release, delivery and post-delivery activities.

The Manufacturing department ensures that operating instructions, including instructions for special processes, are available at the workstation, that production activities, verification results and SPC records are recorded, and that activities for the monitoring and measurement of production processes are implemented and followed.

It is also the responsibility of the Manufacturing department to ensure that the work environment is appropriate for the work being performed and meets statutory requirements. The HR Representative is responsible for compliance with regulatory requirements (MF09).

2.5.3.2 Control Plan

Control plans are developed during quality planning (EG01) and define the development of prototypes, pre-launch and production processes, as applicable. Control plans are available for all production stages of products or parts, including assembly. Pre-Launch control plans heed the outputs of DFMEAs and PFMEAs.

As applicable, control plans specify

- the required controls for manufacturing processes
- the methods used for monitoring applied controls over special characteristics (customer/OPTEK)
- customer-required information
- the reaction plan to be initiated when the process becomes unstable or not capable

With changes of product specifications, or any changes affecting the product, manufacturing processes, inspection activities, logistics, supply sources or FMEAs, control plans are updated by the quality planning team (EG01). If required, customer approval is obtained for the change.
2.5.3.3 Work Instructions

It is the responsibility of the Department Heads to develop and maintain documented work instructions and operating instructions that are necessary for the performance of processes and activities affecting quality of products. These documents are made accessible to personnel at the work place. Work instructions and operating instructions are derived from the output data from quality planning, such as the quality plan or control plan.

2.5.3.4 Verification of job set-ups

Manufacturing is responsible for proper set-ups of production equipment. In case of set-up difficulties, Quality Assurance performs a last-off comparison, as appropriate. First-Offs are approved by Quality Assurance. As applicable, statistical verification methods are used.

2.5.3.5 Preventive and Predictive Maintenance

A master list of machinery and equipment that requires preventive maintenance to ensure continuous process capability is developed and maintained by Manufacturing. Preventive maintenance objectives are established and documented in the first quarter of each year. These objectives are evaluated at least yearly regarding their achievement and opportunities for improvement.

Designated staff in Manufacturing performs required preventive maintenance. The actual maintenance status of each piece of equipment is identified. The maintenance system includes a MIN/MAX - inventory system of frequently used replacement parts and a predictive maintenance analysis that assists in the review of preventive maintenance cycles, maintenance methods and inventory of replacement parts. Equipment and tooling and gauging that are kept in Manufacturing are packaged and preserved according to manufacturer’s guidelines and recommendations.

2.5.3.6 Management of Production Tooling

As applicable, the departments of Manufacturing, Engineering, Toolroom and Quality Assurance are responsible for the design, construction, review and approval of production tooling and fixtures.

- The Toolroom is responsible for the construction of tooling and fixtures (TR01). Established controls are followed to effectively coordinate all activities, including a full dimensional inspection of the tooling and the monitoring of timely completion.
- The status or availability of tooling and fixtures is clearly identified (TR03).
- The Mexico Toolroom is responsible for preventive maintenance (TR03), repair (TR03, TR04), storage and recovery (TR03) of production tooling.
- The set-up of tooling and equipment is the responsibility of Manufacturing (MF07).
• Manufacturing establishes programs for changes of perishable tools in production.
• The Engineering department is responsible for design changes of tooling and fixtures, including engineering change level. As required, these changes are passed on to the Toolroom for implementation.
• Tool modification and revision to documentation are coordinated between Engineering and the Toolroom.
• As required, the Quality Planning Team is involved in the planning of changes to tooling and fixtures.

In the event that design or construction is contracted to outside sources (EG04), a tracking and follow-up system is put in place by Engineering or Manufacturing.

Customer-supplied tooling is inspected and approved by Manufacturing, Toolroom and Quality Assurance.

2.5.3.7 Production Scheduling
Manufacturing is responsible for production scheduling. The production scheduling of custom-made parts is forecasted.

Quarterly, the inventory turnover rate is calculated and corrective action is taken in case that the turnover rate is below the established minimum.

2.5.3.8 Feedback of Information
Based on customer feedback provided by Sales, the Management Representative issues quarterly statistics (QS04) which are analyzed by the heads of Engineering, Sales, Manufacturing and Quality Assurance. Corrective or preventive action is taken as required.

2.5.4 Validation of Processes
Validation of processes for production
Where the resulting process output cannot be verified through monitoring or measurement, the Quality Planning Team validates production processes with the assistance of Manufacturing and Engineering, regarding their ability to achieve planned results.

The quality planning team establishes procedures for the review, approval and requirements of these processes, including - as applicable: criteria for review and approval, approval of equipment and qualified personnel, the use of methods and procedures, required records, and re-validation in case that expected results are not achieved.

Attention is given to special processes where the results cannot be verified through measurement or testing, where deficiencies become apparent when the product is already in use.
2.5.5  Identification of Traceability

Identification and traceability
Designated personnel in Quality Assurance, Manufacturing and Warehouse identify incoming product and material, product and material during production, and product and material in storage with the product identification and inspection status (MF06, QA02, QA03, QA04, WH02).

Using the implemented computerized system in Manufacturing, sensors manufactured by OPTEK are traceable by date codes.

2.5.6  Preservation

2.5.6.1 Preservation of Product
It is the responsibility of Manufacturing and the Warehouse to ensure the proper identification, handling, packaging, storage and protection of product and materials during receiving, handling and storage, shipping and production. This includes constituent parts of a product.

Temperature sensitive products and materials are stored in the temperature-controlled room.

2.5.6.2 Storage and Inventory
During periodic cycle counts, the condition of materials and products in the warehouse is verified to ensure that any deterioration or damage is detected and recorded, and that required corrective action is taken.

The MRP-system in Manufacturing is used to ensure optimized inventory turns over time, minimum inventory levels and appropriate stock rotation (FIFO) of product and raw materials. Manufacturing is responsible for keeping established inventory levels of finished product, using the computerized production scheduling system.

The inventory turnover rate is periodically reviewed and corrective action is taken in the event that the turnover rate is below the established minimum.

When processing shipping orders (217-0014-001), the staff in Warehousing ensures that FIFO is applied.

2.5.7  Monitoring & Measuring of Processes

2.5.7.1 Monitoring and Measurement of Processes
During Quality Management System Planning, and based on statistics of operational performance (QS04) and the achievement of quality objectives, the processes of the Quality Management System are analyzed by the Management Representative and responsible Department Heads regarding their effectiveness. As required, corrective action is implemented to achieve planned results and product conformity, to correct
nonconformities or to improve the operational effectiveness and efficiency of the processes of the Quality Management System.

### 2.5.7.2 Monitoring and Measurement of Manufacturing Processes

To verify process capability and provide additional input for process control, the quality planning team arranges for the monitoring of new and modified manufacturing processes (EG01). Results are documented and include instructions for production processes, verification and maintenance as well as objectives for manufacturing process capability, reliability, maintainability and availability.

Manufacturing ensures that processes are implemented according to control plans and other applicable procedures or documents in order to ensure that process capability and process performance is maintained according to customer part approval process requirements.

Control plans and process flow diagrams are implemented, including adherence to specified measurement techniques, sampling plans, acceptance criteria and reaction plans.

It is the responsibility of Manufacturing to monitor process capability and to ensure that process capability and performance is according to applicable control plans. In case of nonconformity of processes, defined reaction plans are followed.

Important events that are occurring during production, such as down times are recorded.

If identified characteristics on the control plan become unstable or non-capable, the applicable reaction plan is followed. If appropriate, these reaction plans include containment of produced parts or products and 100% inspection. Corrective action is taken as per established procedure QS10 in order to restore required process capability and product quality. If required, these corrective action plans are reviewed with and approved by the customer.

Effective dates of process changes are documented by Manufacturing.

### 2.5.8 Monitoring & Measuring of Product

#### 2.5.8.1 Monitoring and Measurement of Product

It is the responsibility of Quality Assurance to establish and maintain procedures and inspection reports for receiving inspection (QA02), in-process inspection (QA03) and final inspection (QA04) of product and materials.

Product is not released until all specified requirements have been met, unless otherwise approved by an authorized function - and where applicable by the customer.
The warehouse staff performs a visual inspection of outgoing product to ensure that the product and packaging is in good condition and that marking and labeling requirements are met.

In the event that purchased product is released for urgent production prior to inspection and acceptance by Quality Assurance, the product is recorded and controlled in order to permit recall and replacement in case of nonconformity of the product.

Product that does not meet specified requirements is rejected and quarantined per established documented procedure.

As required, Quality Assurance selects accredited laboratories for certain inspection or testing activities. Records of these inspection results are verified, reviewed and maintained.

Inspection results are recorded and records are maintained. These inspection records document acceptance criteria, inspection results, whether the product was accepted or rejected and the inspection authority responsible for the product release.

### 2.5.8.2 Dimensional Inspection and Functional Testing

At least once every twelve months, or as otherwise specified by the customer, Quality Assurance performs a dimensional inspection and functional verification (QA12) for each product specified in control plans. Results are available to the customer upon request.

### 2.5.9 Control of Nonconforming Product

#### 2.5.9.1 Control of Nonconforming Product and Reworked Product

Nonconforming product and product without proper identification is quarantined and controlled according to the documented procedure QA01. The nonconformity of the product is verified and confirmed by Quality Assurance and verification results and recommended disposition or action are recorded. Functions concerned are notified.

Quality Assurance, Manufacturing, Engineering or Sales review and authorizes the release of quarantined product for its final disposition, according to the following options:

- rework to meet specified requirements
- accept with or without repair by concession
- re-grade for alternative applications
- reject or scrap
If the acceptance with or without repair requires the concession of the customer or the approval or permit of a regulatory body or other authority, Manufacturing ensures that the required concession is received prior to initiation of the repair.

Qualified personnel in Manufacturing process rework Orders. Detailed instructions for required rework are available to operators in work instructions.

Reworked product is re-inspected by Quality Assurance.

As appropriate and required, the customer is notified by Sales of the proposed use or repair of nonconforming product. Where applicable, Manufacturing ensures that the reworked product is identified with the actual condition of the product, including the customer’s release authorization.

Records of nonconforming product, including the type of nonconformity, actions taken and concessions obtained are maintained in First Time Quality reports.

In the event that nonconforming product is detected after the product was shipped to the customer, or after its use in production or service, the Engineering department with Manufacturing & Quality Assurance and Sales department analyze the impact of the nonconformity and take appropriate action. As required, the customer is informed and the nonconforming product is recalled.

Nonconforming purchased product and material is returned to the supplier with a Supplier Corrective Action Request (Process Map 009) issued by Supplier Quality Assurance (QA11).

### 2.5.9.2 Customer Waiver

In the event that manufactured product, or purchased product, or manufacturing processes are different from the product or process approved by the customer, or that temporary change to product and processes is required, the request for temporary change or deviation is submitted to Engineering for approval (EG01). As required, customer production part approval is obtained (QA10).

Concessions outside of original design are not permitted by ISO/IEC 80079-34

Manufacturing keeps records of expiration dates and quantities of authorized deviations and ensures that normal production activities are re-instated after expiration of engineering deviations.

Products manufactured and shipped on customer authorization are identified as such on each packaging unit or container.
## Approvals & Revision History

### OPTEK Quality Manual - revision control

<table>
<thead>
<tr>
<th>Page No.</th>
<th>Reference</th>
<th>Revision</th>
<th>Date</th>
<th>Description of Change</th>
<th>Initials</th>
</tr>
</thead>
<tbody>
<tr>
<td>All (1-45)</td>
<td>1st issue</td>
<td>0</td>
<td>2004/Mar/11</td>
<td>No change - first issue</td>
<td></td>
</tr>
<tr>
<td>35, 44</td>
<td>051689</td>
<td>A</td>
<td>2007/May/3</td>
<td>Added prototype process maps #015, #016 to section 2.5.1.1</td>
<td>M.T.</td>
</tr>
<tr>
<td>6, 10</td>
<td>052767</td>
<td>B</td>
<td>2009/March/18</td>
<td>Change Jerry Gallagher to Bob Taber. Delete Magnetoresistive and add UV.</td>
<td>L.T.</td>
</tr>
<tr>
<td>26, 27</td>
<td>052939</td>
<td>C</td>
<td>2009/July/10</td>
<td>Update sections 2.1.1 (under Requirements Determination &amp; Review), 2.2.1 (Design and Development)</td>
<td>L.T.</td>
</tr>
<tr>
<td>1, 7, 10, 26</td>
<td>053405</td>
<td>D</td>
<td>2010/ Feb/23</td>
<td>Updated ref. to ISO/TS 16949 to 2009 rev &amp; ISO9001 to 2008 rev. Updated Corporate Quality Policy – added ITAR Compliance Manual hyperlink in para 2.1.1. Adding numeric revision to all process maps.</td>
<td>R.B.</td>
</tr>
<tr>
<td>7, 8, 17, 23, 26, 43, 46</td>
<td>053950</td>
<td>E</td>
<td>2011/June 16</td>
<td>Updated to include IECEx/EN13980 Baseefa references. Update 003, 005, 006, 007, 012 &amp; 013 Process Maps from HP to SAP.</td>
<td>L.T.</td>
</tr>
<tr>
<td>9, 47</td>
<td>054425</td>
<td>F</td>
<td>2012/August 17</td>
<td>Updated IECEx Certificate of Conformity</td>
<td>L.T.</td>
</tr>
<tr>
<td>All</td>
<td>054919</td>
<td>G</td>
<td>2014/March/28</td>
<td>Change Bob’s title from President/CEO to Vice President/General Manager. Update Process Maps 003, 004, 010, 011 &amp; 012 and operating procedure QS07. Update the TT Electronics logo.</td>
<td>L.T.</td>
</tr>
<tr>
<td>Process Maps 003 &amp; 011</td>
<td>055002</td>
<td>H</td>
<td>2014/July/8</td>
<td>Updated process maps #003 to rev 5 and #011 to rev 3</td>
<td>L.T.</td>
</tr>
<tr>
<td>Process Maps 003 &amp; 011</td>
<td>055157</td>
<td>I</td>
<td>2015/January/28</td>
<td>Updated process maps #003 to rev 6 and #011 to rev 4</td>
<td>L.T.</td>
</tr>
<tr>
<td>Process Map 010</td>
<td>055179</td>
<td>J</td>
<td>2015/April/13</td>
<td>Updated process map #010 to rev 3, only first page of process map changed.</td>
<td>L.T.</td>
</tr>
<tr>
<td>26, 43</td>
<td>055229</td>
<td>K</td>
<td>2015/April/23</td>
<td>Clarify process map #s under section 1.5.1.4 change (Process Map – Process Flow) to (Process Map 010) and under section 2.5.9.1 (Process Map: SCAR Process) to (Process Map 009).</td>
<td>L.T.</td>
</tr>
<tr>
<td>5, 7, 18, 24, 27, 44, 47</td>
<td>055317</td>
<td>L</td>
<td>2015/July/27</td>
<td>Replace all references to EN13980/IECEcOD005 to ISO/IEC 80079-34.</td>
<td>L.T.</td>
</tr>
<tr>
<td>16</td>
<td>055319</td>
<td>M</td>
<td>2015/July/31</td>
<td>Add the maximum interval between management reviews to section 1.2.1.</td>
<td>L.T.</td>
</tr>
</tbody>
</table>
Appendix A: ISO/TS Cross Reference to this Manual

<table>
<thead>
<tr>
<th>ISO/TS16949 Clause #</th>
<th>Description</th>
<th>OPTEK Quality Manual Paragraph</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.1</td>
<td>Quality Management System</td>
<td>1.1.1</td>
</tr>
<tr>
<td>4.1.1</td>
<td>QMS General Requirements</td>
<td>1.1.1</td>
</tr>
<tr>
<td>4.2.1</td>
<td>QMS General Requirements</td>
<td>1.1.1</td>
</tr>
<tr>
<td>4.2.2</td>
<td>Quality Manual</td>
<td>About this Manual</td>
</tr>
<tr>
<td>4.2.2a</td>
<td>Scope</td>
<td>Scope</td>
</tr>
<tr>
<td>4.2.3</td>
<td>Document Control</td>
<td>2.5.1</td>
</tr>
<tr>
<td>4.2.3.1</td>
<td>Engineering Specifications</td>
<td>2.5.1.1</td>
</tr>
<tr>
<td>4.2.4</td>
<td>Records Control</td>
<td>2.5.2</td>
</tr>
<tr>
<td>4.2.4.1</td>
<td>Records Retention</td>
<td>2.5.2.1</td>
</tr>
<tr>
<td>5.1</td>
<td>Management Commitment</td>
<td>1.1.2</td>
</tr>
<tr>
<td>5.1.1</td>
<td>Process efficiency</td>
<td>1.1.2</td>
</tr>
<tr>
<td>5.2</td>
<td>Customer Focus</td>
<td>1.1.3</td>
</tr>
<tr>
<td>5.3</td>
<td>Quality Policy</td>
<td>1.1.4</td>
</tr>
<tr>
<td>5.4.1</td>
<td>Quality Objectives</td>
<td>1.2.2</td>
</tr>
<tr>
<td>5.4.1.1</td>
<td>Quality Objectives - Supplemental</td>
<td>1.2.2</td>
</tr>
<tr>
<td>5.4.2</td>
<td>QMS Planning</td>
<td>1.1.5</td>
</tr>
<tr>
<td>5.5.1</td>
<td>Responsibility, Authority &amp; Communication</td>
<td>1.1.6</td>
</tr>
<tr>
<td>5.5.1.1</td>
<td>Responsibility for Quality</td>
<td>1.1.6</td>
</tr>
<tr>
<td>5.5.2</td>
<td>Management Representative</td>
<td>1.1.7</td>
</tr>
<tr>
<td>5.5.2.1</td>
<td>Customer Representative</td>
<td>1.1.7</td>
</tr>
<tr>
<td>5.5.3</td>
<td>Internal Communication</td>
<td>1.3.1</td>
</tr>
<tr>
<td>5.6</td>
<td>Management Review</td>
<td>1.2.1</td>
</tr>
<tr>
<td>6.1</td>
<td>Provision of Resources</td>
<td>1.1.8</td>
</tr>
<tr>
<td>6.2.1</td>
<td>Human Resources</td>
<td>1.1.9</td>
</tr>
<tr>
<td>6.2.2</td>
<td>Competence, awareness &amp; training</td>
<td>1.1.9.1</td>
</tr>
<tr>
<td>6.2.2.1</td>
<td>Product design skills</td>
<td>1.1.9.2</td>
</tr>
<tr>
<td>6.2.2.2</td>
<td>Training</td>
<td>1.1.9.3</td>
</tr>
<tr>
<td>6.2.2.3</td>
<td>Training on the job</td>
<td>1.1.9.4</td>
</tr>
<tr>
<td>6.2.2.4</td>
<td>Employee motivation &amp; empowerment</td>
<td>1.1.9.5</td>
</tr>
<tr>
<td>6.3</td>
<td>Infrastructure</td>
<td>2.4.1</td>
</tr>
<tr>
<td>6.3.1</td>
<td>Plant, facility &amp; equipment planning</td>
<td>2.4.1.1</td>
</tr>
<tr>
<td>6.3.2</td>
<td>Contingency plans</td>
<td>2.4.1.2</td>
</tr>
<tr>
<td>6.4</td>
<td>Work Environment</td>
<td>2.4.2.1</td>
</tr>
<tr>
<td>6.4.1</td>
<td>Personnel safety to achieve product quality</td>
<td>2.4.2.2</td>
</tr>
<tr>
<td>6.4.2</td>
<td>Cleanliness of premises</td>
<td>2.4.2.3</td>
</tr>
<tr>
<td>7.1</td>
<td>Planning of Product Realization</td>
<td>1.1.10</td>
</tr>
<tr>
<td>7.1.1</td>
<td>Planning of product realization - supplemental</td>
<td>1.1.10.1</td>
</tr>
</tbody>
</table>
7.1.2 Acceptance criteria 1.1.10.2
7.1.3 Confidentiality 1.1.10.3
7.1.4 Change Control 1.1.10.4
7.2.1 Determination of Requirements 2.1.1
7.2.1.1 Customer designated special characteristics 2.1.1
7.2.2 Review of Requirements related to the product 2.1.2
7.2.2.1 Review of Requirements related to the product - Suppl. 2.1.2
7.2.2.2 Organization manufacturing feasibility 2.1.2
7.2.3 Customer Communication 1.3.2
7.3 Design & Development 2.2.1
7.4 Purchasing 2.3.1
7.5.1 Control of Production Provision 2.5.3
7.5.2 Validation of Processes 2.5.4
7.5.3 Identification & Traceability 2.5.5
7.5.4 Customer Property 2.4.3
7.5.5 Preservation 2.5.6
7.6 Control of Monitoring & Measurement Devices 2.4.4

8 Measurement, Analysis & Improvement
8.1 Measurement Analysis & Improvement: General 1.2.3
8.2.1 Customer Satisfaction 1.2.4
8.2.2 Internal Auditing 1.4.1
8.2.3 Monitoring & Measuring of Processes 2.5.7
8.2.4 Monitoring & Measuring of Product 2.5.8
8.3 Control of Nonconforming Product 2.5.9
8.4 Analysis of Data 1.2.5
8.5 Continual Improvement 1.2.6
8.5.2 Corrective Action 1.5.1
8.5.3 Preventive Action 1.5.2

Appendix B: ISO/IEC 80079-34 ATEX Certification References

<table>
<thead>
<tr>
<th>ISO/TS 16949 Clause #</th>
<th>QMS Description</th>
<th>QMS Section</th>
<th>QMS Page #</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.1 Management Processes – QMS</td>
<td>QMS Table of Contents</td>
<td>Table of Contents</td>
<td>5</td>
</tr>
<tr>
<td>4.1</td>
<td>Applicable Standards</td>
<td>Applicable Standards</td>
<td>7</td>
</tr>
<tr>
<td>4.2.4</td>
<td>Records Control</td>
<td>2.5.2</td>
<td>37</td>
</tr>
<tr>
<td>4.2.4.1</td>
<td>Records Retention</td>
<td>2.5.2.1</td>
<td>37</td>
</tr>
<tr>
<td>5.1</td>
<td>Management Commitment</td>
<td>1.2.1.1</td>
<td>18</td>
</tr>
<tr>
<td>5.5.1</td>
<td>Responsibility, Authority &amp; Communication</td>
<td>1.1.6</td>
<td>12</td>
</tr>
<tr>
<td>5.6.1</td>
<td>Management Review Input</td>
<td>1.2.1.1</td>
<td>18</td>
</tr>
<tr>
<td>5.6.2</td>
<td>Internal Auditing</td>
<td>1.4.1</td>
<td>24</td>
</tr>
<tr>
<td>7.1</td>
<td>New Product Requirements</td>
<td>2.1.1</td>
<td>27</td>
</tr>
<tr>
<td>7.2</td>
<td>Customer Waivers</td>
<td>2.5.9.2</td>
<td>44</td>
</tr>
</tbody>
</table>