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Pages 1 to 32

CHECKLIST FOR SURFACE ACOUSTIC

WAVE (SAW) DEVICES

MANUFACTURER AND LINE SURVEY

ESA/SCC Basic Specification No. 2023502

Manufacturer :
Location :
Survey Team Leader :
Date of Survey :
SAW Device Type(s) :



**space components
coordination group**

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		SCCG Chairman	ESA Director General or his Deputy
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SCC

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No. 2023502

PAGE 2

ISSUE 1

DOCUMENTATION CHANGE NOTICE

Rev. Letter	Rev. Date	Reference	CHANGE Item	Approved DCR No.

**SCC**ESA/SCC Basic Specification
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PAGE 3

ISSUE 1

TABLE OF CONTENTS

	<u>Page</u>
1. <u>INTRODUCTION</u>	4
2. <u>SURVEY CHECKLIST</u>	4
2.1 Interview on Arrival of Survey Team	4
2.2 Manufacturer and Survey Team Information	4
2.3 Management Organisation	6
2.4 Quality Assurance System and Organisation	8
2.5 Calibration	10
2.6 Drawing and Change Control	11
2.7 Reliability	12
2.8 Control of Procurement Sources	14
2.9 Control of Incoming Materials	15
2.10 In-process Inspections and Tests	17
2.11 Survey of Manufacturing Line	19
2.11.1 Manufacturing Environment	19
2.11.2 Fabrication of Substrate	20
2.11.3 Metallisation of Substrate	20
2.11.4 Bonding of Substrate	22
2.11.5 Lead-bonding	23
2.11.6 Pre-seal Device Preparation	23
2.11.7 Enclosure of Devices	24
2.11.8 Visual Inspection - General	24
2.11.9 Final Test Area and Screening Facility	24
2.12 Preservation, Packing and Shipping	29
2.13 Summary of Inspection Results	31
2.14 General Observations	32

1. INTRODUCTION

This checklist is intended for use during the initial survey of a Manufacturer's ability to produce high quality articles, his management organisation, production facilities, test facilities and technical know-how. When completed, this checklist should enable the party interested in procurement of the subject components to assess the ability of the Manufacturer concerned to successfully execute a contract for the supply of high reliability space hardware.

2. SURVEY CHECKLIST

2.1 INTERVIEW ON ARRIVAL OF SURVEY TEAM

(a) Introductory Remarks by Team Leader (Explanation of purpose of survey, procedures to be followed, time limitations, etc.):-



(b) Notes (Atmosphere during reception, willingness to co-operate, interest shown, comments on personnel, general remarks):-

2.2 MANUFACTURER AND SURVEY TEAM INFORMATION

(a) Survey requested by :
 Survey Team Leader :
 Team Members :

(b) Key personnel of Manufacturer interviewed:-

Name	Function	Tlph. Ext.
1.		
2.		
3.		
4.		
5.		

 	<p style="text-align: center;">ESA/SCC Basic Specification No. 2023502</p>		<p>PAGE 5 ISSUE 1</p>
---	--	--	---------------------------

(c) Type of Company (Private company, limited company, etc.)

Affiliated with any other company? If so, which:

No. of employees:

- Total number :
- Production :
- Quality Assurance :
- Q.A. Inspection :
- Prod. Engineering :
- Design Engineering :
- Reliability Control :
- Other :

(d) Number of shifts :

(e) Plant area :

(f) General production line :

(1) Device types manufactured:

(2) Will flow diagrams of steps to produce SAW devices be available to Survey Team?

YES NO

Are specifications, if any, referenced in the flow diagrams?

YES NO

(g) Principal Government and industrial customers:-

- 1.
- 2.
- 3.
- 4.
- 5.

(h) The Manufacturer's Quality System is organised in accordance with:

Comments

(i) Manufacturer's Government Service Inspection:

DCAS Inspector, resident/non-resident

(j) National Inspectorate:

(k) Is the Manufacturer's SAW device production

(1) Continuous?	YES	NO
(2) Pilot production?	YES	NO
(3) Advanced R&D, limited?	YES	NO

(l) The Manufacturer has adequate experience in the production of the following hi-rel parts:-

2.3 MANAGEMENT ORGANISATION

(a) What is general policy/attitude of the Management regarding quality/reliability programme?



(b) Which level of Management participates actively in orientating policy towards space component production?

(c) Which organisation, if any, reviews and monitors all work involved in space component production?

(d) Is work related to space components (contracts) regarded as "normal business" or as belonging to the "unique order" category?

(e) What is the general policy concerning proprietary rights?

(f) Has the "Reliability" department the same authority from Management as the "Engineering" and "Production" departments? Does this mean direct responsibility for reliability of products in the line?

 	<p>ESA/SCC Basic Specification No. 2023502</p>		<p>PAGE 7 ISSUE 1</p>
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(g) Has the Q.A. Manager direct authority for implementation of quality policy and actions related to the line?

(h) Does a system exist for the regular supply of quality report summaries to Management?

Does this system lead to (corrective) actions being taken in respect of the production line?

(i) Are key management staff notified of persistent out-of-control conditions?

(j) What is length of service and experience of key management personnel (Q.A., Reliability, Production, Engineering Design)?

(k) How would contract for space components be organised?

(l) How can original requirements from Orderer (Space Agency or end-user) be assumed to be correctly translated into internal instructions?

(m) How can information necessary to the Orderer (corrective actions, deviations, notification of inspections and/or problem areas) be assumed to be issued and channelled to the Orderer?

2.4 QUALITY ASSURANCE SYSTEM AND ORGANISATION

- (a) To whom does Q.A. Manager report?
- (b) Does the company reflect a positive attitude towards Quality Assurance? YES NO

 Comments
- (c) Has the Q.A. group sufficient authority in relation to its position within the company's organisation (see organigram)? _____
 Comments
- (d) Are areas of responsibility within the Q.A. group clearly defined? _____
 Comments
- (e) Are corrective actions to which Q.A. management is committed delegated to responsible staff or does Q.A. management have direct authority regarding the line? _____
 Which?
- (f) Is there a periodic and comprehensive quality data reporting system which covers all operational phases? _____
 Comments
- (g) What is the relationship between Q.A. and Reliability?
- (h) Is a Q.A. manual or equivalent document supplied to all levels of appropriate supervisory personnel? _____
 Is such document kept updated? _____
 Comments

- | | YES | NO |
|---|-----|-----|
| (i) Are written procedures available for identification and positive control of accepted/rejected materials? | ___ | ___ |
| Comments | | |
| (j) What is ratio Q.A. inspectors : personnel directly involved in production? | | |
| (k) Is inspection (acceptance sampling or sorting) performed by Q.A. personnel:- | | |
| On receipt? Sampling ___ Sorting ___ None ___ | | |
| During processing? Sampling ___ Sorting ___ None ___ | | |
| During final testing? Sampling ___ Sorting ___ None ___ | | |
| Comments | | |
| (l) Are written procedures kept and used in areas for:- | | |
| Receiving inspection? | ___ | ___ |
| In-process inspection? | ___ | ___ |
| Fabrication processing? | ___ | ___ |
| Final testing? | ___ | ___ |
| Comments | | |
| (m) Does Q.A. maintain a system of written procedures for statistic controls (control chart, lot plot, etc.) in any of the following areas? | | |
| In-process inspection? | ___ | ___ |
| Fabrication processing? | ___ | ___ |
| Final inspection? | ___ | ___ |
| Comments | | |
| (n) Is Q.A. responsible for determination of need for, and the conducting of, quality training? | ___ | ___ |
| Comments | | |
| (o) Are training programmes provided for special process personnel? | ___ | ___ |
| Comments | | |

- | | YES | NO |
|--|-----|-----|
| (p) Do employees have to pass tests:- | | |
| After training? | ___ | ___ |
| Periodically? | ___ | ___ |
| Comments | | |
| (q) Are production operators provided with visual aids and working instructions? | ___ | ___ |
| Comments | | |

2.5 CALIBRATION

- | | | |
|--|-----|-----|
| (a) Does Manufacturer maintain calibration facilities and standards? | ___ | ___ |
| Is this service purchased? | ___ | ___ |
| If so, from whom? | | |
| (b) Do calibration personnel have written procedures for control and a time schedule for measurement frequency? | ___ | ___ |
| Comments | | |
| (c) Is there an effective calibration record control system? | ___ | ___ |
| (d) Are calibration procedures adhered to and up-to-date? | ___ | ___ |
| Comments | | |
| (e) Are decals used for equipment identification to show that units have been calibrated; when next calibration date is due and calibrator identification? | ___ | ___ |
| Are decals up-to-date? | ___ | ___ |
| (f) Are adjustments of calibrated equipment required to be sealed and tamper-proof? | ___ | ___ |
| (g) Who is in charge of initiating calibration steps? | | |
| User | ___ | ___ |
| Calibration personnel | ___ | ___ |
| Q.A. | ___ | ___ |

- | | YES | NO |
|--|-----|-----|
| (h) Do calibration procedures provide for removal of any equipment not maintained or calibrated according to established schedules?
Comments | --- | --- |
| (i) Have calibrating personnel up-to-date certification records reflecting date, traceability to NBS and identification of calibrator?
(1) Mechanical standard?
(2) Electrical standard? | --- | --- |
| (j) Is modified and/or repaired equipment calibrated prior to release? | --- | --- |

2.6 DRAWING AND CHANGE CONTROL

- | | | |
|---|-----|-----|
| (a) Has Manufacturer adequate written procedures for control of specification and contract changes?
Comments | --- | --- |
| (b) Does Manufacturer's system provide for documented change control guaranteeing availability of required drawing at relevant manufacturing or inspection step?
Do flow documents show current revisions?
Comments | --- | --- |
| (c) Are drawings furnished by ESTEC and contract changes adequately controlled?
Comments | --- | --- |
| (d) Does Q.A. review all drawings and changes therein prior to their becoming effective?
Comments | --- | --- |
| (e) Has Manufacturer established a procedure for notifying his Supplier of changes in drawings?
Comments | --- | --- |
| (f) Are current specification revisions shown on prints of drawings? | --- | --- |

2.7	<u>RELIABILITY</u>	YES	NO
	(a) Is structure of Reliability organisation clearly defined?	___	___
	Has Reliability same authority in respect of the line as Production or Engineering management?	___	___
	Comments		
	(b) Is there a direct feed-back of information between Reliability, Design Engineering and Q.A. groups to ensure timely notification of all relevant data?	___	___
	Comments		
	(c) Does Reliability respond promptly and efficiently to unexpected and/or newly detected failure modes?	___	___
	Comments		
	(d) Are line failures (types and causes) analysed and reported to those responsible for corrective actions?	___	___
	(e) Are corrective actions resulting from failure analysis agreed with the Q.A. group involved or Reliability if parts or process changes must be made?		
	Q.A. Group	___	___
	Reliability	___	___
	Comments		
	(f) Has Reliability right to approve test specifications, data tabulation, parts or process changes?	___	___
	(g) Is there a system for in-process failure analysis?	___	___
	End-item failure?	___	___
	Reporting?	___	___
	Comments		

	YES	NO
(h) Are following items submitted to failure analysis as a matter of routine?		
- Production line rejects	___	___
- Lots with a high rejection rate	___	___
Define:-		
- Items returned by Orderer	___	___
- Items returned by Orderer with special request for failure analysis	___	___
(i) Has Manufacturer a failure analysis laboratory or an equivalent facility?	___	___
Comments		
(j) Are failure analysis procedures:-		
(1) Available?	___	___
(2) In use?	___	___
(3) Adequate?	___	___
Comments		
(k) Is failure analysis equipment:-		
(1) Available?	___	___
(2) In use?	___	___
(3) Adequate?	___	___
Comments		
(l) Are there special personnel for failure analysis?	___	___
Comments		
(m) Are failure analysis reports:-		
(1) Available?	___	___
(2) Adequate?	___	___
Comments		
(n) Has Reliability a programme to ensure reliability of discrete device designs prior to release thereof?	___	___
Comments		

- | | YES | NO |
|--|-----|----|
| (o) Has Reliability access to all pertinent development and production data of discrete devices for analysis purposes?
Comments | — | — |
| (p) Is reliability data available of discrete devices from the line(s) which the Manufacturer wishes to be approved?
Comments | — | — |
| (q) Has Manufacturer an evaluation laboratory for determination of product characteristics? | — | — |
| (r) If Manufacturer has an evaluation laboratory: | | |
| - Does it operate according to an established programme? or | — | — |
| - According to special requests? | — | — |
| Comments | | |
| (s) Give examples of problems investigated by evaluation laboratory | | |
| (t) Are laboratory results available on request? | — | — |
| (u) Are data sheets based on these results? | — | — |

2.8 CONTROL OF PROCUREMENT SOURCES

- | | | |
|--|---|---|
| (a) Has Manufacturer adequate written procedures for purchase control of materials, components and services?
Comments | — | — |
| (b) Has Manufacturer an effective vendor rating system?
Comments | — | — |

	YES	NO
(f) Are accepted materials adequately identified?	___	___
Do documents show evidence of acceptance?	___	___
Comments		
(g) Are rejected materials adequately identified and segregated?	___	___
Comments		
(h) Which materials are subject to limited shelf life limitations?		
Comments		
(i) Are shelf life and cure date materials properly identified and controlled?	___	___
Comments		
(j) Do records indicate traceability of units, lots and sublots to applicable documents (specification, revision letter - if any - and inspection record)?	___	___
Comments		
(k) Are materials stored in a controlled area under the responsibility of an authorised Custodian?	___	___
Comments		
(l) Are suitable inspections and tests, including physical and chemical tests, performed on raw materials?	___	___
Comments		
(m) Are such tests performed:		
- In-house?	___	___
- At other locations?	___	___
Comments		

- | | YES | NO |
|--|------------|------------|
| (c) Does rating system provide for effectiveness of written corrective actions received from Suppliers?
Comments | ___ | ___ |
| (d) Do purchase documents require delivery of test reports if such reports are specified in the relevant ESA contract?
Comments | ___ | ___ |
| (e) Is there a means of channelling information when specification changes require modification of current purchase orders?
Is "Receiving Inspection" notified of changes in purchase orders?
Comments | ___
___ | ___
___ |

2.9 CONTROL OF INCOMING MATERIALS (Performed in situ)

- | | | |
|---|------------|------------|
| (a) Are Manufacturer's written standard inspection procedures adequate for control of incoming materials and services received?
Do inspectors know how and when to apply these procedures?
Comments | ___
___ | ___
___ |
| (b) Are materials received in a controlled area from which removal prior to inspection is impossible?
Comments | ___ | ___ |
| (c) Are materials properly handled and protected during the receiving process?
Comments | ___ | ___ |
| (d) Does Receiving Inspection use drawings and purchase orders?
If so, do these documents show Quality Control review?
Comments | ___
___ | ___
___ |
| (e) Are test reports from Suppliers being reviewed?
Comments | ___ | ___ |

- | | YES | NO |
|---|-----|-----|
| (n) Are storage containers, racks, bins, etc. adequate for type of material stored? | --- | --- |
| Comments | | |
|
 | | |
| (o) Is lot traceability maintained? | --- | --- |
| Comments | | |
|
 | | |
| (p) Is "first in/first out" method applied? | --- | --- |

2.10 IN-PROCESS INSPECTIONS AND TESTS

- | | | |
|--|-----|-----|
| (a) To whom does In-process Q.A. Inspection report? | | |
|
 | | |
| (b) Are inspection and/or operation travellers used sequential to performance and control of all operations and processes? | --- | --- |
| Comments | | |
|
 | | |
| (c) Do travellers refer to inspection procedures? | --- | --- |
| Do inspectors know how and when to use them? | --- | --- |
| Comments | | |
|
 | | |
| (d) Do travellers refer to controlled <u>specifications</u> ? | --- | --- |
| Do specifications show <u>current</u> revision status? | --- | --- |
| Comments | | |
|
 | | |
| (e) Does Q.A. have written in-process procedures to control acceptance of products? | --- | --- |
| Comments | | |
|
 | | |
| (f) Is type and quantity of available inspection equipment adequate for type of work being accomplished? | --- | --- |
| Comments | | |

	YES	NO
(g) Are documentation and instruments used by inspectors subject to calibration control?	___	___
Is calibration evident and up-to-date?	___	___
Comments		
(h) Is there a specific material review procedure?	___	___
Comments		
(i) Do in-process Q.A. inspectors summarise quality experience on the basis of specific process stages?	___	___
Do they issue quality reports on a regular basis?	___	___
Do reports result in assistance and/or action?	___	___
Comments		
(j) Are requests for corrective action issued in writing?	___	___
Are such requests answered?	___	___
Does corrective action ensue?	___	___
Comments		
(k) Does Q.A. maintain any statistic controls (X&R, etc.) in the in-process area?	___	___
Are these controls up-to-date and at individual process stations?	___	___
Comments		
(l) Is lot identification maintained throughout processing?	___	___
Comments		
(m) Are there documents describing in-process manufacturing procedures and controls?	___	___
Comments		

	YES	NO
(n) Are there documents describing in-process inspections?	___	___
Do inspectors know how and when to use them?	___	___
Comments		
(o) Are there specific standards for handling, cleanliness and care of materials, parts and equipment?	___	___
Comments		
(p) Are calibrations evidenced and up-to-date?	___	___
(q) Has Q.A. authority to stop production flow in case of out-of-control conditions?	___	___
Is a written material review procedure in use?	___	___
Comments		
(r) Are records maintained of training and competence of operators for welding, soldering, radiography, radiflo and plating?	___	___
Comments		
(s) Are certified operators identifiable by means of a card or badge on their clothing?	___	___
Comments		

2.11 SURVEY OF MANUFACTURING LINE

This review shall be performed in 2 phases:-

- (1) Identification of the various steps listed in the flow chart to define the corresponding operations and collect all relevant information.
- (2) Actual line survey (indicate if inspection was performed).

If different technologies are applied, the inspection results shall be supplied on separate sheets.

2.11.1 Manufacturing Environment

- (a) Which phases of manufacture are carried out under controlled environmental conditions?

(b) Give details of conditions.

YES NO

2.11.2 Fabrication of Substrate

(a) State type of material used.

(b) State source of material.

(c) State method of optical axis location (if required).

(d) State method of slicing into individual elements.

(e) How are surfaces finished?

(f) How are edges finished?

(g) How are elements cleaned?

Comments

2.11.3 Metallisation of Substrate

(a) Is a SEM available?

	YES	NO
(b) Are controls on photo-resist factors documented?		
(1) Preparation (frequency, chemicals, method)	___	___
(2) Evaluation (specific gravity, viscosity, solids residue, definition of line width, pin holes)	___	___
(3) Storage conditions (temperature, container type)	___	___
(4) Application (mounting on substrate, temperature control, spin rate and duration, acceleration)	___	___
(5) Baking (time, temperature)	___	___
(6) Exposure (contact pressure, light intensity, time)	___	___
(7) Development (time, temperature, inspection, magnification, lighting, rejection criteria)	___	___
(8) Environmental control (temperature, relative humidity, dust count, lighting)	___	___
Comments		
(c) Are controls on masking documented?		
(1) Environmental conditions (relative humidity, temperature, dust count)	___	___
(2) Geometry (width, length, spacing)	___	___
(3) Pin holes (size, density)	___	___
(4) Scratches	___	___
(5) Foreign body contamination	___	___
(6) Edge integrity	___	___
(7) Storage	___	___
(8) Inspection method	___	___
Comments		
(d) Method of metallisation?		
(1) Vacuum chamber deposition	___	___
(2) Electron beam deposition	___	___
(3) Sputtering	___	___
(4) Other	___	___



	YES	NO
(e) Metal adhesion test capability?	—	—
Comments		
(f) Electrical tests on metallisation.		
(1) Are procedures documented?	—	—
(2) Is test equipment available?	—	—
(3) Are personnel trained?	—	—
Comments		

2.11.4 Bonding of Substrate

(a) What material is used?		
(b) How is bond thickness controlled?		
(c) What bonding strength test is employed?		
(d) Documentation control of bonding:-		
(1) Temperature	—	—
(2) Material preparation	—	—
(3) Application technique	—	—
(4) Cleanliness	—	—
(5) Ambient conditions	—	—
Comments		

2.11.5 Lead-bonding YES NO

(a) What material is used?

(b) What lead bond strength test is employed?

(c) Documentation control of bonding:-

- | | | |
|---|-----|-----|
| (1) Temperature | --- | --- |
| (2) Pressure | --- | --- |
| (3) Dwell time | --- | --- |
| (4) Condition of capillary or electrode control | --- | --- |
| (5) Ultrasonic power (if applicable) | --- | --- |
| (6) Ambient conditions | --- | --- |
| Comments | | |

2.11.6 Pre-seal Device Preparation

(a) Are devices cleaned prior to sealing? --- ---

(b) Are devices inspected (100% or less)? --- ---

(c) What internal visual inspection is performed?

(d) What protection is given to cleaned devices?

(e) What segregation of rejected devices is made?

(f) What is final disposition of rejected devices?

Comments

	YES	NO
2.11.7 <u>Enclosure of Devices</u>		
(a) What sealing technique is employed?		
(b) Documentation of sealing process (as applicable):-		
(1) Pre-seal bake (time, temperature, ambient conditions)	___	___
(2) Heat (or power) to produce seal	___	___
(3) Humidity during seal	___	___
(4) Gas-purging of sealed atmosphere (type, pressure, flow rate)	___	___
(5) Welding controls (pressure, power, time)	___	___
(c) Type of leak test (fine, gross)		
(d) Facilities for radiographic inspection?	___	___
Comments		
2.11.8 <u>Visual Inspection - General</u>		
Following relevant operations in Para's. 2.11.1 to 2.11.7:-		
(a) Are visual aids and criteria provided for inspection purposes?	___	___
If so, state for which operations:		
(b) Are visual aids applied to the production line?	___	___
If so, state for which operations:		
(c) Are visual aids and criteria adequate?	___	___
2.11.9 <u>Final Test Area and Screening Facility</u>		
(a) Are they separate operations?	___	___
(b) Are final production tests (see ESA/SCC specification) performed by personnel under Q.A. monitoring? or	___	___
Are they performed by Q.A. personnel?	___	___
Comments		

	YES	NO
(c) Are written inspection and test procedures for product classes on the line available for the final test (Q.A.)?	___	___
Comments		
(d) Do inspectors use assigned stamps to indicate inspection status on materials and accompanying documents?	___	___
Comments		
(e) Are requests for corrective action made in writing?	___	___
Are such requests answered?	___	___
Comments		
(f) Are rejected devices identified and segregated in a controlled area?	___	___
Comments		
(g) Are records of accepted and rejected material maintained?	___	___
Are these records identifiable with such materials?	___	___
Comments		
(h) Are device failures analysed?	___	___
Are device failure analyses summarised and reported by final Q.A.?	___	___
Comments		
(i) Is a summary inspection and test report sent regularly to quality management (lot acceptance, percentage of defects, types of failure)?	___	___
Comments		

	YES	NO
(j) Is a testing laboratory or equivalent facility available for quality assurance purposes?	___	___
Which of the following tests are performed in the laboratory or facility?		
(1) Electrical tests	___	___
(2) Mechanical tests	___	___
(3) Chemical tests	___	___
Comments		
(k) Are statistical controls of device parameter distribution maintained?	___	___
Are they reported to Q.A. or Reliability?	___	___
Comments		
(l) Is an environmental test facility maintained in-house?	___	___
If not, state where:		
Are the following tests performed at this facility?		
(1) Temperature (high, low, cycle)	___	___
(2) Shock (mechanical, thermal)	___	___
(3) Acceleration	___	___
(4) Vibration (fixed, variable)	___	___
(5) Moisture resistance	___	___
(6) Altitude	___	___
(7) Radiographic	___	___
(8) Hermeticity tests	___	___
(a) Fine leak, if applicable	___	___
(b) Gross leak or penetrant dye	___	___
(9) Lead fatigue	___	___
(10) Life tests - operating	___	___
Comments		



	YES	NO
(m) Is available equipment used:		
- For production?	___	___
- In R&D?	___	___
- For Quality Control on a sample basis?	___	___
- For screening?	___	___
(n) Are charts provided for the monitoring of environmental test equipment?	___	___
Comments		
(o) Is test equipment adequate for fulfilment of specification requirements?	___	___
Comments		
(p) Is final external visual inspection performed on 100% of the devices?	___	___
Comments		
(q) Are devices stored in a limited access area?	___	___
Comments		
(r) Are devices adequately identified to Customer requirements?	___	___
Comments		
(s) Are there provisions for lot identification?	___	___
Comments		



YES NO

(t) How many burn-in positions are available:

- At room ambient temperature?

- At specified ambient temperature?

- At specified case temperature (cooled hot plate)?

(u) Does burn-in require soldering of leads?

Comments

(v) What precautions are taken to maintain solderability of leads after burn-in?

Comments

(w) How does Manufacturer ensure that failed devices are separated from processed lots of:

- SCC Level 'B'

- SCC Level 'C'

(x) Has Manufacturer all test equipment necessary to perform all qualification tests:



- In-house?
- In nearby facility?

Specify equipment and its location:

- In remote location

Specify equipment and its location:

		YES	NO
2.12	<u>PRESERVATION, PACKING AND SHIPPING</u>		
	(a) Are there adequate written procedures for control of shipping?	___	___
	Comments		
	(b) Are materials designated for shipment properly identified, handled and protected?	___	___
	Comments		
	(c) Do copies of Customer's purchase order and evidence of inspection acceptance accompany materials from end of final test up to the time of shipment?	___	___
	Comments		
	(d) Do Q.A. personnel perform audits of all outgoing lots?	___	___
	Comments		
	(e) Do shipping documents reflect inspection status or evidence of inspection, identification and similar shipping requirements?	___	___
	Comments		
	(f) Does Manufacturer verify conformity of devices and invoices with purchase order?	___	___
	Comments		
	(g) Does Manufacturer implement special packaging methods for hi-rel devices?	___	___
	If so, which of following methods is used?		
	- Individual packages	___	___
	- Mechanical protection	___	___
	- Environmental protection	___	___
	- Special warning labels	___	___
	(h) Is shipping method designed to allow official inspection by Customs without actual removal of protective material?	___	___
	Comments		

 	<p>ESA/SCC Basic Specification No. 2023502</p>		<p>PAGE 30 ISSUE 1</p>
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(i) Do instructions prohibit the use of substandard packaging methods for shipment of hi-rel devices?	YES —	NO —
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2.13 SUMMARY OF INSPECTION RESULTS

Indicate inspection results per manufacturing and testing area, whereby:

- V = Adequate.
- O = Insufficient or non-adequate.
- = Not checked.
- N/A = Not applicable.

1 2 3 4 5 6 7 8 9 10

Environmental conditions:


- Cleanliness
- Temperature control
- Humidity control
- Occupancy

Procedures available:

- Travellers
 - Calibration
 - Segregation of rejects
 - Inspection evidence
-

Area No.

- 1 =
- 2 =
- 3 =
- 4 =
- 5 =
- 6 =
- 7 =
- 8 =
- 9 =
- 10 =

	ESA/SCC Basic Specification No. 2023502		PAGE 32 ISSUE 1
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2.14 GENERAL OBSERVATIONS (Not to exceed 2 pages)