QM003
Quality Requirements for Suppliers
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Plan & Manage

Performance Evaluation
- Competence, Training and Awareness

Obsolescence Management
- Business Continuity / Disaster Management

Supplier Approval

- Handling Sensitive and Proprietary data
- Counterfeit Product Prevention
- Special Process Approval

Design & Development

- Review of the Requirements for Products and Services
- Design and Development Provision
- Configuration Management
- Deliverable Quality Plan

Make

- Configuration Management
- Process Control and Verification
- Control of Externally Provided Processes, Products and Services
- Control of Equipment, Tools and Software
- Validation of Special Processes
- Production Process Verification
- Application for Production (Deviation) Permit or Concession
- Identification and Traceability
- Traceability and Counterfeit Avoidance
- Workmanship Acceptance Criteria for Surface Engineering
- Foreign Object Debris (FOD) Materials Database

Deliver

- Release of Product and Services
- Shelf Life
- Packaging
- Chemicals and Hazardous Substances
- Munitions
- Legislative Requirements
- Counterfeit Product Prevention

Post Delivery

- Control of Non Conformance
- Record Retention / Destruction Requirement

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1. Foreword

Lockheed Martin’s business depends on a reliable, global network of skilled suppliers that provide the materials, parts and services to make our products and deliver them to our customers, conforming to LMUK-A (Lockheed Martin UK Ampthill) requirements, mission-ready and on time. Goods and services provided by our suppliers have a key impact on the quality of the products, solutions and services we offer our customers.

To maintain a high level of quality we are determined to establish and maintain close and long-lasting relationships with our suppliers working in a zero-defect culture from our supply chain.

QM003 may be distributed to sub-contractors, suppliers and the sub-tier supply chain.
LMUK-A Terms and Conditions of trade shall apply to all contracts unless otherwise agreed.

1.1 Traceability and Counterfeit Avoidance

Traceability and counterfeit avoidance are of paramount importance to Lockheed Martin to avoid safety and operational effectiveness of our products being compromised by defective items. Lockheed Martin rely on suppliers’ stringent controls to ensure traceability is maintained as per requirements and counterfeit risks mitigated. Refer to sections 9 & 10 of this document for detailed requirements.

1.2 Sustainable Supply Chain

LMUK-A define Sustainable Supply Chain Management (SSCM) as “management of our supply base to drive affordability and innovation through social responsibility and environmental stewardship.” The objective of SSCM is to ensure alignment of our supply base’s social, ethical, environmental, safety and health responsibilities with Lockheed Martin’s sustainability commitments. The Lockheed Martin UK Ampthill Ltd Quality Policy can be obtained upon request.

1.3 Conflict Minerals

Conflict minerals are minerals mined in conditions of armed conflict and human rights abuses, and which are sold or traded by armed groups. Suppliers shall be aware of the OECD Due Diligence Guidance. Further information can be found at the Lockheed Martin Conflict Minerals webpage.

1.4 Zero Concession Policy.

LMUK-A expects quality to be built into the Design and Production of our product and this is also the expectation of the supply chain. Conformity first is key to the delivery of world class product to our customers and this culture and mindset needs to be within the supply chain delivering to LMUK-A. As a company LMUK-A relies on the supply chain to deliver product conforming to requirements, free from defects with zero concessions or deviations.

Requests for concession applications will only be accepted under exceptional circumstances. Concession submissions will impact the supplier performance rating.
2. Scope

LMUK-A are required by AS9100 to apply appropriate controls to their direct and sub tier external providers to ensure that requirements are met.

The aim of this document is to formally communicate the Lockheed Martin UK Ltd (LMUK-A) quality requirements to the supply chain.

The supplier must refer to the Contract between LMUK-A and the supplier to identify the applicable version of QM003.

Suppliers shall plan, implement, and control the processes needed to meet the requirements for the provision of products and services to LMUK-A in accordance with this document and the suppliers’ QMS.

2.1 Definitions and Terms

In this Quality Requirements for Suppliers QM003 document, the terms "shall" and "must" mean that the described action is mandatory; "should" means that the described action is expected with some flexibility allowed in the method of compliance; and “may” means that the described action is permissible or discretionary.

The term “supplier” means vendor, supplier of goods and services, sub-contractor and distributor. Questions concerning this manual should be directed to your respective LMUK-A Supply Chain representative or Supplier Quality Engineer (SQE).

The term “Contract” refers to the legal agreement between LMUK-A and the Supplier for the provision of goods and/or services in which the requirement to comply with a version of QM003 has been incorporated.

3. Order of Precedence

Any inconsistencies in this document shall be resolved in accordance with the following descending order of precedence: (1) the Contract including any special terms and conditions; (2) the drawing, design data and any approved concession deviation (3) QM003.

4. Quality Conditions (Q Code)

Additional Quality Conditions required by NATO & Military contracts above those detailed within QM003 basic requirements and ISO9001 are defined by quality condition (q code) QM003-A & QM003-B which will be stated on LM’s purchase order and/or Sub-Contract.

These will only apply when indicated by the delivery quality condition (Q Code) on the purchase order or other documentation associated with the Contract. See Diagram 1 – Additional Flow-down requirement selection flowchart and Appendix 1 – Additional AQAP Guidance notes.

<table>
<thead>
<tr>
<th>Q Code</th>
<th>Description</th>
<th>Flow-Down Requirement in Addition to ISO9001 &amp; QM003</th>
</tr>
</thead>
<tbody>
<tr>
<td>QM003-A</td>
<td>Quality Requirements for NATO Contracts</td>
<td>AQAP 2110 / 2105 Applies</td>
</tr>
<tr>
<td>QM003-B</td>
<td>Quality Requirements for Non-Military Contracts</td>
<td>ISO 10005 / 10007 Applies</td>
</tr>
<tr>
<td>QM003-C</td>
<td>Quality Requirements for Business Supplies</td>
<td>No additional requirement</td>
</tr>
</tbody>
</table>
Diagram 1 - Additional Flow-down requirement selection flowchart

- **QM003-A** Quality Requirements for NATO Contracts
- **QM003-B** Quality Requirements for Non-Military Contracts
- **QM003-C** Quality Requirements for Business Supplies

START

Para. 6.1 Supplier has ISO9001 certification?

Delivery Conditions Apply?

Para. 6.1 Quality Requirement exceptions apply?

Para. 6.1.4 Any exceptional circumstances?

YES

NO

QM003-A

QM003-B

QM003-C

Supplier has AS9100 certification?

YES

NO

AQAP 2310

AQAP 2110

Supplier has AS9100 certification?

NO

YES

ISO 10007

Para 8.2 Deliverable Quality Plan required?

NO

YES

Development Software?

YES

NO

AQAP 23110

AQAP 2105

ISO 10005

QM003 General Requirements

NO additional quality requirements

Use alternative supplier
5. Sensitive and LMUK-A Proprietary Data

5.1 Handling Sensitive and Proprietary data

Sub-tier suppliers and sub-contractors used by the supplier that have access to any sensitive or LMUK-A proprietary data must be authorised with a Non-Disclosure Agreement (NDA) in place with LMUK-A.

LMUK-A proprietary and customer technical data must only be shared with 3rd party suppliers who have:

- Been approved by LMUK-A and the owner of the technical data
- Confirmed in writing (e.g., hardcopy letter, email with return address header) that they are authorized to receive such data and they understand the implications of and requirements for handling sensitive and proprietary technical data
- Only authorised personnel shall have access to restricted data and the data shall be controlled in such a way as to prevent unauthorized transmission or access

Suppliers that require Restricted and Official Sensitive Classification data shall have a procedure in place for the control, handling and monitoring of such data. Any loss or potential compromise of any classified material must be reported to LMUK-A without delay.

Lockheed Martin Proprietary Information (LMPI) shall be handled in accordance with the LMUK-A terms and conditions.

Where a supplier is identified on a Technical Assistance Agreement (TAA) or Manufacturing Licence Agreement (MLA), the organisation must complete a Non-Disclosure Agreement (NDA) when requested by LMUK-A and shall continue to maintain access controls in accordance with the NDA and any Technology Control Plan (TCP) that LMUK-A and the organisation enter into.

LMUK-A reserve the right to issue an NDA where LMUK-A deem sensitive information will be shared with the supplier.

5.2 Record Retention / Destruction Requirement

Suppliers shall retain records relating to processing, testing, calibration, manufacture, supply, traceability and certification for a minimum of 7 years from end of Contract date unless otherwise stated by Contract or statutory requirements (i.e. 10 years for EU declaration of Conformity & CE Marking as defined in Journal 2016/C 272/01).

All OFFICIAL + LMPI documents must be returned to Lockheed Martin UK Ampthill either:

a. When they are no-longer required as part of the sub-contract; or
b. At the end of the sub-contract.

Requests to destroy programme-related OFFICIAL information locally on the sub-contractor’s site must be passed to the Lockheed Martin UK, Ampthill Programme/Project Manager and Lockheed Martin UK UK Ampthill, Security Office for authorisation in accordance with LMUK-A Contract conditions

At a minimum, all OFFICIAL hard copy information must be destroyed using a cross-cut shredder which makes the reconstitution of the material highly unlikely. Unwanted OFFICIAL information/material that cannot be destroyed in such a way shall be returned to the originating Authority.
6. Supplier Approval

6.1 Supplier Approval Minimum Requirements

The minimum quality requirement for suppliers of goods supplied, work performed and services provided to LMUK-A **shall** be Quality Management System (QMS) certification to ISO9001 by a **UKAS** (or national equivalent) accredited certification body.

Suppliers that provide goods and services that are used in projects for aviation and space applications **should** be certified to AS9100 for QM003-A & QM003-B Contracts.

Distributors **shall** be certified to AS9102 and Calibration Laboratories **shall** be certified to ISO 17025 or equivalent for QM003-A & QM003-B Contracts.

6.1.1 Exceptions

LMUK-A may approve exceptions to the above requirements if any of the following criteria are met:

- The supplier is mandated by our customer
- The supplier is the manufacturer of a single sourced product mandated by our customer
- The supplier is the only distributor of a product mandated by our customer
- The supplier provides goods or services that have no direct or indirect effect on the goods and services we provide our customer

6.1.2 Supporting Documentation

Documents required to complete the supplier approval process are:

- Form **F0189 Supplier Quality Assessment** and / or F0183 Supplier Assessment Questionnaire
- QMS certification
- Confidentiality or non-disclosure agreement (NDA) if applicable
- Inclusion on a Technical Assistance Agreement (TAA) if applicable
- Deliverable Quality Plan (See section 8.2)

(Form F0189 is available on the Lockheed Martin UK website.)

6.1.3 Security Assessment

Suppliers working at OFFICIAL-SENSITIVE/LEGACY RESTRICTED, will be required to undergo a Security Assurance visit and Risk Assessment by LM Security personnel prior to the exchange of any sensitive information.

Following the initial Security Assurance visit, the supplier will be required to complete an Annual Security Questionnaire which will be evaluated to ensure that there has been no change to the Risk Assessment.

Suppliers may be subject to subsequent, follow up Security Assurance visits where required.
6.1.4 Exceptional Circumstances

Where the supplier approval criteria or listed exceptions cannot be met, depending on the product, its application, value and criticality, special authorisation may be granted where evidence of compliance can be provided.

This may include LMUK-A audit to a set of alternative quality requirements.

6.1.5 Scope of Approval

- Suppliers approved for use will be allocated to the LMUK-A Supplier database stating the scope detail on their approval
- Suppliers shall not conduct work for LMUK-A outside their scope of approval unless authorised by LM Quality department through audit i.e. special process audits or LM specific approvals

6.2 Maintenance of Approved Status

6.2.1 Site Visits and Supplier Audits

- Where appropriate, suppliers shall be subject to on-site audit and / or site visit by Lockheed Martin
- In some instances, LMUK-A will be unable to raise a contract until completion of successful supplier audit
- Scheduled verification audits, site visits and business to business meetings shall be supported when required
- If specified on the Contract, Verification at Source acceptance inspection and witness testing by Buyer Quality Assurance representative shall be required prior to shipment of product from your facility. The Seller shall provide a notice period of 7 days prior to which components will be available for source inspection. To notify and arrange email sqa_ampthill.fc-amphill@lmco.com and the nominated supply chain representative identified on the Contract

6.2.2 Approval updates – Supplier Responsibilities

It is a requirement of the conditions of supply into LMUK-A that the contractor / supplier fully understands and adheres to the following. It is the supplier’s responsibility to ensure:

- LMUK-A shall be provided up-to-date copies of Quality Management System certification including scope of certification
- LMUK-A shall be informed by the approved supplier when approval bodies are changed, and certificates are re-issued or revoked
- LMUK-A shall be informed by the approved supplier when certificates scopes are planned to be amended which would affect work currently undertaken or scheduled for future delivery. This would also include any change of address
- LMUK-A shall be informed if due to any circumstance suppliers Special Process skill base alters i.e. coded welder’s certification lapses – in this instance the LMUK-A Special Process Auditor must be informed

To notify LMUK-A of any updates email sqa_ampthill.fc-amphill@lmco.com or the nominated LMUK-A supply chain representative identified on the Contract.
6.2.3 Right of Access

Suppliers and their sub-suppliers shall provide to LMUK-A, their customer, regulatory authorities and the Government Quality Assurance Representative (GQAR):

- The right of access to facilities where parts of the contracted activities are being performed including sub-suppliers’ premises
- Information pertaining to the fulfilment of requirements in the Contract
- Unrestricted opportunity to evaluate supplier compliance with this document
- Unrestricted opportunity to conduct verification of product conformity to Contract requirements
- Assistance for evaluation, verification, validation, testing, inspection or release of the product to verify that Contract requirements have been accomplished at the supplier’s or sub-supplier’s premises
- Working area and facilities
- The necessary equipment available for reasonable use for performing verification
- Supplier and/or sub-supplier’s personnel for operation of verification equipment as required
- Access to information and communication facilities
- The necessary supplier documentation, to confirm product conformance to specification
- Copies of necessary documents, including those on electronic media
- Confirmation of capacity constraints
7. Special Processes

7.1 Special Process Approval

The use of Special Processes at suppliers and sub-tier suppliers requires approval by one of the following methods:

- Third party accreditations (e.g. NADCAP)
- Prior Lockheed Martin Special Process Approval (specific to Special Process)

Nadcap Welding approval requires additional scope coverage and as such must be audited by Lockheed Special Processes Supplier Quality Engineer (SQE).

Lockheed Martin reserve the right to audit suppliers Special Processes completed in house and at sub-tier suppliers / sub-contractors.

- If there are 0 non-conformances during LMUK-A audit approval is valid for 3 years.
- If non-conformances are identified approval is valid for 2 years once non-conformances are resolved before re-audit is required

Refer to section 7.2 for Special Process Validation requirements that suppliers shall complete.

Special processes include but are not limited to the following table:

<table>
<thead>
<tr>
<th>Coatings</th>
<th>Elastomer Seals</th>
<th>Electronics</th>
<th>Fluid Distribution Systems</th>
<th>Non-Metallic Materials</th>
<th>Sealants</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thermal Spray</td>
<td>Plate Seals</td>
<td>Printed Circuit Board (PCB) Manufacture</td>
<td>Hose Manufacturing</td>
<td>Resin Manufacturing</td>
<td>Hole / Slot Sealing</td>
</tr>
<tr>
<td>Vapour Deposited Coatings</td>
<td>Fabric / Textile Reinforced Seals</td>
<td>PCB Assembly (Incl. Soldering)</td>
<td>-</td>
<td>Prepreg Manufacturing</td>
<td>Wire Bundle Sealing</td>
</tr>
<tr>
<td>Diffusion Coatings</td>
<td>O-Rings</td>
<td>Cable and Harness Assemblies</td>
<td>Fitting and Coupling Manufacturing</td>
<td>Adhesive Film Manufacturing</td>
<td>Joggle Sealing (Pre – Packed and Injection)</td>
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<tr>
<td>-</td>
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<td>-</td>
<td>Liquid displacement or drain path sealing</td>
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<tr>
<th>Composites</th>
<th>Heat Treating</th>
<th>Materials Testing</th>
<th>Nonconventional Machining</th>
<th>Non-Metallic Materials Testings</th>
<th>Sealants</th>
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<tr>
<td>Prepreg</td>
<td>Brazing</td>
<td>Chemical Analysis</td>
<td>Electrochemical Machining</td>
<td>Mechanical Testing</td>
<td>EMI Seal Caps</td>
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<td>Resin Film Infusion (RFI)</td>
<td>Carburizing</td>
<td>Metallography</td>
<td>Electrical Discharge Machining</td>
<td>Chemical Testing</td>
<td>Sealing with Seal Caps</td>
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<tr>
<td>Metal Bonding</td>
<td>Gas/ion/Plasma Nitriding</td>
<td>Micro Indentation Testing</td>
<td>Laser Beam Machining</td>
<td>Thermal Testing</td>
<td>Fillet or Bead Sealing</td>
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<td>Core Processing</td>
<td>Hot Isostatic Pressing</td>
<td>Corrosion Testing</td>
<td>Laser Part Marking</td>
<td>Flammability Testing</td>
<td>Fay or Interfay Surface Sealing</td>
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</thead>
<tbody>
<tr>
<td>Electroplating</td>
<td>Rotational Friction / Inertia Welding</td>
<td>Penetrant Flaw Detect</td>
<td>Coordinate Measuring Machines (CMM)</td>
<td>Shot Peening</td>
<td>Battery Cell Manufacture</td>
<td>Forging</td>
</tr>
<tr>
<td>Electroless Plating</td>
<td>Torch / Induction Brazing</td>
<td>Anodise Flaw Detect</td>
<td>Laser Tracker</td>
<td>Peen Forming</td>
<td>Manual Peening</td>
<td></td>
</tr>
<tr>
<td>Anodising</td>
<td>Flash Welding &amp; Laser Welding</td>
<td>Magnetic Particle Inspection</td>
<td>Articulating Arm</td>
<td>Glass Bead Peening</td>
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<td></td>
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<tr>
<td>Chemical Conversion Coatings</td>
<td>Electron Beam Welding</td>
<td>Ultrasonic Testing</td>
<td>Mass Airflow Measurement of Turbine Parts</td>
<td>Automated Peening</td>
<td>Battery Cell Array Assemblies</td>
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<tr>
<td>Passivation</td>
<td>Resistance Welding</td>
<td>Radiographic Inspection Testing</td>
<td>-</td>
<td>Flapper Peening</td>
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<tr>
<td>Painting &amp; Dry-Film</td>
<td>Fusion Welding</td>
<td>Eddy Current Inspection Testing</td>
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<tr>
<td>Etching &amp; Chemical Cleaning</td>
<td>Evaluation of Welds</td>
<td>-</td>
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</tr>
</tbody>
</table>
7.2 Validation of Special Processes

Validation of a special process is critical and allows suppliers to demonstrate that the process provides a repeatable and reproducible output conforming to requirements.

A special process is a process that generates outputs that cannot be measured, monitored, or verified non-destructively or cost effectively. Therefore, deficiencies cannot be detected until after products are in use.

- Special processes **must** be periodically validated in order to prove that they can generate planned results
- Periodic validation is usually performed by the use of test coupons, verification tests, system accuracy tests or personnel qualification tests
- Suppliers and supplier sub-contractors providing special processes **shall** have a documented process control schedule (Process Control Document (PCD), Process Control Flow Chart (PCFC), job card traveller or similar) suitable of meeting all requirements prior to the commencement of production including all preparatory treatments, post treatments, processing, significant surfaces, tests and all other processes and treatments
- Suppliers **shall** ensure validation of sub-contracted Special Processes
- Refer to section 11.1.1 for paint system adhesion testing requirements
8. Quality Management Requirements

8.1 Review of the Requirements for Products and Services

The supplier shall ensure that they have the ability to meet LMUK-A’s requirements for products and services.

The supplier shall conduct a review before committing to supply products and services (commonly known as contract review).

If the review identifies any anomalies that could jeopardise the quality conformance or delivery of the Contract the supplier shall inform the nominated supply chain representative identified on the RFQ/contract.

This review will cover but not be limited to:

- Scope of certified approval against what product or service is being requested
- Technical ability i.e. can equipment or employee skills meet the requirements of the drawings.
- Capacity constraints
- Statutory and regulatory requirements
- Contract or order requirements differing from those agreed at tender
- **Drawing pack** i.e. tolerance, datum’s and geometric tolerancing, material requirements (ensuring the material is available in size and condition stated), special processes, specific drawing notes including adherence to standards and specifications quoted within the context, destructive and non-destructive testing requirements i.e. mechanical, electrical, software etc.
- Design and verification – if undertaking this requirement for LMUK-A, understand the conditions of the Contract as highlighted in para 8.3
- Reference documentation - it is the responsibility of suppliers to obtain, review, work to and maintain current or contracted issues of specifications and standards from appropriate sources
- Additional Resources – should the requirement for fixturing, hard gauging, specialist test equipment, specialised training etc be identified *this must be communicated to LMUK-A Supply Chain*. It is not acceptable if risk is identified and no action is undertaken or communicated to LM due to timescales or financial constraints. This will also apply to sub-contractors undertaking work on the product
- Supplier selection of sub-contractors. LMUK-A must be informed if sections of work are to be subcontracted. LMUK-A reserve the right to audit that supplier if it is deemed a perceived risk to contractual requirements (see 8.6). Special Processes are covered in section 7.

8.2 Deliverable Quality Plan

The quality plan is considered the key document which shall define all relevant standards and procedures to ensure that work is completed with zero defects to the required level of quality. The supplier must ensure that their own personnel are aware of the existence, purpose and content of the quality plan.
Suppliers shall submit a Deliverable Quality Plan (DQP) to LMUK-A in accordance with AQAP 2105 for QM003-A & ISO 10005 for QM003-B Contracts which describes the framework in which the Contract will be accomplished and is subject to approval by LMUK-A quality department.

- The DQP submission date shall be agreed with the LMUK-A Supplier Quality Representative post Contract award but shall be prior to manufacture of products.
- A revised DQP shall be submitted to LMUK-A when significant changes occur that invalidate the original submission.
- LMUK-A form F0389 Deliverable Quality Plan template may be used.

If only providing COTS/MOTS items to LMUK-A, review the application & extent of the DQP with your LMUK-A Supplier Quality Representative.

8.3 Design and Development Provision

If undertaking design and development work for LMUK-A, the supplier shall conform to the requirements defined within ISO9001 as a minimum, and AS9100 or AQAP 2110 for quality condition QM003-A.

8.4 Configuration Management

A disciplined Configuration Management process ensures that products conform to their requirements and are identified and documented in sufficient detail to support the product life cycle.

Configuration Management is critical to ensure the product and system interfaces remain compatible.

For Quality Conditions QM003-A and QM003-B the supplier shall plan, implement, and control a process for configuration management as appropriate to the products and services supplied to ensure the identification and control of physical and functional attributes through the product life cycle. This shall be in accordance with DEF STAN 05-57 for QM003-A & ISO 10007 for QM003-B Contracts.

For suppliers outside the United Kingdom configuration management in accordance with ACMP 2100 is acceptable for Quality Condition QM003-A.

8.5 Process Control and Verification

 Suppliers shall demonstrate confidence that their processes have been carried out as planned and the conformity of those products and services can be demonstrated.

For QM003-A & QM003-B Contracts Process Control & Verification activities shall be detailed in the Deliverable Quality Plan (see 8.2) and agreed with both parties. The requirements shall be flowed down the supply chain as necessary (see 8.6).

Examples of how to demonstrate process control include:

- Value Stream Mapping
- Process Flow Diagrams -identifying key characteristics, inspection stages, processes, frozen operations if identified by LMUK-A (no changes allowed unless agreed by LMUK-A delegate), associated documentation. Examples of compliant layouts may be requested from LMUK-A.
- PFMEA – Process Failure Mode Effect Analysis. Critical in analysis of the process flow showing anticipation of risks and actions to nullify those risks to the process
- Control plans – detailing the stages of the process where process monitoring, inspection and special process controls are required
- Inspections plans – identifying by whom (level of trained operators), with what (equipment’s to be used), how (standard operation) and the frequency of how those checks/inspection will be carried out
- Use of controlled fixtures and hard gauging

If Key Characteristics are specified, suppliers **shall** demonstrate the control of variation in the process by the use of process capability measurement, statistical process control, Measurement Systems Analysis studies i.e. Gauge Repeatability and Reproducibility (R&R) and Machine Capability Report (CmK).

### 8.6 Control of Externally Provided Processes, Products and Services

The supplier, as the recipient of the Contract, **shall** be responsible for meeting all requirements, including goods supplied, work performed, and services provided by the supplier’s sub-tier suppliers (also known as sub-suppliers or subcontract suppliers).

Where the supplier intends to sub-contract work or service normally undertaken by the supplier, a **written agreement shall be in place between LMUK-A and the supplier** indicating the reason for the sub-contract and the sub-tier sub-contractor to be used.

When the supplier uses sub-tier sources for goods supplied, work performed and services provided for LMUK-A, the supplier **shall** flow down to its sub-tier sources all of the applicable technical and quality requirements contained in the LMUK-A contract. This will:

- ensure that externally provided processes remain within the control of their own quality management system
- define both the controls that it intends to apply to an external supplier and those it intends to apply to the delivered product

LMUK-A representatives, customers and/or end users **shall** be allowed access to the sub-supplier’s plant and facilities for the purpose of surveillance and inspection.

For QM003-A & QM003-B Contracts suppliers **shall** confirm that Sub-Tier suppliers have QMS certification to ISO 9001 by a UKAS (or national equivalent) accredited certification body.

In exceptions where Sub-Tier suppliers do not hold ISO 9001 accreditation, the supplier **shall** liaise with their LMUK-A Supplier Quality Contact to evaluate where evidence of compliance can be provided.

### 8.7 Control of Equipment, Tools and Software

Equipment, tools and software programs used to monitor, measure, automate or control production processes **shall** be validated by the supplier prior to release for production and **shall** be maintained and calibrated (where applicable).

Traceability **shall** be maintained to national standards by supplier whether those items are calibrated internally or externally.
Storage requirements shall be defined for production equipment or tooling in storage including any necessary periodic preservation or condition checks.

Items that have an extended replacement period shall be detailed in a risk avoidance document that will detail the supplier’s disaster recovery plan in such an event – see section 20.

8.8 Competence, Training and Awareness

The supplier shall ensure personnel processing orders or performing work affecting conformity to product or service are trained and aware of the relevance and importance of their activities in relation to meeting the requirements of LMUK-A Contracts and associated documentation.

The supplier shall as a minimum produce a skills matrix which details the training undertaken by suppliers’ staff/operators in relation to the processes specific to LMUK-A product.

The skills matrix shall be maintained by supervisory/management level and demonstrate control of those activities. The skills matrix procedure shall also detail how risks associated with highly skilled staff are covered i.e. sickness, leave, succession planning etc. and the business decision on how that risk will be covered as well as capacity constraints.

8.9 Production Process Verification

When indicated on the purchase order or Contract the supplier shall produce a First Article Inspection Report (FAIR) to verify the manufacturing process using a representative item from the first manufacturing run of a new part or assembly.

- The FAIR shall be produced in accordance with AS9102
- A revised full FAI or partial FAI shall be completed when changes occur that invalidate the original results as per AS9102 section 4.6 requirements
- FAIRs shall be submitted electronically prior to delivery of goods to fai.fc-ampthill@lmco.com

The FAIRs shall include all certification indicating conformity of materials (including raw material mill test certificates), special processes, calibration, testing and personnel training qualification where applicable.

Guidance on how LM require First Articles to be completed is detailed in LM’s First Article Inspection Guidebook. It is strongly advised this is reviewed and its requirements understood.

8.10 Performance Evaluation

Performance evaluation shall be in accordance with ISO 9001 paragraph 9.

The supplier shall evaluate the performance and the effectiveness of the quality management system and retain appropriate documented information as evidence of the results. This information shall be accessible to LMUK-A upon request.

8.10.1 Monitoring, Measurement, Analysis and Evaluation

8.10.1.1 General

Comply with ISO 9001:2015 clause 9.1.1 requirements.

8.10.1.2 Customer Satisfaction

For QM003-A & QM003-B Contracts suppliers **shall**:

- Create production process performance metrics that monitor the following as a minimum unless otherwise agreed:
  1) Process yield rates (% scrap, % rework)
  2) Right First Time
  3) Statistical Process Control (if applicable on items with Key Characteristics)

- Implement appropriate Key Performance Indicators for quality and delivery performance

- Monitor performance metrics and use to drive process improvement initiatives

**8.10.1.3 Analysis and Evaluation**

Comply with the requirements defined in ISO 9001:2015 clause 9.1.3

**8.10.2 Internal Audit**

Comply with the requirements defined in ISO 9001:2015 clause 9.2

**8.10.3 Management Review**

Comply with the requirements defined in ISO 9001:2015 clause 9.3
9. Identification and Traceability

Traceability of the entire supply chain is of paramount importance to LMUK-A, including traceability to Original Equipment Manufacturer (OEM) where requirement applies for QM003-A & QM003-B Contracts.

9.1 Serialisation and Part Marking

Serialisation and part marking identification shall be in accordance with the Contract, design data, drawing or any contractually agreed specification or standard.

9.2 Traceability to Source/Origin of Raw Material

Where the delivery quality conditions are QM003-A or QM003-B and/or any applicable Quality Plan requires demonstration of traceability and design provenance through the supply chain, the supplier shall include in any relevant sub-contract the requirement for certification from its sub-tier suppliers.

Where QM003-A or QM003-B has been incorporated into the Contract, the supplier shall ensure that ‘Full Traceability’ is maintained within their own activities and throughout their sub-tier supply chain. The Supplier must provide traceability evidence to LMUK-A on request.

‘Full Traceability’ requirements for QM003-A & QM003-B Contracts are as follows:

<table>
<thead>
<tr>
<th>Category</th>
<th>Traceability Requirements</th>
</tr>
</thead>
<tbody>
<tr>
<td>Raw Material</td>
<td>Lot traceable to manufacturers part no, lot no, date code with Certified Mill Test Report or Material Test Report</td>
</tr>
<tr>
<td>Manufactured Parts</td>
<td>Traceability shall be maintained for all product throughout production from raw material to finished product (including product quantities, split orders, nonconforming product etc.)</td>
</tr>
<tr>
<td></td>
<td>Raw material used shall be Lot traceable to manufacturers part no, lot no, date code with Certified Mill Test Report or Material Test Report</td>
</tr>
<tr>
<td></td>
<td>Requirements specified for COTS, MOTS &amp; Mechanical/Electrical parts also apply where applicable.</td>
</tr>
<tr>
<td>Commercial off the Shelf (COTS)</td>
<td>Traceability through supply chain to Original Equipment Manufacturer (OEM)</td>
</tr>
<tr>
<td>Modified off the Shelf (MOTS)</td>
<td>Traceability shall be maintained for all product modifications throughout production from material to finished product (including product quantities, split orders, nonconforming product etc.)</td>
</tr>
<tr>
<td></td>
<td>The originating COTS item(s) shall have Traceability through supply chain to Original Equipment Manufacturer (OEM)</td>
</tr>
<tr>
<td></td>
<td>Requirements specified for Raw Material &amp; Mechanical/Electrical parts also apply where applicable for item modifications.</td>
</tr>
<tr>
<td>Mechanical / Electrical Parts</td>
<td>Lot traceable to Original Equipment Manufacturers part no, lot no, date code</td>
</tr>
</tbody>
</table>
10. Counterfeit Product Prevention

The supplier shall not deliver counterfeit or suspected counterfeit parts. The supplier shall inform LMUK-A of any suspected counterfeit parts immediately on discovery within the supplier’s supply chain.

The supplier shall have a defined and documented policy for the avoidance of counterfeit materiel, including an Anti-Counterfeiting Management Plan (ACMP) in accordance with DEF STAN 05-135.

The purpose of the supplier’s policy shall be to develop a robust process to prevent the delivery of counterfeit commodities and to control commodities identified as counterfeit.

Policies to AS5553 or AS6174 are also deemed acceptable providing the requirements of DEF STAN 05-135 are met. Semi-conductor distributors shall be certified to AS6081.

Product delivered by the Supplier that is subsequently identified as counterfeit by LMUK-A will be scrapped by LMUK-A and will not be returned to the Supplier. The Supplier shall replace any delivered counterfeit items at the Supplier’s own cost.
11. Preservation of Product

The supplier shall preserve the product during internal processing, storage and delivery to the intended destination.

11.1 Workmanship Acceptance Criteria for Surface Engineering

Unless otherwise stated, the following workmanship acceptance criteria shall be used:

- Supplied product with surface finishes for functional or cosmetic applications shall meet the requirements of the drawing (or referenced specification) for surface conditions i.e. uniform in appearance, free from blisters (adhesion), pits, nodules, scratches, stains. This includes but is not limited to electroplated, conversion coated, anodised, painted, mechanically finished and passivated surfaces.

11.1.1 Paint System Performance

Where products are painted, sufficient conditions shall be met to ensure effective adhesion is maintained.

For QM003-A & QM003-B Contracts this shall comply with one of the following acceptance criteria:

- ISO 2409 Table 1 Class 0
- ASTM 3359 Test Method A (X-Cut): 5A (No Peeling or removal)
- ASTM 3359 Test Method B (Cross Cut): 5B (0% Removed).

Adhesion testing shall be performed on test coupons representative of product substrate and process. Frequency of testing shall be adequate to validate the paint process output. Test records must be maintained traceable to the product.

Test reports shall be submitted in FAI packs when FAI is requested (ref. section 8.9) in accordance with the standard used for testing, and also be available upon request.

Refer to ISO 2409 paragraph 12 and ASTM 3359 paragraphs 9 & 14 for test report requirements relative to testing criteria.

11.2 Foreign Object Debris (FOD)

The supplier shall establish a process to detect and prevent Foreign Object Debris for QM003-A & QM003-B Contracts. This shall be in accordance with AS9146 or NAS412.

As a minimum the process shall include:

- FOD process review
- Training of FOD practices
- Material handling and product protection
- Tool / hardware accountability
- Lost items search and documentation process
- Physical entry control into FOD critical areas
- Inspection for foreign objects prior to closing apertures and compartments during assembly

The supplier shall review and apply FOD controls where appropriate for QM003-C Contracts.
11.3 Shelf Life

Goods and products containing items with finite shelf life shall have the expiry date identified on the product and the delivery documentation. The remaining shelf life must be a minimum of 80% of the total shelf life for the material at time of delivery unless otherwise specified.

11.4 Packaging

The supplier shall adequately plan and ensure that products are protected, packaged and labelled to a standard that will provide adequate protection against damage, deterioration, corrosion, contamination and loss.

The supplier shall ensure that the product packaging is labelled to a standard that provides adequate identification and traceability of the product.

The use of approved industry standard labelling and bar-coding shall be in accordance with any contractually agreed packaging specification.

If the contract specifies specific packaging requirements or standards these shall be complied with.

11.4.1 Moisture Sensitive Level (MSL)

Moisture sensitive components shall be packaged in accordance with IPC/JEDEC J-STD 033. The Moisture Sensitivity Level (MSL) must be clearly identified on the outer packaging.

11.4.2 Electrostatic Discharge (ESD)

Where appropriate, suppliers shall provide adequate protection measures against ESD damage to goods and LMUK-A property. This shall be in accordance with MIL-STD-1686 or ANSI/ESD S20.20. Electronic Components shall be handled, packaged and supplied in accordance with BS EN 61340-5-1 or as specified in the Contract conditions with LMUK-A.
12. Materials Database

LMUK-A maintain a Materials Database that specifies approved material grades for use based on a classification code system.

Where a material classification code is specified on drawing or specification, the Supplier must refer to the Lockheed Martin UK Ampthill Materials Database on the Lockheed Martin UK website to identify acceptable material grades for use.

When relevant, a specific Materials Database version reference shall be referenced within the Contract. Only the specific Materials Database version referenced on the Contract shall be used by the Supplier.

The materials database shall only be used where a related material classification code is specified on the drawing or contracted specification. It shall not be used for legacy drawings that specify a specific material specification.

All historical versions of the Materials Database version are available on the Lockheed Martin UK website.
13. Control of Non Conformance

13.1 Root Cause Analysis (RCA)

When nonconformities occur, the supplier must perform Root Cause Analysis (RCA) and corrective action activities to prevent recurrence of the problem.

LMUK-A recommends that the suppliers Improvement teams use industry standard root cause analysis tools to aid in identifying these issues i.e. 5 why methodology, 8D & Cause and Effect Diagram (Ishikawa or fishbone). Refer to AS13000 & ARP9136 for guidance.

Nonconformances on delivered hardware to Lockheed Martin are viewed as escapes by the Supplier, and as such will impact the Supplier’s performance rating. The Suppliers performance rating may influence LMUK-A’s future source selection decisions.

It is imperative that suppliers react to nonconformances and provide responses within agreed timeframes. Where timescales cannot be met the supplier shall inform the LMUK-A Supplier Quality Representative identified on the Supplier Corrective Action Report (SCAR) form immediately.

LMUK-A will inform the supplier of nonconformities that are highlighted at any stage of LMUK-A’s process flow including, but not limited to, trials and subsequent service. The supplier shall respond to the SCAR when raised. The SCAR is structured around the 8D process which details the requirement for the following:

- Problem statement
- Containment Action (in production, in stores, in transit, delivered product)
- Root Cause Analysis
- Corrective Action
- Implement Corrective Action
- Define and Plan Preventative action to prevent recurrence
- Review of Implementation or actions

Unless otherwise communicated by your LMUK-A Supplier Quality representative, SCARs shall be processed to the following timescales by the supplier.

- Supplier has 5 working days to acknowledge receipt, undertake containment action and provide containment response to LMUK-A
- Supplier then has a further 25 calendar days to respond with a detailed corrective action plan on the LM Supplier Corrective Action Form A-0108
- Supplier will submit on or before the agreed verification date, evidence of the implemented corrective & preventative action. This evidence will allow the LMUK-A Supplier Quality Engineer to close the SCAR
- Should the SCAR be rejected by LMUK-A the supplier will have a further 5 working days to re-submit for approval and closure
14. Application for Production (Deviation) Permit or Concession

Suppliers *shall* generate the Production Permit or Concession in accordance with LMUK-A Form F0045, or their own form provided it incorporates the requirements listed in DEF STAN 05-61 Part 1.

This *shall* include the proposed corrective action to eliminate the cause and prevent recurrence.

14.1 Production Permit

Production Permits (Deviations) are considered permission to produce an item that deviates from design data. This *may* be because of design anomalies, material availability issues or other unforeseen reasons prior to manufacture.

- Completed production permits *shall* be submitted to the supply chain representative indicated on the Contract / purchase order
- *All Production permits must be referenced on the applicable certificate of conformity* quoting the permit approval number provided by LMUK-A
- Any production prior to production permit approval is entirely at the supplier’s own risk
- Products delivered against a LMUK-A approved permit are considered as conforming.

14.2 Concessions

Requests for concession applications will only be accepted under exceptional circumstances. Concession submissions will impact the supplier performance rating.

Prior to submission of concession application mitigation actions *shall* be reviewed by the supplier to remove the need for non-conformance and be available upon request by LMUK-A. If concession is still viewed as necessary, a discussion *shall* be held with the LMUK-A supply chain representative indicated on the Contract / purchase order before submission.

* The concession *must* quote the LMUK-A Contract / purchase order(s) reference, serial numbers or batch affected or, if required, the time span applicable. The concession will NOT be considered for acceptance by LMUK-A if these conditions are not met
* Completed concession request forms *shall* be submitted to the LMUK-A supply chain representative indicated on the Contract
* Delivery of nonconforming product *shall* not occur unless an approved concession is in place. *All concessions must be referenced on the applicable certificate of conformity* using the concession approval number provided by LMUK-A.
15. Release of Product and Services

Suppliers shall supply conforming goods and services on time in full (OTIF) including all required correct documentation and certification where applicable.

Certification refers to any document that states the goods or services meet or conform to specification or contract requirements. These include, but are not limited to; Certificate of Conformity, Certificate of Compliance, Certificate of Analysis, Certificate of Attestation and Certificate of Calibration.

The certifying document shall be deemed as an authorised contractual guarantee that the goods and services reference on the certificate meet drawing, specifications, technical data and contract requirements. A signed copy or digital signature will be acceptable, but Certificates must be traceable to a certifying supplier quality representative or authorised supplier company official.

15.1 Supplier Documentation (Certificate of Conformity Requirements)

The following data/information shall be included on each certification document (normally referred to as a CoC or Release Note)

* Certificate or delivery unique identifier / Certification / Delivery Note number
* Certificate Date
* Purchase order / contract number
* Drawing number and / or part number and revision (as stated on Purchase Order or Sub-Contract)
* Batch unique identifier (Batch number / Lot number / Date code / Serial number)
* Quantity
* Supplier Name and Address
* Statement that goods and / or services conform to the specified requirements
* Original Manufacturer’s name, part number and lot / date code (when applicable)
* LMUK-A Reference numbers for all approved Production Permits / Concessions applicable
* Reference to current First Article Inspection Report (where applicable)
* Reference to the Quality Management System release

Suppliers shall ensure the correct documentation is supplied with products and services. This is further illustrated in Diagram 2 – Delivery documentation requirement flowchart.

15.1.1 Certificate of Conformity

For deliveries that apply quality condition QM003-A & QM003-B (see section 4 - Quality Conditions (Q Code) a certificate of conformity shall be supplied with delivered goods or services that meet the above Certificate of Conformity requirements. Please read section 17 which further expands the requirements for Chemicals and Hazardous Substances.
15.1.2 Delivery Advice Note

The following data/information **shall** be included on each Delivery Advice Note when required:

* Delivery Advice Note number
* Release date
* Purchase order / contract number
* Quantity
* Supplier Name and Address
* Drawing number and / or part number (as stated on Purchase Order or Sub-Contract)
* Batch unique identifier (Batch number / Lot number / Date code / Serial number) if applicable. This is not mandated if detail captured on supporting C of C provided for QM003-A/B Contracts.

15.1.3 Calibration and Test Certification

Where calibration and test certification are issued to LMUK-A information **shall** include:

* The calibrated test apparatus / instrument / standard used. These will be traceable to UKAS or the national equivalent from sources other than the UK
* Calibration / test specification used to include tolerances and criteria
* Items outside specified limits will be identified, especially if the item has undergone authorised repair to bring it into specification

Calibration and test certification **shall** be submitted electronically prior to shipment of item to calibration.fc-amphill@lmco.com. Delivery note **shall** be provided with physical delivery.

15.1.4 Completeness of Supplied Documentation

Certification documentation supplied to the requirements of any LMUK-A contract will be rejected and deemed not complete should it transgress any of the following:

* Certification supplied with CofC is illegible i.e. faint, blurred or ambiguous
* Certification supplied with concession/production permit whose approval is outstanding
* Incorrect / different material or subcontracted special process certification being referenced that do not tie up with FAI documentation
* Alternative material and or subcontracted special processes – will be rejected if authorised certification is not attached to the CofC i.e. production permit or concession approved with the prior agreement of LMUK-A.

15.1.5 Late Deliveries / Short Deliveries

If non-delivery, short or late deliveries are anticipated, suppliers **shall** immediately notify the LMUK-A supply chain representative indicated on the Contract.
Diagram 2 – Delivery documentation requirement flowchart

* First Article Inspection Reports shall be submitted electronically prior to shipment of goods to fai.fc-ampthill@lmco.com

* Calibration and test certification shall be submitted electronically prior to shipment of item(s) to calibration.fc-ampthill@lmco.com
16. Legislative Requirements

Suppliers *shall* conform to all applicable Environmental, Health and Safety legislation as applicable to the Contract and retain appropriate compliance data as defined by legislation.

Where specified in legislation or Contract appropriate compliance data *shall* be supplied to customer (LMUK-A).
17. Chemicals and Hazardous Substances

Nothing in this section shall reduce or limit any statutory duty or legal obligation of LMUK-A or the supplier.

17.1 Safety Data Sheets

If the supplier is required, under, or in connection with the contract, to supply articles or components of articles that, in the course of their use, maintenance, disposal, or in the event of an accident, may release hazardous materials or substances, they shall provide to LMUK-A a list of those hazardous materials or substances with a subsequent Safety Data Sheet.

Safety data sheets (SDS) provide information on chemical products that help users of those chemicals to make a risk assessment. They describe the hazards the chemical presents, and give information on handling, storage and emergency measures in case of accident.

By law suppliers of chemicals must provide an up to date safety data sheet if a substance is classified as dangerous in accordance with the Classification, Labelling and Packaging (CLP) Regulation 1272/2008.

17.2 Registration, Evaluation, Authorisation and Restriction of Chemicals (REACH)


The regulation aims to better protect human health and the environment by ensuring the risks associated with chemicals are measured and understood. REACH calls for registration of chemicals above specific thresholds that are imported and/or manufactured within the European Economic Area (EEA), chemical disclosure requirements for articles that are imported and/or manufactured within the EEA, and the complete restriction of some chemicals from the EEA altogether.

REACH applies to substances manufactured or imported into the EU in quantities of 1 tonne or more per year. Generally, it applies to all individual chemical substances on their own, in preparations or in articles.

The supplier shall disclose such information to LMUK-A for the purpose of compliance with the REACH regulation and comply with the requirements of Regulation (EC) No 1907/2006.

Lockheed Martin has established a Corporate REACH Program Office within its Global Supply Chain Operations organisation that is fully conversant with the requirements of REACH to ensure compliance throughout the Corporation and within our supply chain.

For more information please contact the Lockheed Martin REACH Program Office at reach.info@lmco.com.

17.3 Lead, Asbestos and Radioactive Substances

Special regulations apply to Lead, Asbestos and radioactive substances. In addition, refer to DEFCON 624 for Asbestos. Adequate packaging must be provided to prevent exposure of LMUK-A staff to these substances in accordance with the relevant Health and Safety Executive (HSE) Approved code of practice (ACOP).
17.4 Restriction of Hazardous Substances (RoHS)

RoHS stands for Restriction of Hazardous Substances. RoHS originated in the European Union and restricts the use of specific hazardous materials found in electrical and electronic products (known as EEE). All applicable products in the EU market after July 1, 2006 must pass RoHS compliance.

Directive 2011/65/EU lays down rules on the restriction of the use of hazardous substances in electrical and electronic equipment (EEE) with a view to contributing to the protection of human health and the environment, including the environmentally sound recovery and disposal of waste EEE.

RoHS is a CE-marking directive. As such, all manufacturers of electrical/electronic products must comply with RoHS before the CE mark can be applied on their products and introduced to the EU market.


Importers & Distributors of electrical and electronic equipment shall be fully cognisant of the requirements and comply with EU Directives 2011/65/EU, 2017/2102 & 2105/863.
18. Munitions

18.1 Weapons

Suppliers transporting weapons or weapon component parts must hold a current Home Office approval to transport goods controlled under the Firearms Act 1968, Section 5 (as amended).

Home Office Guidance can be found here.

18.2 Explosives

For explosive items, the supplier shall contact Head of Safety Services at LMUK-A, 3 business days prior to delivery.

Any delivery & storage of explosives must be confirmed by LMUK-A that it is covered by LMUK-A explosives licence prior to delivery.

Release documentation for explosive items must include a copy of the National Competent Authority Classification and a Certificate of Correctness signed by the supplier’s authorised person.

Release documentation must include the National Competent Authority Classification Number, United Nations (UN) Number, Hazard Division and Net Explosive Quantity.

All packages must bear a UN Mark and be classified and labelled in accordance with UK Government Statutory Instrument 1994 No. 669 (The Carriage of Dangerous Goods by Road and Rail [Classification, Packaging and Labelling] Regulations 1994).

UK Government Statutory Instrument 2014 No. 1638 (The Explosives Regulations 2014, Health and Safety) shall apply to all explosive items.

Packaging shall be in accordance with DEFCON 130 Packaging for Explosives.
19. Obsolescence Management

Obsolescence Management is ‘the co-ordinated activities to direct and control an organisation with regard to obsolescence’. Obsolescence Management seeks to reduce the risk of obsolescence through undertaking planned activities to reduce the frequency of obsolescence issues and/or reduce the impact when an item becomes obsolete.

Suppliers **shall** notify LMUK-A of any pending obsolescence, the relevant last time buy date and last time ship date at least 6 months prior to the last time buy date.

Suppliers should refer to IEC 62402 for further guidance on Obsolescence Management.

20. Business Continuity / Disaster Management

For QM003-A & QM003-B contracts suppliers **shall** have in place a business continuity plan. This **should** be in accordance with ISO 22301. This includes requirements to plan, establish, implement, operate, monitor, review, maintain and continually improve a documented management system to protect against, reduce the likelihood of occurrence, prepare for, respond to, and recover from disruptive incidents when they arise.

Essentially this is a management plan that ensures no disruption to the supply of goods to LMUK-A should the business fall foul of environmental circumstances such as fire, flood, power failure etc. This will include but not be limited to safety stocks of goods, fire protection of tooling/Jigs, safeguarding essential key machinery, off site holding of key software etc.
21. Useful Resources

21.1 International Aerospace Quality Group

It is highly recommended that the free resources provided by the Society of Automotive Engineers (SAE) International Aerospace Quality Group (IAQG) in the form of the Supply Chain Management Handbook (SCMH) is utilised to its full potential by all suppliers.

This online document contains invaluable training and guidance material on every element of Aviation, Space and Defence (AS&D) requirements including first article inspection, configuration management, quality plans, counterfeit management and contract review.

21.2 Lockheed Martin Support

If there are any questions in relation to this document or requirements the applicable LMUK-A Supplier Quality representative shall be contacted.

Alternatively contact sqa_ampthill.fc@lmco.com.

21.3 Lockheed Guidance Documents & Templates

Guidance documentation & templates (including First Article Inspection Guide & Deliverable Quality Plan Template) are available on the Lockheed Martin UK Suppliers webpage.
Appendix 1 – Additional AQAP Guidance notes

AQAP 2110 EDITION D Quality Requirements for Design, Development and Production
(ref QM003-A – Quality Requirements for NATO Contracts)

Allied Quality Assurance Publications (AQAP) are standards for quality assurance systems that have been developed by the North Atlantic Treaty Organization (NATO). AQAP 2110 Quality Assurance Requirements for Design, Development and Production defines requirements, which, if applied appropriately, provide confidence in the supplier's capability to deliver products that conform to LMUK-A contract requirements.

The common ISO 9001 baseline inherently makes AS9100 and AQAP 2110 appear almost identical. It is acceptable for a supplier to offer a QMS that complies with the provisions of AS9100 as a satisfactory response to the QMS requirements of AQAP 2110, under two conditions:

- The supplier formally states that, “All AS9100 requirements applicable to the organization are applicable to this contract”;
- The supplier formally states that, “No exclusions to AS9100 taken by the organization shall in any way diminish, alter, or relieve the AQAP 2110 requirements of this contract”.

These formal statements shall be made in the Deliverable Quality Plan (DQP). See BS EN 9137 or ARP9137 Guidance for the Application of AQAP 2110 within a Quality Management System for further information.

AQAP 2120 Quality Requirements for Production

AQAP 2130 Quality Requirements for Inspection and Test

AQAP 2210 Supplementary Software Quality Assurance Requirements
(as per QM003-A – Quality Requirements for NATO Contracts)

AQAP 2210 is intended for use with AQAP 2110 as a software specific and project-oriented supplement and defines the requirements for the Software Quality Management Activities as related to the Project to be documented in a Software Project Quality Plan. These activities are based on the supplier’s software quality system. The publication also requires the evaluation of the software quality management activities to ensure their effectiveness.

AQAP 2105 Deliverable Quality Plan
(as per QM003-A – Quality Requirements for NATO Contracts)

Where required suppliers shall submit a Deliverable Quality Plan (DQP) in accordance with AQAP 2105 which describes the framework in which the contract will be accomplished and is subject to approval by LMUK-A quality department.

The quality plan is considered as the key document which shall define all relevant standards and procedures to ensure that work is completed successfully to the required level of quality.

The supplier must ensure that their own personnel are aware of the existence, purpose and content of the quality plan. Form F0389 Deliverable Quality Plan template may be used to meet this requirement.

AQAP 2131 Quality Requirements for Final Inspection
AQAP 2131 does not apply within the scope of this document.
AQAP 2310 Quality Requirements for Aviation, Space and Defence Suppliers

AQAP 2310 is invoked when suppliers hold AS9100 (or equivalent) certification where AQAP 2110 does not apply.

QUALITY PLAN (ISO 10005)
(as per QM003-B – Quality Requirements for non-military contracts)

Where required suppliers shall submit a (Deliverable) Quality Plan (DQP) in accordance with AQAP standards or as detailed with the contractual requirements which describes the framework in which the contract will be accomplished and is subject to approval by LMUK-A quality department.

The quality plan is considered as the key document which shall define all relevant standards, procedures and specific customer requirements to ensure conformance to requirements.

The supplier must ensure that their own personnel are aware of the existence, purpose and content of the quality plan. Form F0389 Deliverable Quality Plan template may be used to meet this requirement.

ISO 10007 CONFIGURATION MANAGEMENT PLAN
(as per QM003-B – Quality Requirements for non-military contracts)

Where required suppliers shall prepare and operate a Configuration Management Plan, which describes the application of configuration management to the contract in accordance with AQAP standards or as detailed with the contractual requirements.
Appendix 2 – Glossary of Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation / Acronym</th>
<th>Definition</th>
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<tbody>
<tr>
<td>A-0108</td>
<td>Supplier Corrective Action Report Form – structured report to suppliers informing them of nonconformities that have been highlighted</td>
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<tr>
<td>ACMP</td>
<td>Anti-Counterfeiting Management Plan</td>
</tr>
<tr>
<td>COTS</td>
<td>Commercial Off the Shelf</td>
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<tr>
<td>DEF-STAN</td>
<td>Defence Standards</td>
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<td>DQP</td>
<td>Deliverable Quality Plan</td>
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<td>F0045</td>
<td>LMK-A form for Application for a Deviation Permit or Concession – complies with DEF-STAN 05-61 iss.6</td>
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<tr>
<td>F0183</td>
<td>Supplier Assessment Questionnaire – required to complete supplier approval</td>
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<td>F0189</td>
<td>Supplier Assessment Report – may be required to complete supplier approval</td>
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<tr>
<td>F0389</td>
<td>Deliverable Quality Plan Template – optional template that suppliers may use</td>
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<tr>
<td>FAI</td>
<td>First Article Inspection</td>
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<tr>
<td>FAIR</td>
<td>First Article Inspection Report</td>
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<tr>
<td>FOD</td>
<td>Foreign Object Debris</td>
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<tr>
<td>GQAR</td>
<td>Government Quality Assurance Representative</td>
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<td>LM</td>
<td>Lockheed Martin</td>
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<tr>
<td>LMPI</td>
<td>Lockheed Martin Proprietary Information</td>
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<td>LMK-A</td>
<td>Lockheed Martin UK Ampthill Limited</td>
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<td>MLA</td>
<td>Manufacturing Licence Agreement</td>
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<td>MOTS</td>
<td>Modified Off the Shelf</td>
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<td>MSL</td>
<td>Moisture Sensitive Level</td>
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<td>NDA</td>
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<td>OEM</td>
<td>Original Equipment Manufacturer</td>
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<td>PCD</td>
<td>Process Control Document</td>
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<td>PCFC</td>
<td>Process Control Flow Chart</td>
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<td>Process Failure Mode Effect Analysis</td>
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<td>QM003</td>
<td>This document – LMK-A’s Quality Requirements for Suppliers</td>
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<td>QMS</td>
<td>Quality Management System</td>
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<td>RCA</td>
<td>Root Cause Analysis</td>
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<td>SQE</td>
<td>Supplier Quality Engineer</td>
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<td>SSCM</td>
<td>Sustainable Supply Chain Management</td>
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<tr>
<td>Supplier</td>
<td>Vendor, supplier of goods and services, sub-contractor and distributor</td>
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<tr>
<td>TAA</td>
<td>Technical Assistance Agreement</td>
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<td>TCP</td>
<td>Technology Control Plan</td>
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<tr>
<td>UKAS</td>
<td>The UK’s National Accreditation Body, responsible for determining, in the public interest, the technical competence and integrity of organisations such as those offering testing, calibration and certification services</td>
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### Appendix 3 – Standards Referenced

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<tr>
<td>ACMP 2100</td>
<td>Configuration Management Contractual Requirements</td>
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<td>ANSI/ESD S20.20</td>
<td>Protection of Electrical and Electronic Parts, Assemblies and Equipment (Excluding Electrically Initiated Explosive Devices)</td>
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<td>AQAP 2105</td>
<td>NATO Requirements for Quality Plans</td>
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<td>AQAP 2110</td>
<td>NATO Quality Assurance Requirements for Design, Development and Production</td>
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<td>AQAP 2120</td>
<td>NATO Quality Assurance Requirements for Production – Superseded by AQAP 2110 Edition D (2016)</td>
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<td>AQAP 2130</td>
<td>NATO Quality Assurance Requirements for Inspection and Test - Superseded by AQAP 2110 Edition D (2016)</td>
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<td>AQAP 2131</td>
<td>NATO Quality Assurance Requirements for Final Inspection</td>
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<td>NATO Quality Assurance Requirements for Aviation, Space and Defence Suppliers</td>
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<td>ARP9136</td>
<td>Aerospace Series – Root Cause Analysis and Problem Solving (9S Methodology)</td>
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<td>Guidance for the Application of AQAP 2110 within a 9100 Quality Management System</td>
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<td>AS5553</td>
<td>Counterfeit Electronic Parts; Avoidance, Detection, Mitigation, and Disposition</td>
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<td>AS6081</td>
<td>Fraudulent/Counterfeit Electronic Parts: Avoidance, Detection, Mitigation, and Disposition – Distributors</td>
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<td>AS6174</td>
<td>Counterfeit Materiel; Assuring, Acquisition of Authentic and Conforming Materiel</td>
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<td>AS9100</td>
<td>Quality Management Systems – Requirements for Aviation, Space and Defense Organizations</td>
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<td>AS9102</td>
<td>Aerospace First Article Inspection Requirement</td>
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<td>Foreign Object Damage (FOD) Prevention Program – Requirements for Aviation, Space, and Defense Organizations</td>
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<td>ASTM3359</td>
<td>Standard Test Methods for Rating Adhesion by Tape Test</td>
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<td>BS EN 61340-5-1</td>
<td>Electrostatics. Protection of Electronic Devices from Electrostatic Phenomena. General Requirements</td>
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<td>Standard for Handling, Packing, Shipping and Use of Moisture/Reflow Sensitive Surface Mount Devices</td>
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<td>ISO2409</td>
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<td>General Requirements for the Competence of Testing and Calibration Laboratories</td>
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### 22. QM003 Document Changes

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<td>Complete re-write of Quality Requirements for Suppliers</td>
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<td>5.1, 5.2, 5.3 &amp; 5.6</td>
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<td>Part marking as per purchase order, design data or drawing added</td>
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<td>The supplier shall ensure that full traceability is maintained and can be provided on request throughout the sub-tier supply chain added</td>
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<td>13/03/2017</td>
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<td>Certificates must be traceable to the certifying quality representative or company official on request added</td>
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<td>Reference to raw material certification where applicable and name of authorised certifying quality representative or company official removed</td>
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<td>Shelf life requirement increase to 80% unless otherwise specified</td>
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<td>Figure 2 Delivery Documentation Requirement Flowchart added</td>
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<td>Requests for deviation permit or application for concession amended in line with current process</td>
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<td>31/10/2017</td>
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<td>Restructure of Document to align with ISO 9001 and AS9100 requirements</td>
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<td>05/01/2018</td>
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<td>30/05/2018</td>
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<td>4.11.2</td>
<td>Addition of Production Permit term</td>
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<td>Appendix 2</td>
<td>First Article Requirement clarification on flowchart</td>
<td>Craig White</td>
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<td>03/07/2018</td>
<td>7</td>
<td>Appendix 3</td>
<td>New section added to Appendix 3 to include Special Process Document References &amp; update to Appendix 2</td>
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<td>27/02/2019</td>
<td>8</td>
<td>3.2 &amp; Appendix 3, Multiple</td>
<td>Re-write of Special Process Approvals and amendment to table &amp; Hyperlinks to LM web resources updated and obsolete hyperlinks removed</td>
<td>Craig White</td>
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<td>21/12/2020</td>
<td>9</td>
<td>Multiple</td>
<td>Substantially re-written and restructured.</td>
<td>Jon Hood / Will Cullen</td>
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