IECEx OPERATIONAL DOCUMENT

IEC System for Certification to Standards relating to Equipment for use in Explosive Atmospheres (IECEx System)

IECEx Certified Equipment Scheme –

Guidelines on the Management of Assessment and Surveillance programs for the assessment of Manufacturer’s Quality Systems, in accordance with the IECEx Scheme.
Introduction

The IECEx System provides for the assessment and surveillance of Manufacturer’s Quality Management Systems and the issue of an IECEx Quality Assessment Report in accordance with ISO/IEC 80079–34.

The purpose of this document is to provide guidance upon the methodology, thereby, providing the opportunity for IECEx Certification Bodies to conduct assessments and audits of manufacturer’s quality systems in a uniform manner.

This document is based on work conducted by the European Notified Bodies Group for ATEX and in turn is based upon ISO 19011:2002 Guidelines for quality and/or environmental management systems auditing.

NOTE ISO 19011: 2002 uses the term “audit” throughout but IECEx 02 uses the separate terms “assessment” and “surveillance” and “audit”. For the purpose of this document the term “audit” is considered to be an activity that comprises part of the processes of applicable to both “assessment” and “surveillance”.

This version was issued to replace references to OD 005-1 with ISO/IEC 80079–34: 2011 in accordance with ExTAG/247A/INF. Other changes are indicated by a red margin bar.

This Edition 3.1 as circulated as ExMC/1264/DV and subsequently approved by the 2017 ExMC Meeting (via Decision 2017/42 that incorrectly refers to Edition 3.2) replaces Edition 3.0 upon publication.

This REDLINE VERSION displays changes as compared to Edition 3.0 with side bars and added text shown as text and deleted text shown as text.
1 Scope
This document provides guidance on the fundamentals of conducting assessment and surveillance visits as a means of verifying a manufacturer’s compliance with the IECEx quality system requirements.

2 Normative references
ISO 9000: 2005 Quality management systems -- Fundamentals and vocabulary
ISO19011: 2002 Guidelines for quality and/or environmental management systems auditing.
ISO/IEC 80079-34: 2011 Explosive atmospheres – Part 34: Application of quality systems for equipment manufacture

3 Terms and Definitions

3.1 Audit
An activity based on a systematic, independent and documented process for obtaining records, statements of fact or other relevant information and assessing them objectively to determine the extent to which specified requirements are fulfilled.

NOTE: Whilst “audit” applies to management systems, “assessment” applies to conformity assessment bodies as well as more generally. The assessment process includes auditing.

3.2 Product Audit
An audit (3.1) to determine whether the product is in compliance with the type described in the IECEx Assessment and Test Report.

3.3 Audit Criteria
Set of policies, procedures or requirements used as a reference.

3.4 Audit Evidence
Records, statements of fact or other information, relevant to the audit criteria (3.3) and which are verifiable.

NOTE Audit evidence can be qualitative or quantitative.

3.5 Audit Finding(s)
Result of the evaluation of the collected audit evidence (3.4) against audit criteria (3.3).

NOTE Audit findings can indicate either conformity or nonconformity with audit criteria.

3.6 Audit Conclusion(s)
Outcome of an audit (3.1), reached by the audit team after consideration of the audit objectives and all audit findings (3.5).
3.7 Audit Client
Organisation or person requesting an audit (3.1).

3.8 Auditee
Organisation being audited.

3.9 Auditor
Person with the competence (3.15) to conduct an audit (3.1).

3.10 Audit Team
One or more auditors conducting an audit (3.1).

NOTE 1 One auditor of the audit team is appointed as audit team leader.
NOTE 2 The audit team can include auditors-in-training and, where required, technical experts.
NOTE 3 Observers can accompany the audit team but do not act as part of it.

3.11 Technical Expert
Person who provides specific knowledge or expertise to the lead auditor with respect to the subject being audited.

NOTE 1 Specific knowledge or expertise includes those on the organisation, process, product, or activity to be audited, as well as language or cultural guidance.
NOTE 2 A technical expert does not act as an auditor in the audit team.

3.12 Audit Programme
Set of one or more audits (3.1) planned for a specific time frame and directed toward a specific purpose.

3.13 Audit Plan
Description of the on-site activities and arrangements for an audit (3.1).

3.14 Audit Scope
Extent and boundaries of an audit (3.1).

NOTE The scope typically includes a description of physical locations, organisational units, activities and processes, as well as the time period covered.

3.15 Competence
Demonstrated capability to apply knowledge and skills.

3.16 Initial Assessment
Activities related to the notification of a manufacturer to determine whether the manufacturer and applicable manufacturing locations(s) and production sites(s) meet all the requirements of the relevant clauses of the specified standard necessary for granting notification as to whether they have effectively implemented, including documentation review, site audit at the manufacturers’s premises, manufacturing locations(s) and production site(s), preparation and
consideration of the audit report and other relevant activities necessary to provide sufficient information to allow a decision to be made as to whether notification shall be granted. Audit results shall be recorded on a Quality Assessment Report (QAR).

3.17 Surveillance
Surveillance of a manufacturer’s quality system takes place on a regular basis as defined in this document. The surveillance audit should be product based with audit results being recorded on an approved Surveillance Audit Report.

The purpose of surveillance programmes is to:
- Verify that the approved quality system and associated product quality plans, continues to be implemented; and:
- To consider the implications of any changes to the system, initiated as a result of changes in the manufacturer’s operation; and
- To confirm continued compliance with ISO/IEC 80079-34.
- To evaluate any addition manufacturing, suppliers, sub-suppliers where critical requirements of ISO/IEC 80079-34 are being performed.

3.18 Re-Assessment
To verify overall continuing effectiveness of the manufacturer’s quality system in its entirety. In most cases it is unlikely that a period greater than three years for periodic re-assessment of the manufacturer’s quality system would satisfy this requirement. The re-assessment should provide for a review of past performance of the system over the period of the notification. The re-assessment program shall take into consideration the results of the above review and shall at least include a review of the quality system documents and a site audit (which may replace or extend a regular surveillance audit). It shall at least ensure
- the effective interaction between all elements of the system;
- the overall effectiveness of the system in its entirety in the light of changes in operations;
- demonstrated commitment to maintain the effectiveness of the system.

Upon the acceptance of the assessment, the Certification Body shall. Issue an IECEx Quality Assessment Report (QAR) and have this registered on the official IECEx Website www.iecex.com.

4 Principles of auditing
Auditing is characterised by its reliance on a number of principles. These make the audit an efficient and reliable tool in support of management policies and controls, providing information on which management can act to improve its performance. Adherence to these principles is a prerequisite for audit conclusions that are relevant and sufficient, such that auditors working independently from one another will reach similar conclusions in similar circumstances.
Three of these principles relate primarily to personal characteristics of the auditors themselves. These are:

i. **Ethical conduct - the foundation of professionalism**
   The role of the auditor is one of trust, integrity, confidentiality and discretion.

ii. **Fair presentation - the obligation to report truthfully and accurately**
   Audit findings, audit conclusions and audit reports reflect truthfully, accurately and completely the audit activities. Any unresolved or diverging opinions between the audit team and the auditee and any obstacles encountered are reported.

iii. **Due professional care - application of reasonable care in auditing**
   Auditors exercise a degree of care appropriate to the importance of the task they perform and to the confidence placed in them by audit clients and other interested parties. Having the necessary competence is an important prerequisite.

The remaining two principles of auditing relate primarily to the audit process. An audit is by definition independent and systematic and these characteristics are closely linked to the following two principles of auditing:

iv. **Independence - the basis for the impartiality and objectivity of the audit conclusion**
   Audits are objective and independent. Audit team members are free from bias and conflict of interest.

v. **Evidence - the rational basis for reaching audit conclusions**
   Audit evidence is verifiable. It is based on samples of the information available, since an audit is conducted during a finite period of time and with finite resources. However, the use of sampling is appropriate to the confidence placed in the audit conclusions.

5 Managing an audit programme

5.1 Introduction
An IECEx Certification Body having a need to conduct audits shall implement and manage an efficient and effective audit programme. The purpose of an audit programme is to assist the IECEx Certification Body in providing the resources necessary to facilitate the conduct of individual audits on a complete and timely basis.

The IECEx Certification Body’s management shall grant the authority for managing the audit programme.

Managing the audit programme includes:
   a) establishing the objectives and extent of the audit programme;
   b) establishing the responsibilities, resources and procedures;
   c) ensuring the implementation of the audit programme;
   d) monitoring and reviewing the audit programme;
   e) ensuring that appropriate audit programme records are maintained.
Figure 1 - illustrates the application of the Plan-Do-Check-Act cycle to the management of an audit programme

Note: The numbers in this and all subsequent figures refer to the relevant clauses of this document.

5.2 Audit programme objectives and extent

5.2.1 Objectives of the audit programme
The objective of the audit programme is to establish that the manufacturer is complying with ISO/IEC 80079-34, and with respect to the certificates listed or to be listed on the IECEx On-line system. Depending on their size, nature and complexity, organisations may meet the objectives in a single audit.
The objective can be based on consideration of:
   a) conformity of the product with the type described in the IECEx Test Report and with the requirements of the IEC Standard, which apply to it.
   b) management priorities;
   c) commercial intentions;
   d) management system requirements supporting conformity of the product;
   e) legal and contractual requirements;
   f) supplier evaluation;
   g) customer requirements;
   h) potential risks to the organisation.

5.2.2 Extent of the audit programme
An audit programme can vary in size, nature and complexity. The extent of the programme will be influenced by:
   a) scope, objective, duration and frequency of each audit to be conducted;
   b) size, nature and complexity of the organisation audited;
   c) the number, importance, complexity, similarity and locations of the activities to be audited;
   d) standards, legal and contractual requirements, policies, procedures and other audit criteria;
   e) accreditation and registration/certification;
   f) the results of previous audits or a previous audit programme review;
   g) language, cultural and social issues;
   h) concerns of interested parties;
   i) significant changes to any organisations, activities or functional areas.

5.2.2.1 Scope of initial assessment, surveillance and re-assessment

a) Initial assessments and re-assessments:
For initial assessments and re-assessments all requirements of ISO/IEC 80079-34 apply. These assessments may be conducted in whole or part at the manufacturer’s premises, as well as other manufacturing location(s)/production site(s) if applicable.

It is the CB (Certification Body) responsibility to determine what level of assessment is required for suppliers / sub-suppliers that are responsible for critical operations and requirements laid down in ISO/IEC 80079-34. Examples of critical operations include, but are not limited to:
   • Design and development changes
   • Assembly / sub-assembly of product
   • Routine Testing/Final testing of product
   • Labelling of product
   • Inspection and control of critical operations.
“Storage” and “handling” are activities included in the definitions of “Manufacturing Location” and “Production Site.” Storage and handling are critical operations only in cases where improper storage or handling of parts or assemblies may potentially impact the certification or safety of the final product. It is not the intent of this clause to require assessment of locations whose sole function is the storage of finished goods inventory after final assembly and packaging.

It is not the intent of this clause to require CB’s to audit suppliers or sub-suppliers. It is however the responsibility of the CB’s to evaluate if suppliers / sub-suppliers responsible for requirements of ISO/IEC 80079-34 have the necessary quality systems and quality plans in place to insure the requirements of ISO/IEC 80079-34 are being met. Examples of evaluations of suppliers / sub-suppliers can include:
- Review of a current ISO certificate with the proper scope
- Audits performed by the supplier
- Audits performed by another CB
- Audits by the CB
- Components covered by the IECQ Scheme (www.iecq.org)

b) Surveillance:
The surveillance visits at the manufacturer’s premises shall include:
- the system maintenance elements, which are internal audit, management review and preventive and corrective action;
- customer complaints;
- changes to the documented system;
- areas subject to change;
- selected elements of the certification/registration standard;
- other selected areas as appropriate.
- identification of any new manufacturing locations/production sites
- review of suppliers and/or sub-suppliers performing critical operations.
- re-assessment is defined in clause 3.18 of this document.

NOTE: Where a manufacturer has a certified quality system with an appropriate scope, certified by a recognised body (Type A or C manufacturer), then the scope of assessment/surveillance may be restricted to assessing the effectiveness of that quality system with respect to the certificates listed or to be listed upon the on-line notification system.

5.2.2.2 Audit programme for IECEx Certification Bodies
5.2.2.2.1 Initial and Re-assessments
IECEx Certification Bodies have essentially 3 types of customer, concerning initial or re-assessments:

TYPE (A) a manufacturer requiring an initial assessment or re-assessment prior to the issue of an IECEx QAR, where the manufacturer has a certified
quality system, for example, to ISO 9001:2008 requirements, with an appropriate scope certified by a recognised body. (The scope does not necessarily need to include protection concepts)

TYPE (B) a manufacturer requiring an initial assessment or re-assessment prior to the issue of a QAR, where the manufacturer does not have a certified quality system for example, to ISO 9001:2008 requirements, or a certified quality system with an inappropriate scope. (The scope does not necessarily need to include protection concepts)

TYPE (C) a manufacturer requiring initial assessment or periodic surveillance, where the manufacturer has also contracted the IECEx Certification Body in its capacity as an ISO 9000 Certification provider to integrate the two activities i.e. an IECEx Quality Assurance assessment and a certified quality system for example, to ISO 9001:2008 requirements. (The scope does not necessarily need to include protection concepts)

IECEx Certification Bodies are required to take into consideration the factors listed in clause 5.2.2 of this document. The ‘complexity’ factor given in 5.2.2 b) is one of the major issues for IECEx Certification Bodies in that products include equipment (electrical and non electrical) along with protective systems, components and devices required for contributing to the safe functioning of equipment and/or protective systems, all of which covers a wide range of technologies.

Therefore as a guide to the amount of resource (in auditor days) that should be spent conducting initial assessments, based upon the number of protection concepts, number of Ex personnel located onsite, and if there is a Certified Quality System in place the following table(s) are provided. These tables are only supplied as guidelines. It is the IECEx auditor’s responsibility to ensure that the appropriate time is spent to ensure that audits cover all requirements of ISO/IEC 80079-34 and any issued certificates on the online system.

WITH CERTIFIED QUALITY SYSTEM
(Manufacturing Types (A) and (C))

<table>
<thead>
<tr>
<th>No. OF PROTECTION CONCEPTS LISTED ON MANUFACTURER’S APPLICATION OR IECEx ON-LINE CERTIFICATE(S)</th>
<th>25 OR LESS Ex PERSONS ON SITE</th>
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WITHOUT CERTIFIED QUALITY SYSTEM
(Manufacturing Type (B))

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<tr>
<th>No. OF PROTECTION CONCEPTS LISTED ON THE MANUFACTURER’S APPLICATION OR IECEx CERTIFICATE(S)</th>
<th>25 OR LESS Ex PERSONS ON SITE</th>
<th>100 OR LESS Ex PERSONS ON SITE</th>
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<td>3 - 4</td>
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*The above times do not include the time spent by technical experts.

Protection concepts as referenced in the above tables are represented and identified by the different technologies, i.e., Ex d, Ex i, Ex n etc., as used for equipment certified under the IECEx Scheme.

These guidance times may be varied at the auditor’s discretion provided that all protection concepts are subject to adequate assessment / surveillance. For example the duration may be extended where there is not a common language, a large number of certificates are involved or reduced if processes are similar for several different products but with a minimum of one auditor day for an initial assessment.

5.2.2.2.2 Surveillance Assessments
Surveillance assessments shall be carried out as follows:-

a) For manufacturers without a certified ISO 9001: 2008 Quality System, surveillance audits should be conducted every 12 months.

b) Manufacturers with a Qualified ISO 9001: 2008 Quality System, surveillance audits should be carried out every 18 months.

Additional surveillance / assessments may be conducted at the discretion of the IECEx Certification Body based on the manufacturer’s / manufacturing location’s / production site’s performance and results of previous surveillance / audit results. It is recommended that the time frame for audit intervals begin when the initial assessment report is issued.

5.3 Audit programme responsibilities, resources and procedures
5.3.1 Responsibilities
Responsibility for managing an audit programme should be assigned to (an) individual(s) who has (have) a general understanding of audit principles, of
auditor competence and the application of audit tools and methods. They should also have technical and business understanding relevant to the activities to be audited.

Those assigned the responsibility for managing the audit programme should:

a) define, implement, maintain, and improve the audit programme;
b) identify and provide resources for the programme.

5.3.2 Resources
When identifying resources for the audit programme, consideration should be given to:

a) financial resources necessary to develop, implement, manage and improve audit activities;
b) audit tools and methods;
c) availability of auditors and technical experts;
d) processes to achieve and maintain auditor competence, and to improve auditor performance;
e) auditor competence appropriate to the particular audit programme objectives;
f) time, travel and other auditing needs.

5.3.3 Procedures
Audit programme procedures should be established and should address, for example:

a) planning and scheduling audits;
b) assuring the competence of auditors and audit team leaders;
c) selecting appropriate audit teams;
d) conducting audits;
e) performing audit follow-up.

5.4 Audit programme implementation
The implementation of an audit programme should include:

a) documenting the audit programme and communicating it to relevant parties;
b) co-ordinating and scheduling audits and other audit programme activities;
c) establishing and maintaining a process for the initial evaluation of auditors and the on-going evaluation of training needs and continuing professional development of auditors;
d) ensuring the appointment of audit teams;
e) providing required resources to the audit teams;
f) ensuring the conduct of audits according to the audit programme;
g) ensuring the control of records of the audit activities;
h) ensuring review and approval of audit reports, and ensuring their distribution to the audit client and other specified parties;
i) ensuring audit follow-up, if applicable.
5.5 Audit programme records
Records of all IECEx audits shall be maintained by the IECEx Certification Body to demonstrate the operation of the audit programme and shall include:

a) results of audit programme review;
b) audit records, such as:
   - audit plans;
   - audit reports;
   - nonconformity reports; and
   - corrective and preventive action reports;
c) audit personnel records, covering subjects such as performance evaluation, audit team selection and training.

5.6 Audit programme monitoring and reviewing
The effectiveness of an IECEx Certification Body’s auditing programme shall be monitored by the ExCB itself and at appropriate intervals reviewed and included as part of their internal quality system review, to assess whether its objectives and those of the IECEx Scheme have been met. This audit programme review should be carried out to assess the audit programme effectiveness and identify opportunities for improvement.

Monitoring should be carried out by using performance indicators that measure, for example:

- the ability of the audit teams to meet audit objectives;
- conformity with audit programmes and schedules;
- feedback from audit clients, auditees and auditors;
- time to close audit programme corrective actions.

This audit programme review should consider for example:

- results and trends from monitoring;
- conformity with procedures;
- evolving needs and expectations of interested parties;
- audit records;
- alternative or new auditing practices.

Results of the audit programme review can lead to corrective actions and improvement of the IECEx Certification Body’s audit programme.

6 Audit Activities
6.1 Introduction
This clause contains guidance on managing and conducting audits, including the selection of audit team members.
After completion of an audit, audit follow-up actions can take place (see clause 6.8). Figure 2 provides a flowchart of the audit process as described in this clause.

![Flowchart of the audit process](image)

**Figure 2 – Illustrates where QARs fit into an example Manufacturer’s process**

In this example, 3 QARs are issued – the Manufacturer (A), a Production Site (B), and a Manufacturing Location (C). It is also possible that supplier audits will be performed, per ISO/IEC 80079-34.

Graphics courtesy of Microsoft Visio
6.2Initiating the audit
6.2.1Audit objectives, scope and criteria
Within the overall objectives of an audit programme, an individual audit should be based on defined objectives, scope and criteria.

The objective for the audit programme is to establish that the manufacturer is complying with ISO/IEC 80079-34 with respect to all the certificates listed on the online notification system, and kept up-to-date by the IECEx Certification Body planning and conducting the audit programme. An individual audit that is part of a surveillance programme may address only part of a quality management system.

In preparing for an audit, the following shall be communicated to the auditee and recorded on the job file:

• The audit scope describes the extent and boundaries of the audit in terms of factors such as physical locations, organisational units, activities and processes to be audited and, where relevant, the time period covered by the audit.
• The audit criteria, i.e. ISO/IEC 80079-34
• Dates of the audit, as agreed with the auditee
• Membership of the audit Team

6.2.2Feasibility of the audit
Those responsible for managing the audit programme should determine the feasibility of the audit, taking into consideration such factors as:
a) sufficient and appropriate information for planning the audit;
b) adequate co-operation from the auditee;
c) availability of time and adequate resources.

Upon receipt of an application from a manufacturer for an IECEx Certificate of Conformity, the IECEx Certification Body should:
1. ensure that the documentation as required by ISO/IEC 80079-34 as appropriate and the relevant IEC Standard is provided.
2. ascertain that sufficient and appropriate information regarding the manufacturer is available e.g. size, location, status of quality system.

An IECEx Certification Body is NOT permitted to issue an IECEx Quality Assessment Report (QAR) until a document review and site assessment have been satisfactorily completed.

Where products to be covered by an IECEx Certificate of Conformity are to be manufactured in different locations or countries, the IECEx Certification Body to whom the application for an IECEx Certificate of Conformity has been lodged
(known as ExCB ‘A’) may engage other IECEx Certification Bodies to perform sites visits of the various manufacturing locations under the direction and control of ExCB ‘A’.

An IECEx Certification Body is permitted to amend the scope of an existing QAR (without a site assessment visit) where the technology/processes are declared similar to those already covered.

6.2.3 Establishing the audit team
When the audit has been declared feasible, an audit team should be established and an audit team leader is appointed taking into account the competence needed to achieve the objectives of the audit. When there is only one auditor, the auditor should perform all applicable duties of an audit team leader.

Those responsible for managing the audit programme and/or the audit team leader, in consultation with the audit client and, if necessary, the auditee, should identify the resources necessary.

When deciding the size and composition of the audit team, consideration should be given to the following:

a) audit objectives, scope, criteria, location(s) and estimated duration;
b) the overall competence of the audit team needed to achieve the objectives of the audit;
c) requirements from accreditation/certification bodies, as applicable;
d) the language of the audit and understanding of the auditee’s social and cultural characteristics either through their own skills or through the support of a technical expert;
e) the need to assure the independence of the audit team from the activities to be audited and to avoid conflict of interest;
f) the ability of the audit team members to interact effectively with the auditee and to work together.

The process of assuring the competence of the audit team should comprise the following steps:

1) identifying the knowledge and skills needed to achieve the objectives of the audit;
2) defining the criteria by which knowledge and skills are to be evaluated;
3) selecting the audit team such that all of the knowledge and skills needed to conduct the audit and to achieve the audit objectives are present in the audit team. If not fully covered by the auditors in the team, the overall competence may be satisfied by including technical experts in the team.
Technical experts should operate under the direction of an auditor. The IECEx Certification Body should evaluate and record the required competencies of technical experts.

Both the audit client and auditee have a right to request the replacement of particular team members on reasonable grounds which should be communicated to those responsible for managing the audit programme. Any decisions to replace team members should be taken by those responsible for managing the audit programme. Examples of reasonable grounds can be conflict of interest situations (such as an audit team member having been a former employee of the auditee or having provided consultancy services) or previous unethical behaviour.

6.2.4 Initial contact with the auditee
The initial contact with the auditee can be informal or formal. Depending on the audit situation, those responsible for managing the audit programme or audit team leader should consider the following, as appropriate:
   a) contacting the auditee to establish communication channels;
   b) providing information on proposed timing and audit team composition;
   c) requesting documents, including records, if needed, and
   d) making arrangements for the audit.

Any need for accompanying persons such as observers or guides for the audit team should be mutually agreed.

6.3 Document Review
Relevant management system documents, including records, from the auditee, including any previous audit reports, should be reviewed to determine the conformity of the system components or processes, as documented, with audit criteria. The review, normally by the audit team leader or by one or more auditors assigned by the audit team leader, should take into account the size, nature and complexity of the organisation, and the objectives and scope of the audit. A preliminary on-site visit is an option to get a good overview of available information.

If the auditee’s management system documentation is found to be inadequate, such that it is not commensurate with the audit scope or criteria, the audit client, those responsible for managing the audit programme and the auditee should be informed. Further resources should not be expended on the audit until such concerns are resolved to the satisfaction of those responsible for managing the audit programme in consultation with the audit client, the audit team leader and, if appropriate, the auditee.
6.4 Preparing for the on-site audit activities
6.4.1 Planning the on-site audit activities

NOTE The following guidance may not be fully applicable to unexpected visits.

The audit team leader should prepare a plan for the on-site audit activities. This plan should provide necessary information to the audit team, auditee and audit client. It also facilitates scheduling and co-ordination of the audit activities.

The level of detail provided in the audit plan, should be adapted to suit the scope and complexity of the audit. The details can for example differ between initial and subsequent surveillance visits.

The audit plan includes the audit objectives and scope; which shall include a product audit and should include:

a) the audit criteria and any reference documents;
b) the dates and places where the on-site audit activities are to be conducted;
c) the identification of the organisational and functional units and processes to be audited;
d) the expected time and duration for audit on-site activities, including meetings with the auditee’s management and audit team meetings.

The audit plan can also include, as appropriate:

e) the identification of the sites, activities, and management system processes that are essential to meeting audit objectives in order to allocate appropriate resources to critical areas of the audit;
f) the identification of the auditee’s key representative participating in the audit;
g) the working and reporting language(s) of the audit where this is different from the native language of the auditor(s) and/or the auditee;
h) the identification of roles and responsibilities of the audit team members and any accompanying persons;
i) the audit report topics (including any methods of nonconformity gradings), format and structure, expected date of issue and distribution;
j) logistic arrangements (travel, on-site etc.)
k) matters related to confidentiality;
l) any arrangements for audit follow-up actions.

The plan is to be presented to the manufacturer prior to conducting all certification audits but may not be necessary for all surveillance audits.

Any objections by the auditee should be resolved among the audit team leader, the auditee, the audit client and ExCB management before continuing the audit.

The audit plan should be sufficiently flexible to permit changes, such as any changes in emphasis, which can become necessary as the on-site audit activities
progress. Any revised audit plan should be agreed among the parties concerned before continuing the audit.

6.4.2 Audit team work assignments
When an audit is conducted by an audit team comprising more than one auditor, the audit team leader, in consultation with the audit team, should assign to each team member responsibility for auditing specific management system processes, functions, sites, areas or activities. Such assignments should take into account the need for auditor independence, competence and efficient use of resources as well as different roles and responsibilities of auditors, auditors-in-training, and technical experts. Changes to the work assignments can be made to ensure the achievement of the audit objectives.

The audit team members should review the relevant information related to their audit assignments and prepare any work documents necessary for those assignments.

6.4.3 Work documents
Work documents used by the audit team for the purpose of reference and/or recording the proceedings of the audit can include:

a) audit procedures, checklists and audit sampling plans;

b) forms for recording information, supporting evidence, records of audit findings and meetings.

The use of work documents, such as checklists and forms, should not restrict the extent of audit activities.

Work documents, and any records resulting from their use, should be retained, at least until audit completion. The requirements for the retention of documents, including records, after audit completion, are described in clause 6.6.3. Those records involving confidential or proprietary information should be suitably safeguarded at all times by the audit team members.

6.5 On-site audit activities
6.5.1 Opening meeting
An opening meeting should be held to confirm the audit plan, clarify how the audit activities will be undertaken, and to establish communications. The opening meeting should be held with the auditee's management or, where appropriate, those responsible for the functions or processes to be audited. The opening meeting should also include opportunity to the auditee to ask any questions.

The meeting should be formal and records of the attendance should be kept. The meeting should be chaired by the audit team leader and the following items considered, as appropriate:
6.5.2 Roles and responsibilities of guides
Where guides are assigned, they should assist the audit team and act on the request of the audit team leader. Their duties can include ensuring that rules concerning site safety and security procedures are known and respected by the auditors, and they can also witness the audit on behalf of the auditee. Guides should not influence or interfere with the conduct of the audit except where, with the agreement of the auditor, the guide can provide clarification or assist in establishing correct information.

6.5.3 Collecting and verifying information
Figure 3 provides an overview of the process steps from collecting information to reaching audit conclusions.
Information relevant to the audit objectives, scope, and criteria, including information relating to interfaces between functions, activities, and processes should be collected during the audit. It should be verified by the auditor(s) and can then be considered to be audit evidence. Audit evidence should be identified as such and recorded.
NOTE The audit evidence will inevitably be only samples of the information available, since an audit is conducted during a finite period of time and with limited resources. There is thus an element of uncertainty inherent in all audits, and those acting upon the audit conclusions should be aware of this uncertainty.

The sources of information chosen can vary according to the scope and complexity of the audit and can include:

a) interviews;
b) observations of activities and the surrounding work environment and conditions;
c) documents, including, for example, policy, objectives, plans, procedures, instructions, product certificates and notifications, specifications, drawings, contracts orders;
d) records, such as inspection records, minutes of meetings, reports or logbooks on customer complaints and other relevant communication from external parties, audit reports, monitoring programmes and results of measurements;
e) data summaries, analyses, metrics and performance indicators;
f) records of the basis of relevant auditee's sampling programmes and the procedures for ensuring effective quality control of sampling and measurement processes;
g) reports from other sources, for example, customer feedback, external reports and vendor supplier ratings;
h) computerised data bases and web sites.

**Practical Help – Interviews**

Interviews are one of the important means of collecting information and should be carried out in a manner adapted to the situation and person interviewed. However, the auditor should consider the following:

a) interviews should be held with persons from different levels and functions, and especially with persons performing activities or tasks within the scope of the audit;
b) whenever possible, the interview should be conducted during normal working hours and at the normal workplace of the interviewed person;
c) every attempt should be made to put the interviewed person at ease prior to the interview;
d) the reason for the interview and any note taking should be explained;
e) interviews can be initiated by asking the persons to describe their work;
f) the results from the interview should be summarised and reviewed with the interviewed person;
g) questions that bias the answers (leading questions) should be avoided;
h) the interviewed persons should be thanked for their participation and cooperation.
6.5.4 Audit findings
Collected audit evidence should be evaluated against the audit criteria to generate the audit findings. An audit finding can indicate either conformity or nonconformity with audit criteria. If so decided, audit findings can be graded in accordance with the audit plan.

An audit team should meet as needed to review the audit findings at appropriate stages during the audit.

Conformities should be summarised to at least indicate locations, functions, processes, or requirements that were audited.

Individual audit findings of conformity should also be recorded and supported with audit evidence.

Nonconformities should be recorded and supported by audit evidence. Nonconformities should be reviewed with an appropriate auditee representative to obtain acknowledgement of the audit evidence. The acknowledgement indicates that the audit evidence is accurate, and that the nonconformity is understood. Every attempt should be made to resolve any divergence of opinion concerning the audit evidence and/or findings, and unresolved points should be recorded.

The audit team leader shall determine whether it is possible for the auditee to correct any nonconformities raised, prior to the completion of the audit.

6.5.5 Communication during the audit
Dependent upon the scope and complexity of the audit, it can be necessary to make formal arrangements for communication during the audit.

An audit team should confer at least daily in order to exchange information, assess audit progress, and reassign work between auditors as needed.

During the audit, the audit team leader should periodically communicate the status of the audit and any concerns to the auditee and audit client, as appropriate. Any evidence collected in the audit that suggests a significant risk exposure should be reported immediately to the auditee and, as appropriate, to the audit client.

Where the available audit evidence indicates that the audit objectives are unattainable, the audit team leader should report the reasons to the audit client and the auditee to determine the appropriate action. Such action can include
reconfirmation of the audit plan, termination of the audit or a change in the audit objectives.

Any concern about an issue outside the audit scope should be noted and reported to the audit team leader, for possible communication to the audit client and auditee. Any need for changes in the audit scope which may become apparent as on-site auditing activities progress should be reviewed with and approved by the audit client and, as appropriate, the auditee.

6.5.6 Preparation for the closing meeting
The audit team should confer prior to the closing meeting in order to:

a) review the audit findings and any other appropriate information collected during the audit;
b) prepare a list of audit findings, if appropriate;
c) reach consensus on the audit conclusions;
d) agree on roles and tasks for the closing meeting;
e) prepare recommendations, if specified by the audit objectives;
f) discuss subsequent audit follow-up, if appropriate.

In many instances a simplified approach can be taken for the audit team review, depending on the audit objectives and scope and the audit team size.

6.5.7 Closing meeting
A closing meeting should be held to present audit findings and conclusions in such a manner as to ensure that they are understood and acknowledged by the auditee, and to agree, if appropriate, on the time period for the auditee to present any corrective action plan.

The meeting should be formal and records of attendance should be kept. The meeting chaired by the audit team leader should be held with the auditee’s management and those responsible for the functions audited.

Any unresolved diverging opinions relating to audit findings and/or conclusions between the audit team and the auditee should be discussed and if possible resolved. If not resolved, both opinions should be recorded.

6.6 Reporting on the audit
6.6.1 Audit report preparation and content
The audit team leader should be responsible for the preparation and contents of the audit report.

The audit report should provide a complete, accurate, concise and clear record of the audit and should contain audit conclusions on issues such as the following, if within the audit objectives and scope:
extent of conformance of the quality system to the requirements of ISO/IEC 80079-34;

the effective implementation and maintenance of the quality system, relevant to the requirements of ISO/IEC 80079-34;

the ability of management review process to ensure the continuing suitability, adequacy, and effectiveness of the quality system, relevant to the requirements of ISO/IEC 80079-34;

The audit report should also include, or make reference to the following:

a) the identification of the organisational and function units or processes audited;
b) the identification of the manufacturer and audit client;
c) the identification of audit team members;
d) the date(s) and place(s) the on-site audit activities were conducted;
e) the audit criteria, and, if applicable, a list of reference documents, against which the audit was conducted, e.g. ISO/IEC 80079-34;
f) the audit findings.

The audit report can also include or reference, as appropriate:

g) the agreed audit objectives, scope and plan;
h) the time period covered by the audit;
i) the identification of the auditee's key representatives participating in the audit;
j) a summary of the audit process including any obstacles encountered;
k) a statement of the confidential nature of the contents;
l) a distribution list for the audit report;
m) confirmation that the audit objectives have been accomplished within the audit scope in accordance with the audit plan;
n) any agreed follow-up action plans;
o) any unresolved diverging opinions between the audit team and the auditee;
p) areas not covered, although within the scope.

6.6.2 Report approval and distribution

Every attempt shall be made for the audit report to be submitted to the audit client within one month from the date of the audit. The audit team leader should be responsible for monitoring this. If this is not possible, the audit team leader shall inform the audit client of the reasons for the delay and a revised issue date then agreed.

The audit report shall be dated, and reviewed by the relevant ExCB Manager and approved by a nominated representative within the ExCB.

The audit report should then be distributed as follows:

• Original to the audit client;
• A copy to be retained by the IECEx Certification Body

When the audit report has been issued to the audit client, the ExCB conducting the audit shall register the QAR on the official IECEx On-Line System, www.iecex.com

Note: Refer to Operational Document OD 011-2 for guidance regarding On-Line issue of QARs.

The confidentiality of the audit report should be respected and appropriately safeguarded by the audit team members and all report recipients.

6.6.3 Retention of documents
The audit report should be retained for a period not less than 10 years by the IECEx Certification Body, or longer at their own discretion.

Unless required to do so by law, the IECEx Certification Body, audit team and those responsible for managing the audit programme should not disclose the contents of documents, any other information obtained during the audit, or the audit report, to any other party without the explicit approval of the audit client and, where appropriate, the approval of the auditee. If disclosure of the contents of any audit document is required, the audit client and auditee should be informed as soon as possible.

6.7 Audit completion
The audit is completed when all activities in the audit plan have been finalised and the approved audit report has been distributed to the manufacturer with a copy retained by the IECEx Certification Body.

6.8 Audit Follow-up
6.8.1 Review
The IECEx Certification Body should review the lead auditors audit report, any non conformities raised and the manufacturer’s responses, and then the IECEx Certification Body should approve the report.

The IECEx Certification Body should have a unified method of addressing the actions required following an audit.

The following rating system is provided for guidance.

<table>
<thead>
<tr>
<th>RATING</th>
<th>DEFINITION</th>
<th>ACTION FOLLOWING AN INITIAL ASSESSMENT OR RE-ASSESSMENT</th>
<th>ACTION FOLLOWING A SURVEILLANCE VISIT</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Where a quality system fully meets the</td>
<td>Issue new or updated</td>
<td>Issue updated IECEx</td>
</tr>
<tr>
<td>RATING</td>
<td>DEFINITION</td>
<td>ACTION FOLLOWING AN INITIAL ASSESSMENT OR RE-ASSESSMENT</td>
<td>ACTION FOLLOWING A SURVEILLANCE VISIT</td>
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<tr>
<td></td>
<td>requirements or where there are only very few minor nonconformities. Also where compliance of the product is observed during a product audit.</td>
<td>IECEx QAR</td>
<td>QAR</td>
</tr>
<tr>
<td>B</td>
<td>Where the quality system has a series of minor nonconformities Also where compliance of the product is observed during a product audit.</td>
<td>Issue new or updated IECEx QAR upon receipt of satisfactory documentary evidence supporting effective corrective action which is then subject to verification at the next surveillance visit</td>
<td>Issue updated IECEx QAR upon receipt of an acceptable corrective action plan, which is subject to verification at the next visit.</td>
</tr>
<tr>
<td>C</td>
<td>Where the quality system has major nonconformities and, or there is a non compliant product observed during the product audit.</td>
<td>Issue new or updated IECEx QAR only after a satisfactory follow-up visit has verified that the corrective actions have been effectively documented and implemented.</td>
<td>Issue updated IECEx QAR only after a satisfactory follow-up visit has verified that the corrective actions have been effectively documented and implemented. Should the manufacturer fail to take timely and effective corrective action, then the IECEx Certification Body reserves the right to suspend or cancel the IECEx Certificate of Conformity.</td>
</tr>
<tr>
<td>D</td>
<td>Where the quality system has many major nonconformities which may include non compliant product observed during the product audit</td>
<td>Issue new or updated IECEx QAR only after a further complete assessment of the quality system has been satisfactorily completed.</td>
<td>Suspend the IECEx Certificate of Conformity pending a further complete assessment to re-establish the effectiveness of the quality system. This is to be followed by surveillance visits at a frequency which maintains confidence in the effectiveness of the quality system.</td>
</tr>
<tr>
<td>E</td>
<td>Where there is no quality system or a system that has serious deficiencies rendering it ineffective</td>
<td>Close the application, no IECEx QAR to be issued or re-issued</td>
<td>Cancel the IECEx Certificate of Conformity and inform other IECEx Certification Bodies</td>
</tr>
</tbody>
</table>
6.8.2 Issue of QAR
A Quality Assessment Report (QAR) shall be valid for a period not exceeding 3 years.

IECEx Certification Bodies shall register the QAR on the IECEx Website in accordance with Operational Document OD 011 Part 2. This On-line registration provides the following summary information, which is publicly available.

- QAR reference number
- ExCB conducting the audit
- Manufacturer and audit location(s)
- IECEx Certificates of Conformity linked to the QAR
- Comments of the ExCB if any

In accordance with OD 011-2, the online QAR summary should include the Manufacturer’s name in the “Manufacturer” field, and the location(s) audited in the “Site(s) Audited” field. The ExCB shall consider whether a separate QAR is needed for each site audited.

The “Comments” field of the QAR summary should be used to explain the relationships between sites audited, in cases when different critical operations take place at different locations. One exception to this is for Case 2 or Case 3 Trade Agents per OD 203, when the relationship between the trade agent and OEM is to be kept confidential. In these cases the details are to appear in the QAR but not in the online QAR summary.

In addition, IECEx Certification Bodies shall inform the IECEx Secretary of any IECEx Certificates of Conformity that are to be suspended or cancelled. This will ensure that other IECEx Certification Bodies are informed of all Certificates that are issued, suspended or cancelled.