**GE-AVIATION JOINT AFFILIATES SUPPLIER REQUIREMENTS FOR CHARACTERISTIC ACCOUNTABILITY, VERIFICATION AND QUALITY PLANNING**

**Specification Number: S-SPEC-5**

**aeDMS #: S-1007**

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*This specification is in addition to and in no way limiting, superseding, or abrogating any contractual obligation as required by the applicable procurement document.*

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**Applicability:**

The applicable portions of this specification apply to suppliers of the following GE Aviation Joint Affiliates entities when flowed down via purchase order or long term agreement:

* GE Aviation Systems LLC
* GE Aviation Systems Ltd
* Unison Engine Components Inc.
* Unison Engine Components Ltd
* Unison Industries, LLC
* Middle River Aircraft Systems

1. REFERENCED DOCUMENTS

AS9102 Aerospace First Article Inspection Requirements

S-1005 GE-AJA Supplier Quality System Requirements

GT1005-2 Source Problem Report

GT1005-3 Supplier Nonconformance Material Report

GT1007-1 Part Number Accountability

GT1007-2 Product Accountability: Raw Material, Special Process, Functional Testing

GT1007-3 Characteristic Accountability, Verification and Compatibility Evaluation

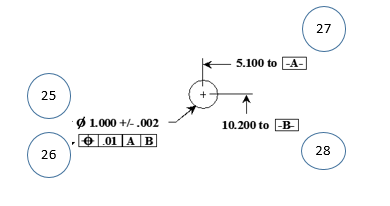
GT1007-4 Product Audit Part Family Designation

GT1007-5 Product Audit Proposal/Completion

GT1007-6 Product Validation Checklist

1. PURPOSE
2. To ensure that all GE Aviation Joint Affiliates (hereinafter referred to as GE-AJA accountable characteristics of a product are addressed by the supplier in the manufacturing and quality plans and that planning includes controls adequate to ensure continued conformance of these characteristics.
3. To provide requirements for documenting the results of First Article Inspection (FAI) and evaluations of changes after initial FAI documentation.
4. APPLICABILITY AND USE
5. This specification defines characteristic accountability and verification (CAV) and quality planning for initial approval and ongoing production. This applies to product supplied to GE-AJA by GE-AJA Suppliers (reference Appendix E for FAI applicability guidance),
   1. Unless otherwise specified in the purchase order, this specification does not apply to:
      1. Procured standard catalog items, COTS or deliverable software
      2. Unique single run production orders, not intended for ongoing product (for example, out of production spares)
      3. Development and prototype parts that are not considered part of the first production run
6. The supplier is responsible for performing characteristic accountability and verification and development of the quality plan in accordance with this specification. This shall be completed and GE-AJA QR approved before shipment of production hardware. GE-AJA reserves the right to witness the supplier’s inspections and/or tests to determine the degree of conformance.
7. FAI documentation shall contain the elements of forms GT1007-1, GT1007-2 and GT1007-3. Any exceptions shall be approved by the GE-AJA Quality Representative (GE-AJA QR). FAI documentation (and supporting elements as described in paragraph F) shall be considered a quality record and shall be retained in accordance with S-1005. All forms shall be completed either electronically or in permanent ink. All forms and supporting FAI documentation shall be in English (native/non-English language may be included). In the event of any inconsistency between the supplier’s native language and translations to or from English, the English language meaning shall control.
8. This specification applies to GE-AJA drawings and specifications for all levels of parts within an assembly, assemblies, sub-assemblies, and detail parts, including castings and forgings, modifications to standard catalog or Commercial-Off-the-Shelf (COTS) items, and to all Suppliers who are responsible for producing accountable characteristics of the product. Suppliers who receive the Purchase Order/Contract from GE-AJA are responsible for flow down of the applicable requirements of the latest issue of this specification to their sub-tier suppliers. If the FAI is completed by an independent 3rd party, the GE-AJA supplier remains fully responsible for the completeness and accuracy of the FAI.
9. In the event of conflict, requirements in other sourcing specifications or quality documents referenced on the Purchase Contract take precedence over the requirements in this document.
10. Data shall be recorded in the Units of Measure specified on the drawing.
11. The Drawing Revision entry on form GT1007-1 is the revision applicable to the First Article part shipped. GT1007-2 and GT1007-3 do not need to be changed or submitted if a drawing revision does not affect accountable characteristics reported on those forms.
12. For any part or assembly with a lapse in manufacturing of 24 months or more (between the completion of the manufacturing cycle of the last part produced and the start of a new part), a new full FAI shall be completed per this specification. The FAI shall be approved by the GE-AJA QR prior to shipment
13. For any part or assembly with a lapse in manufacturing of less than 24 months that are ordered to a revision other than the revision listed on the FAI, a delta/partial FAI shall be completed per this specification to account for all drawing and Part List changes that have occurred. This FAI shall be approved by the GE-AJA QR prior to shipment.
14. The supplier shall ensure secure transmission of FAI data package. Email is not an acceptable transmission method. Transmission via eDistrib is preferred. Link to eDistrib: <https://apps.geae.com/edistrib/index.jsp>
15. ESTABLISHMENT OF ACCOUNTABLE CHARACTERISTICS
16. Suppliers are responsible for all accountable characteristics, including those generated by their sub-tier suppliers (e.g. inspection data, test data, Acceptance Test procedures, etc.). If sub-tier suppliers do not account for their characteristics, the prime supplier is responsible for initiating a separate FAI document or including the characteristics in their FAI document.
17. A ballooned drawing shall be generated and accountable characteristics numbered for the manufactured part. Suppliers are responsible for ballooning the drawing themselves using the guidelines in this Section. The supplier has the ultimate responsibility for ensuring completeness and accuracy of the Characteristic Accountability. Each ballooned characteristic shall be recorded on GT1007-3.
18. Characteristic numbers shall be assigned to each of the following drawing features (unless otherwise required or specified by GE-AJA QR.
    1. Dimensional features, with the following notes:
       1. Reference dimensions are not considered accountable characteristics and need not be ballooned.

ii. Basic dimensions should be referenced in relation to the respective measurable characteristic. (See Figure 1 below for an example).



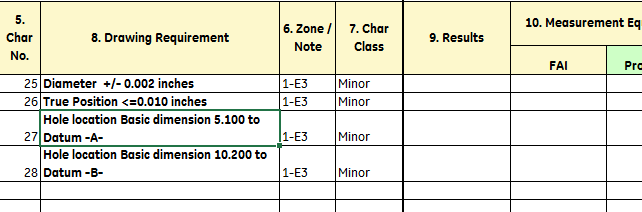


Figure 1 (for reference only)

* 1. Specifications
     1. Accountable characteristics produced by specification definition/requirements defined on the drawing shall be identified and listed for characteristic accountability and first article inspection on GT1007-3. However, the supplier shall evaluate any specifications that are referenced within those specifications listed on the drawing for additional accountable characteristics.
     2. The supplier is responsible for verifying the revision, accuracy, completeness for characteristic breakout of the specification as they apply to their FAI.
     3. Unless otherwise approved by GE-AJA QR, specification characteristics and results shall be included as part of the FAI package.
     4. Accountability requirements regarding characteristics identified within process/special process specifications (for example, processing parameters such as heat treat temperature) will be defined by the GE-AJA QR.
  2. Drawing Notes

1. Measurable features within notes.
2. Non-measurable characteristics within notes.
3. Sequence of Operations.
4. Test/functional parameters. If a Test Procedure is defined, the document number/note will be ballooned and must be included on GT1007-2.
   1. Tables (on drawing) - Except as follows
5. Nondestructive Evaluation (NDE) tables can be ballooned as one characteristic unless otherwise requested by the GE-AJA QR.
6. Configuration control tables – Only cells associated with the part number being reported shall be ballooned.
7. Information tables such as datum tables need not be ballooned.
   1. Supplementary views, tabulated features and alternate methods of manufacture views
8. When required for the part number being reported, all characteristics shall be ballooned.
9. When not required for the part number being reported, a single characteristic may be used.
   1. Find numbers (i.e. item numbers) called out on the drawing (in the notes and/or the body of the drawing)
10. When working to an issued drawing plus Changes in Design (CID), the CID number(s) shall be listed on GT1007-1.
11. When design requirements are in a DPD format and drawing information is not available or provided in the GE-AJA 2D drawing, the supplier shall establish a process to extract the accountable characteristics and extract, verify and include the DPD accountable characteristics in the CAV/FAI.
12. FIRST ARTICLE INSPECTION

The supplier shall have a process to plan for FAI activities prior to the first production run and activities to be performed through the FAI process. Those responsible for the FAI activities shall be identified. The First Article Inspection part or parts shall be representative of a production run.

1. When possible, the First Article Inspection shall be performed using an independent gaging method rather than the normal product acceptance plan. Production gaging and test equipment may be used for first article inspection in cases where it is the only method of accurately checking a characteristic.
2. Inaccessible characteristics: A characteristic inaccessible at final inspection may receive first article inspection when accessible during the process in lieu of disassembly/destruction. The supplier shall ensure that subsequent processing does not cause the characteristic to become nonconforming or unintentionally alter the characteristic.
3. Characteristics that cannot be evaluated non-destructively on a finished part may be re-evaluated using component parts prior to final assembly or by using hardware not useable because of reasons unrelated to the characteristic being re-evaluated.
4. Non-measurable characteristics- results such as or “Conform”, “Verified” or “For Information Only” shall be entered.
5. This paragraph is no longer used.
6. Multiple characteristics (e.g. bolt circle, dovetail slots, radii): Provide variable data for all occurrences of every characteristic or minimum/maximum readings along with the number of measurements taken unless additional data is required by the GE-AJA QR.
7. Continuous characteristics (e.g. radius along circumference, weld seams, edge breaks, surface finish, slot dimension, wall thickness and continuous features invoked by specifications): Measure a sufficient number of locations over the total extent of the characteristic to ensure total conformity. Provide variable data for all measurements or minimum/maximum readings along with the number of measurements taken unless additional data is required by the GE-AJA QR.
8. Characteristics that are identified as non-conforming during first article inspection shall be managed as follows:

a. The FAI shall be marked as “Not Complete”

b. The non-conformance shall be documented in accordance with S-1005 and the non-conformance document number shall be referenced in the GT1007-3 results.

c. Any nonconforming characteristic found on the FAI shall be subject to 100%

characteristic evaluation until justification for an alternate acceptance plan per Appendix A is approved by the GE-AJA QR.

d. Corrective action(s) shall be implemented and a partial FAI performed for all affected characteristics on the next production run, after implementation of the associated corrective action(s). If the partial FAI does not clear all identified non-conformances, the FAI is still “Not Complete” and the requirement to complete the FAI is still in effect. NOTE: A full FAI may be done in lieu of a partial FAI

1. The supplier shall:
   1. Review Manufacturing process documentation (e.g. routing sheets, work instructions, etc.) to ensure all operations are complete as planned
   2. Review Material certifications for compliance
   3. Verify approved Special Process sources are used, if applicable, and the correct specification(s) are documented
   4. Verify that Key Characteristic requirements have been met, as applicable
   5. Verify part specific gages and/or tooling is qualified and traceable, as applicable
   6. Verify all design characteristic requirements are accounted for, are uniquely identified and have inspection results traceable to each unique identifier
   7. Review completed GE-AJA approved test procedure data sheets, if applicable
   8. Verify test equipment (including any Special Test Equipment i.e. STE), if applicable, has been identified and if required, approved by GE-AJA QR.
2. SUPPORTING ELEMENTS OF FAI/CAV

Items in this paragraph are required supporting elements of the FAI Data Package unless otherwise directed by the GE-AJA QR (reference paragraph C.3).

1. Photo of part marking (if required by GE-AJA QR)
2. Manufacturing routing sheets
3. Completed GT1005-3 Supplier Nonconforming Material Report approved by GE-AJA QR as applicable
4. Completed GT1005-2 Source Problem Report (SPR) approved by GE-AJA QR for interpretation/specification options
5. Certificates of conformance for all parts, material, tests and sub-tier special processes. Certificates shall be traceable to the documented information on GT1007-2 or GT1007-3 forms
6. Ballooned drawing
7. Special Process Technical Plans (as required by specification)
8. Inspection Reports (if applicable)
9. Completed Test Procedure Data Sheets, if applicable
10. Completed FAI Checklist
11. QUALITY PLANNING AND ACCEPTANCE PLANS
12. Manufacturing and quality planning shall be in place before final acceptance of deliverable hardware to ensure that all accountable characteristics are included in the plans. See [Appendix A](#_Appendix_A:_) for a recommended quality planning process map. Where supplier/sub-tier product and process drawings exist that contain GE-AJA accountable/design characteristics, the supplier shall complete a compatibility assessment. Reference paragraph J.
13. The adequacy of the measurement system shall be considered when selecting inspection equipment for first article inspection and ongoing production. See [Appendix B](#_Appendix_B:_) for a recommended measurement and test equipment selection process.
14. The required acceptance plan is 100% inspection of each characteristic on every piece manufactured, except when implementing a Product Acceptance Plan per [Appendix A](#_Appendix_A:_), Table 1.
    1. Documented justification for less than 100% inspection is required and shall be available upon request by the GE-AJA QR. The documented justification shall be referenced in the FAI.
15. Data utilized for process capability calculations shall be representative of the planned process and shall not include rework or work outside the normal process. Statistical methods defined in [Appendix C](#_Appendix_C:_) are the preferred methods to use for Cpk calculations.
16. The operation where each characteristic is verified shall be recorded on the FAI. Consideration should be given as to whether subsequent operations could affect the final characteristic.
17. Measurement of characteristics for product acceptance, whether completed during manufacturing or at final inspection, shall be performed by qualified inspectors or certified operators. Certification of operators shall be achieved through a supplier certification program that meets the requirements of [Appendix D](#_Appendix_D:_).
18. Measurement of characteristics for product acceptance shall be performed using measurement and test equipment meeting the requirements of S-1005 (Calibration).

1. CHANGE MANAGEMENT

All requirements of this specification apply to accountable characteristics impacted by any of the changes listed below including those invoked by drawing specifications. The supplier shall notify the GE-AJA QR when a change occurs (as defined in a-e below) to determine if a full/partial FAI must be submitted to and approved by GE AJA. The FAI may require additional characteristic accountability as deemed necessary by the change. When the submission of a full/partial FAI is not required, the supplier shall conduct an internal full/partial FAI.

Changes to a configuration of a previously approved part (i.e. -001 to -002, P01 to P02, G01 to G02): Note, ALL changes; additions, deletions, and modifications of characteristics shall be accounted for and submitted to GE-AJA QR for approval.

Drawing or specification changes that do not change the part or assembly number. Note, ALL changes; additions, deletions, and modifications of characteristics shall be accounted for.

Process changes (including sub tier changes): Inspection method and/or frequency for affected characteristics of any process change shall be evaluated for impact.

Product Acceptance Change:

Change in product acceptance plan. (see [Appendix A](#app_A))

Change in production inspection equipment (e.g. from micrometer to functional gage).

Changes to the point of inspection relative to the manufacturing process (e.g. move from final inspection to in-process).

A repeat (full or partial) [first article inspection](#FAI) shall be considered when any of the following events occur that could affect part characteristics: (This paragraph highlights some repeat [FAI](#FAI) scenarios.)

A change in inspection methods or measurement equipment. Note: Refer to [Appendix B](#app_B), “Measurement and Test Equipment”.

Relocation of a process and equipment within a [Source](#Source) (for example, changing a printed circuit board assembly from one line to another; moving a reflow oven to another area/section of the building)

A change to numerical control programs. Note: Refer to S-1005, “Requirements For Suppliers Software Quality Assurance Programs”.

A natural or man-made event that adversely affects characteristics.

Any change in the process or process sequence (examples: tooling, fixtures) or material that could potentially affect part characteristics.

The most current approved quality plan shall match the inspection method and frequency being used at the supplier and/or sub-tier suppliers. Supplier shall develop a plan to update FAI when manufacturing plans or inspection plans are revised.

Supplier shall have a process to ensure all engineering and manufacturing changes to the manufacturing planning are reviewed against the current quality plan. Changes shall be submitted as required.

1. PRODUCT/PROCESS AUDIT

Supplier Product/Process Audit: This section defines the minimum requirements of product audits. These requirements shall be incorporated into a Supplier Product Audit procedure.

1. The purpose of this audit is to verify that the established process controls and product acceptance plans continue to provide conforming material.
2. This audit is a full FAI for a fully processed production part. It is also an evaluation of the supplier’s planning and procedures to ensure compliance with the requirements of this specification. Evaluation shall include variable results, inspection equipment, and the current acceptance plan per [Appendix A](#_Appendix_A:_). Use of GT1007-6 (Product Validation Checklist) is encouraged.
3. Whenever possible, the re-evaluation shall be performed using a method of acceptance measurement independent of the planned acceptance measurement method. In cases where the production method of acceptance is the most accurate (e.g. CMM), it may be used as long as program verification or an independent check is completed. Characteristics that cannot be evaluated non-destructively on a finished part may be re-evaluated using component parts prior to final assembly or hardware that is not useable because of reasons unrelated to the characteristic being re-evaluated.
4. The supplier shall handle Product Audit non-conforming findings per the current revision of S-1005, Quality Systems Requirements.
5. Any finding during product audit will subject the supplier to additional audits at the GE-AJA QR’s discretion.
6. Product Audit Family designations and parts assigned shall be submitted to the GE-AJA QR for review and approval using form GT1007-4 (Product Audit Part Family Designation). Product audit Planning and Family designations will be reported to GE-AJA QR through the Support Central Workflow on an annual basis. If the Support Central Workflow cannot be used by the supplier, GT1007-4 must be submitted to the GE-AJA QR at the beginning of the audit year. NOTE: Families defined either too broadly or too restrictively can defeat the purpose of a product audit, which is to evaluate effectiveness of the processes used to manufacture parts.
7. A minimum of one part per part family shall be audited annually. The supplier part selection for audits shall be reviewed and approved by the GE-AJA QR through the workflow. Any changes to the plan shall be approved by the GE-AJA QR. Once the audits are complete, audit completion information shall be reported through the workflow. If the workflow cannot be used by the supplier, GT1007-5 must be submitted to the GE-AJA QR by the end of the audit year or when all product audits are complete.
8. Parts that have been audited should not be re-audited until all parts in the family have been completed. (Exception may be made for parts with quality issues, high volume parts or for parts not in production when the audit is performed).
9. The supplier shall retain Product Audit documentation including the completed FAI package. If the workflow was not used by the supplier, the Product Audit documentation must also include forms GT1007-4 and GT1007-5.
10. Exception to Product/Process Audit Requirements: 100% lot-by-lot testing performed by a certified Test Laboratory may satisfy the requirements of the product audit. This applies to raw material and to processes that generate a certification of conformance for every manufacturing lot. Processes that are verified by a certified Test Laboratory but do not get a certification for every manufacturing lot (e.g. EDM, Laser, Heat Treat) require a re-certification and shall meet the requirements of this specification.
11. SUPPLIER DESIGNED PRODUCTS

All requirements of this specification apply to the characteristics defined by the GE-AJA or GE-AJA Customer drawings unless specifically noted below.

Where supplier/sub-tier product and process drawings exist that contain GE-AJA accountable characteristics, the supplier shall complete a compatibility evaluation.

Product design specification documentation shall be completed only on the following:

Acceptance/Inspection Tests defined in the Quality Assurance Provisions section of the design specification.

Identification and Part Marking requirements of the design specification or specifications referenced therein.

As a minimum, the Supplier's system shall assure that characteristics defined by the Supplier/Sub-tier drawings are accounted for, documented, and controlled. The format(s) shall be defined by the Supplier and may be subject to review and disapproval by GE-AJA Quality Representative. The system shall include the following:

The documented format(s), defined by the Supplier, shall include the same elements as shown on form GT1007-3 under the headings; Characteristic Accountability, Inspection/Test Results, and Product Acceptance.

Changes to supplier's or sub-tier’s drawing, manufacturing or quality plan shall be documented and approved under requirements defined by the supplier system for their characteristics.

First Article Inspection Package (FAI) Requirements for characteristics defined by GE-AJA Supplier/Sub-tier drawing(s).

GE-AJA Drawing(s) and Characteristics: FAI package shall include all items required by Section F and the following items:

Results from product acceptance test and inspection requirements.

Evidence of GE-AJA engineering approval of applicable Test Procedures.

Supplier/Sub-tier Drawing(s) and Characteristics: First Article Inspection package shall include the following items that are to be retained at the Supplier facility unless otherwise directed by the procurement document:

First Article Inspection results.

Nonconformance document(s) referenced for accepting nonconforming characteristic(s).

Referenced exhibits, e.g. functional test reports, evidence of part marking, certifications, etc.

NOTE: A copy of all GE-AJA approved Test Procedures shall accompany first article data.

1. REMOVED

**DEFINITIONS (See SAE AS9102 and S-1005 for additional definitions)**

ACCOUNTABLE CHARACTERISTICS (equivalent to Design Characteristic as defined in AS9102): Those dimensional, visual, functional, mechanical, and material features or properties, which describe and constitute the engineering definition of the article and can be measured, inspected, tested, or verified to determine conformance to the engineering definition or Digital Product Definition (DPD) requirements. Dimensional features shall include those features defined by the engineering definition such as target-machined (or forged/cast) dimensions on forgings, castings, and weld/braze joint preparation necessary for acceptance of finished joint. Material features or properties shall include processing variables and sequences that are specified by the engineering definition (e.g., heat treat temperature, fluorescent penetrant class, ultrasonic scans, sequence of welding, heat treat, etc.).

Accuracy Ratio: The ratio between the total M&TE (Measurement & Test Equipment) Accuracy and the total part tolerance.

Calibration Tolerance: Total permissible variation or limits allowed for calibration of M&TE (Measuring and Test Equipment).

CERTIFIED OPERATOR – An operator who has fulfilled all the qualifications, training and testing requirements for their assigned job description per the supplier OAP (Operator Acceptance Plan). Certified Operators may verify characteristics which are inspected at the point of generation. See [Appendix D.](#_Appendix_D:_)

CERTIFIED TEST LABORATORY- a GE-AJA approved independent test laboratory facility.

Characteristic Tolerance: Difference between upper and lower limits of a part characteristic.

CHANGE IN DESIGN (CID)- an engineering document which changes the content of the product definition. Could also be known as NOR (Notice of Revision), ECO (Engineering Change Order, RN (Revision Notice) or other names.

COMPATIBILITY EVALUATION: An evaluation of supplier/sub-tier product and process drawings containing GE-AJA engineering definition, to ensure that they specify the same engineering definition as the GE-AJA engineering definition.

CORRELATION: A characteristic that has been verified by an operator is re-verified by a different operator/inspector using the same gage type and results are equivalent within acceptable tolerance band.

CUSTOMER: The term customer, as used in this procedure, can mean external end users or internal customers.

DIGITAL PRODUCT DEFINITION (DPD) REQUIREMENTS: requirements of any digital data files that disclose, directly or by reference, the physical or functional design requirements. Reference AS9102 for additional information on DPD.

ENGINEERING DEFINITION: Design engineering requirements as documented within the drawing, drawing notes, specifications on the drawing, or referenced specifications including digital product definition (DPD) requirements, if applicable.

Feasible: Capable of being performed, within constraints (e.g., delivery, cost, technical) as agreed between GE-AJA and the Supplier.

FIND NUMBER: Find number or item number refers to the ordinal number that gives an ID tag to one of the constituents in a parts list (list of materials, bill of materials). Thus "fasten using Find Number 7 (or item number 7)" refers to a fastener that is listed as number 7 in the parts list or bill of material (BOM).

FIRST ARTICLE INSPECTION (FAI): A complete, independent, and documented physical and functional inspection process to verify that prescribed production methods have produced an acceptable item as specified by engineering drawings, DPD, planning, purchase contract, engineering specifications, and/or other applicable design documents

FIRST ARTICLE INSPECTION (FAI) REPORT: The forms and package of documentation for a part number or assembly, including FAI results, per this specification

GE AVIATION JOINT AFFILIATES QUALITY REPRESENTATIVE (GE-AJA QR): A GE employee or authorized representative with the authority to represent GE-AJA Sourcing Quality.

INSPECTOR – An individual who inspects and verifies characteristics, but does not generate the characteristics.

INTERPRETATION OR SPECIFICATION OPTION: A documented process by the manufacturing source to submit a request for a drawing interpretation, specification interpretation, selection of a specification option, or report a possible drawing error, or a producibility proposal.

Key Characteristic (kc): The select few, measurable features of a specific part/drawing/specification/process where variation can significantly impact customer satisfaction, manufacturability, durability or performance.

Measuring and Test Equipment (M&TE): All devices used to measure, gage, test, inspect or otherwise examine items to determine compliance with drawing or specification requirements.

Measurement System Analysis (msa): Method to define and document the amount of variation in the process due to the measurement system. It is a tool that evaluates the measurement system’s performance on specific characteristics in the process and under conditions that occur in the process.

Nonconformance Document: A document used for disposition of nonconforming characteristics, entered into a Nonconformance Report (NCR) System or similar system.

OPERATOR ACCEPTANCE PLAN (OAP): Supplier plan which defines the requirements, procedures and individual responsibilities for the certification of operators. See [Appendix D](#_Appendix_D:_)

OPERATOR – The individuals who physically perform the process. These individuals can be referred to as ‘Individual Process Owners’, ‘Technicians’, ‘Process Team Members’, or by other terminology suitable for the organization’s program focus and cultural and customer environment.

PART FAMILY – A group of parts with similar processes, materials, complex form, and tolerances, which have been produced by similar manufacturing methods.

Process Capability: The performance of which a process is capable, with all the effects of assignable cause variation removed. Process capability is typically quantified as + or – 3 standard deviations about the process mean.

Process Stability: A process that is operating with only chance causes of variation present is said to be statistically stable.

PRODUCT ACCEPTANCE: Verification that characteristics of a part meet the engineering definition.

Product/PROCESS Audit: Evaluation of any or all accountable characteristics for conformance, independent of Product Acceptance evaluation. Also includes an appraisal of the Supplier’s system to ensure stable processes are in place that continually generates conforming characteristics.

PURCHASE CONTRACT: Purchase Order, Purchase Agreement or other Purchase document

Sigma Value: A statistical measurement, indicating the probability of producing a part characteristic within the drawing limits. The sigma value represents “Z”, the number of process standard deviations between the process mean and the nearest specification limit.

Single Purpose M&TE/Gage: Gage which is designed to accommodate specific part configurations (e.g. airfoil guillotine gages).

SOURCE CHANGE: A change in manufacturing source or the addition of an alternate manufacturing source for a complete part.

SPECIAL PROCESS TECH PLANS: A technical plan is a part-specific, process-specific document required by a Drawing Specification, used to demonstrate source capability to meet the technical requirements of the special process.

Standard M&TE/GagE: M&TE that is not controlled by a tool drawing, i.e., commercially available.

STATISTICAL CONTROL: A quantitative condition which describes a process that is free of assignable/special causes of variation, e.g., variation in the central tendency and variance. Such a condition is most often evidenced on a control chart.

Supplier: Sources (including distributors, warehouses,) other than GE-AJA, who supply material, parts, processes, or services for incorporation into GE-AJA products.

TEST PROCEDURE: Documented procedure describing the functional test methodologies, environmental conditions, equipment, specified values and tolerances.

VERIFICATION – Confirmation through objective evidence that specified requirements have been fulfilled.

# Appendix A: QUALITY PLANNING and ACCEPTANCE PLANS

**GENERAL GUIDELINES**

A1. Supplier product engineering and manufacturing should be involved in quality planning. Collaboration of quality, engineering and manufacturing is expected at the following times:

* PO review
* Initial product development
* Initial quality plan development
* Supplier engineering, inspection, process and/or manufacturing changes (including sub-tier changes)
* A change of inspection frequency is being substantiated
* GE-AJA drawing revision
* 24 month lapse in production

A2. Key, Critical and/or Major characteristics should be considered for continued 100% inspection.

A3. It is recommended that accountable characteristics be inspected at the earliest possible step in the manufacturing process if subsequent process steps will not alter the characteristic.

A4. Following is a recommended map for the Quality Planning process followed by recommended checklist items for each step.

A5. Table 1 shall be followed when selecting Acceptance plans.

**Appendix A: Quality Planning Process Map**

**Evaluate Part Drawing**

**Purchase Contract Issued**

**On-going Process Monitoring and Control Plan**

**Validate Quality Plan**

**Produce Part/ Execute Plan**

**Define Verification Plan and Inspection Method**

**Develop Manufacturing Plan**

**Appendix A: Checklist associated with each step of Process**

**Purchase**

**Contract Issued**

Verify PO matches the quote

Review PO, quality requirements, engineering requirements, remarks, and Customer specific requirements.

Verify if the part is new or previously manufactured

Verify current revision of drawing

Identify if any Change in Design (CID) are issued but not included within the drawing.

Obtain engineering parts list

Is Ballooned drawing available

**Evaluate Part Drawing**

Review engineering part list

Review latest revision of drawings, CID’s as applicable

Identify required specifications and verify current revisions

Identify any drawing or manufacturing issues

Identify stack-up concerns

Review part quality history and discuss with GE-AJA QR nonconformance and escape history

For an existing PN, review internal and sub-tier quality history

Review lessons learned for similar parts with similar manufacturing processes

Request engineering models, Mylar etc. as needed

Extract accountable characteristics from DPD

Identify education and training needs for applicable supplier personnel

Submit Interpretation/Specification option as required (based on issues identified)

**Appendix A: Checklist (cont.)**

**Develop Manufacturing**

**Plan**

After receiving responses to Interpretation/Specification option, identify risk abatement plans for open issues. Discuss with GE-AJA QR.

Initiate FAI

Balloon the drawing

Verify ballooned drawing includes all accountable characteristics

Request specifications not available on site (as needed)

Review datum/transfer datum system.

* Does the datum system control movement?
* Is the datum system repeatable?
* Is order of precedence maintained

Identify key manufacturing characteristics

Develop proposed manufacturing plan sequence. Ensure operation sequence complies with engineering drawing

Develop sequence of steps within each operation

Identify operation step where each characteristic is generated

Verify special processes sources are current and approved (whether performed in house or at sub-tier)

Ensure fixture has needed controls: fixture height, size, tolerances, etc. Error proofing should be considered.

Ensure fixture set-up has needed controls. Error proofing should be considered.

**Define Verification Plan and Inspection Method**

Determine where each accountable characteristic is verified. If an accountable characteristic is verified ‘in process’, evaluate the effect of subsequent processing (including manual benching).

Select appropriate acceptance plan for each accountable characteristic: 100% Evaluation, special process, process parameter, variable data charting, die/mold, fixture/tool, software/numeric, & component/accountable characteristic stack-up. (See Table 1)

Enter the acceptance plan for each characteristic into FAI or equivalent.

Evaluate need for MSA on new gaging techniques. Refer to Appendix B

For single purpose gages or functional gages, error-proof the gage and verify the gage meets the engineering requirements.

Develop detailed inspection process sheets, including visual cell techniques

Verify an Operator Acceptance Plan exists if applicable (See Appendix D)

Define and execute necessary training for operators

**Appendix A: Checklist (cont.)**

**Produce Part/ Execute Plan**

Ensure raw material, processes, equipment, and operators are production ready

Verify gaging method meets minimum requirements of Appendix B

Verify selected gage can be used with geometry / fixture combination

Verify gaging method is understood by those performing the inspection.

Ensure each accountable characteristic is verified by an inspector or certified operator (See appendix D)

For accountable characteristics requiring CMM inspection, verify CMM set-up and routines/programs satisfy engineering requirements

Where single purpose or functional gages are used, perform independent inspections of accountable characteristics. Ensure that the gage correlates to the independent inspection.

Apply statistical analysis if applicable

**Validate Quality Plan**

Ensure all accountable characteristics are included in the quality plan

Ensure FAI part is representative of the defined production process

Complete FAI and quality plan

Ensure quality plan is reconciled to final engineering drawing

If required, complete the “Frozen Process” package

Evaluate the manufacturing/inspection process for improvements (evaluation to be done by product engineering, manufacturing, and quality)

Select appropriate acceptance plan for each accountable characteristic: 100% Evaluation, special process, process parameter, variable data charting, die/mold, fixture/tool, software/numeric, & component/accountable characteristic stack-up. (See Table 1)

**On-going Monitoring and Control Plan**

Maintain On-going monitoring for reduced inspection per Table 1

Evaluate quality plan periodically. Correct /update quality plan as required

Update FAI as the process or quality plan changes (including subtier changes)

Execute product audit plan per S-1007

| Table 1 –Acceptance Plans | | | | |
| --- | --- | --- | --- | --- |
| **Product Accept Plan/Description** | | **Application** | **Requirements** | |
| **Initial Approval** | **On-going Monitoring** |
| **1. 100% Evaluation**  Control by 100% EVALUATION OF ACCOUNTABLE CHARACTERISTICS  Measure all occurrences of every accountable characteristic on every piece manufactured. This plan ensures accountable characteristic conformance through direct measurement of the characteristic on all parts to determine the conformance requirements. | Plan required when justification does not exist for another plan. | | No additional approval data is required to justify this plan. | Consider independent verification of the characteristic and the measurement technique.  Consider Error-proofing. |
| **2. Special Process Control/Evaluation**  This plan ensures accountable characteristic conformance through the control of the input of parameters as generated by a special process. | Applies to special processes when the characteristic is entirely controlled by the process. | | Some Processes may require GE –AJA Certifying Agent Approval.  Establish control parameters through specific correlation studies, i.e. part or specimen cut-up, or through historical process knowledge.  If a specimen is used, provide evidence that the specimen, represents the product as processed. | Evaluation may entail lot by lot or periodic testing proposed by the supplier in the justification for less than 100% evaluation.  Plan shall be in compliance with the applicable engineering specification/s.  Lot traceability shall be maintained through the manufacturing cycle. |

| Table 1 –Acceptance Plans | | | |
| --- | --- | --- | --- |
| **Product Accept Plan/Description** | **Application** | **Requirements** | |
| **Initial Approval** | **On-going Monitoring** |
| **3. Process Parameter Control**  This plan ensures characteristic conformance through the control of the input parameters as generated by a non-special process. | Examples (setup, feeds, and speeds)  For example, surface finish on a grinder where feeds and speeds are not software controlled and areas of features that may be inaccessible without destructive evaluation. | Establish control parameters through specific correlation studies, i.e. part or specimen cut-up, on-part evaluation, or through historical process knowledge.  If a specimen is used, provide evidence that the specimen represents the product as processed. | A monitoring plan shall be proposed by the supplier in the justification for less than 100% evaluation.  Periodic evaluation or additional testing may be required. |
| **4. Variable Data Charting/ SPC (e.g. Process Maintenance through Statistical Process Control)**  The output of a process is statistically monitored to ensure characteristic conformance through verification of the process stability. Generally graphical output is used. | This method may be employed when it can be shown that the output from a process is stable and the capability is sufficient. | - Use standard SPC techniques (Appendix C) to establish control limits. Data should include normal variation that is characteristic of routine production such as different operators and work shifts  - Define a data collection and plotting plan that ensures the ability to capture process shifts or other indications of loss of process stability. The classification of characteristics, rate of production, stability and complexity of process and method of control should be considered in selecting the frequency | If the process gives evidence of violating statistical stability, investigation and corrective action shall be performed. 100% evaluation shall be put in place until stability is revalidated.  - Stability measures such as control limits, limits on first/last piece, etc. shall be reevaluated whenever substantive changes are made to the process. When such limits are modified, the associated capability measure shall also be recalculated.  See Appendix C for signs of process drift or instability |
| **5. Die/Mold Control**  This plan ensures accountable characteristic conformance through the control of the geometry and wear factors for the Die/Mold used to generate the characteristic. | **Appropriate** where a relationship exists between the geometry of Die/Mold being used to generate the accountable characteristics and the final product.  **Not appropriate** if there are removable parts on the die/mold for which assembly cannot be error proofed. | Validate the ability of the Die/Mold to generate the characteristic through verification of the characteristic in the first run of the process.  Per process in Appendix C, ensure process is statistically capable of producing characteristics in conformance with Engineering Requirements. | -Verify correct Die/Mold is being used  -For each set-up of operation, first piece verification shall be completed to ensure proper setup.  - Periodically verify pieces and/or the die or mold to identify wear or shifts that could impact part conformity. Interval shall be documented as justification for less than 100% inspection along with Cpk Value.  -Where rework/repair of the Die/Mold affects product conformance, re-verification shall be performed  -When wear of the Die/Mold is a factor, monitoring shall include periodic inspection of the part.  -Visually Inspect the Die/Mold periodically for damage and wear.  **For Sheet Metal Forming:**  In addition  - The last piece of a lot, run, or work shift (whichever occurs first) shall be verified to ensure no change has occurred that would affect conformity.  -Ensure parts are identified to the lot, run or work shift until last piece has passed verification. |
| **6. Fixture/Tool Control**  This plan ensures characteristic conformance through the control of the cutting tool and/or the fixture. | **Appropriate** where a relationship exists between the geometry of Fixture/Tool being used to generate the characteristics and the final product.  **Not appropriate** if the feature is related to datums.  **Not appropriate** if there are removable parts on the fixture for which assembly cannot be error proofed. | Conduct first article inspection for the established cutting tool/fixture combination. Inspection of the cutting tool is not an acceptable alternative to inspecting hardware for FAI.  Planning should include identification of Fixture/Tool including ancillary parts  Per process in Appendix C, ensure process is statistically capable of producing characteristics in conformance with Engineering Requirements.  **Fixture:**  Establish a plan for on-going monitoring. (e.g. periodic calibration of fixture) | Inspect all potentially affected part accountable characteristics after any modifications or rework to the fixture.  Visually check Fixture/Tool for wear, distortion, damage, loose parts, etc. on a periodic basis.  Changes to the Fixture, Tool, or Process require re-verification of capability.  **Cutting Tool:**  Verify first and last characteristic controlled by the Tool/Process.  Verify characteristics on the first piece of a new work shift/operator change.  NOTE: If it is not feasible to inspect actual part features during production (e.g. inaccessible characteristics), inspection of the cutting tool may be an accepted alternative |

| **Table 1 –Acceptance Plans** | | | |
| --- | --- | --- | --- |
| **Product Accept Plan/Description** | **Application** | **Requirements** | |
| **Initial Approval** | **On-going Monitoring** |
| **7. Software/Numerical Control**  (All aspects of Software Control apply per S-1005)  This plan ensures conformance of accountable characteristics through programmed aspects of a machine (i.e., control of the cutter path of a machine tool.) | **Appropriate** for those characteristics that are generated through software/numerical control  **Not appropriate** if the characteristic is affected by fixture/part set up and the fixture set up is not controlled.  **Note**: If operator offset is required, the characteristics affected by the offset shall be verified on the first part produced after the offset adjustment. | Per process in Appendix C, ensure process is statistically capable of producing characteristics in conformance with Engineering Requirements.  Verify and approve the NC program using an independent method.  Assign unique program numbers and list the controlled program in the manufacturing planning.  Identify the characteristics that will be accepted by the NC program.  Establish a plan for on-going monitoring | Once software program has been proven to generate conforming hardware, all changes to the program shall be under revision control.  Whenever the program is revised, process shall be re-qualified in accordance with the Initial Approval. Monitoring may have to be adjusted based on the change being made.  Verify correct setup for each use, including cutting tool/probe.  Periodically verify pieces to identify process shifts that could impact part conformity. |
| 8. Component/Characteristic Stack-Up This plan ensures characteristic conformance through control and verification of engineering characteristics at lower drawing levels such that assembly of the components into the product result in conformance to the next higher-level engineering characteristics. | Plan is employed for the acceptance of characteristics generated by assembly of two or more components | Functional or engineering analysis showing that the higher-level characteristic will meet print given lower level characteristics are sufficiently controlled | Provide for periodic confirmation that the higher-level characteristics are meeting print requirements.  Changes to sub-components, sub-component processes or sub-component control plans require re-evaluation. |

# Appendix B: MEASUREMENT AND TEST EQUIPMENT

B1. The supplier is required to have a system in place to evaluate their measurement and test equipment (M&TE) through Measurement Systems Analysis (MSA). This system shall ensure that the M&TE utilized effectively evaluates part characteristics.

B2. The purpose in conducting a MSA is to estimate the variation in the measurement system. Once quantified, it can determine if the level of variation can be tolerated or if actions must be taken to improve the measurement system or control effects of this variation.

B3. A MSA should be considered for situations such as the following:

B3.1 A characteristic is defined as a key, major or critical on the Engineering Drawing.

B3.2 A major process improvement effort is being initiated.

B3.3 The characteristic being evaluated has failed a product audit due to gage concern.

B3.4 The characteristic has experienced an inspection escape, such as a delivered nonconformance to an internal or external customer.

B3.5 The known process capability (+/- 3 sigma spread) exceeds the engineering requirement.

B3.6 Recent changes in gage design, measurement method or measurement personnel.

B4. The M&TE accuracy ratio for single purpose measurement equipment for effective characteristic evaluation is minimally 10:1. The M&TE accuracy ratio for standard measurement equipment is minimally 4:1. If the MSA or other knowledge of a gage’s accuracy ratio proves this equipment does not meet this ratio, action is required. To assure characteristic conformance, typical actions could entail (particularly for part characteristics near a tolerance limit):

B4.1 Use of a different, more accurate type of gage.

B4.2 Reducing the part characteristic acceptance limits to be tighter than the drawing tolerance.

B4.3 Conducting redundant evaluation of the characteristic with an alternate means of inspection.

B4.4 Modification of the part manufacturing process to reduce the characteristic variability.

B4.5 Analysis of the results of inadequate or questionable MSA studies to determine root cause and action to take. NOTE: Typical root causes are wrong gage for task, operator not following planning, inadequate planning, work area not suitable, material relaxes or changes due to environment, method changes when shift changes, etc.

B5. Exceptions to paragraph B4 requirements must be supported by data and/or studies to assure effective control of production parts. The GE-AJA QR must be notified if gaging cannot meet paragraph B4 requirements above.

B6. It is recommended that equipment used for measurement purposes be of sufficient accuracy to measure one decimal point beyond Engineering requirements (i.e. if the drawing requires 3 decimal points (.000), the equipment should be capable of reading to 4 decimal points (.0000)). All significant digits beyond the required accuracy capability should be truncated.

B7. Other Recommendations:

B7.1 An accuracy ratio of 10:1 is recommended for key, major and critical characteristics.

B7.2 Direct measurements are preferable to calculated measurements.

B7.3 Use radius gages for radii that are classified as minor only.

B7.4 A No-Go plug or pin should not be used for detecting over maximum conditions for characteristics generated by processes that may produce elongated holes.

# Appendix C: Preferred Statistical Methods

NOTE: Throughout this document, any reference to Cp and Cpk implies long-term capability. It should be noted that some statistical software programs (e.g. Minitab) or statistical publications might refer to Pp and Ppk as long-term capability and Cp and Cpk as short-term capability.

Use the flow diagram below to determine what statistical method is appropriate for the characteristic data sample.

Variable or Attribute Data?

Variable

Attribute

Conduct Normality Test per Paragraph C1

Calculate Cpk per Paragraph C2

Calculate Cpk per Paragraph C3

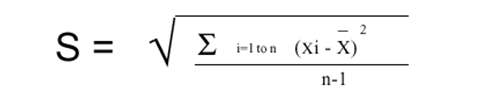
Calculate Cpk per Paragraph C4

Normal

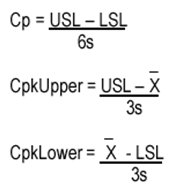
Non-Normal

C1. Normality Test – Using a minimum of 25 data points, create a histogram of the characteristic measured. If the histogram is bell (or lump) shaped and fairly symmetric, the sample is normally distributed and should have the Cpk calculated using methods described in section II. If the histogram is not bell shaped or is asymmetric, the sample is non-normal and should have the Cpk calculated using methods described in section III. If still unsure about the normality of the sample, run a statistical test for normality (Wilks-Shapiro, Chi-squared, etc…)

C2. Cpk Calculations for Normal Data – Sample standard deviation, s, is calculated using the following equation:



The Process Capability Ratio is calculated as follows:



Cpk is reported as the smaller value of CpkUpper and CpkLower.

C3. Cpk Calculations for Non-Normal Data – You can treat your data as attribute data and use the method described in section C4 below. Contact the purchaser for assistance.

C4. Cpk Calculations for Attribute Data – Use one of the following methods depending on the criteria listed below.

Method 1 – Inspect each characteristic in your sample and identify as defective or non-defective. Then calculate p(defective) = (number defective)/(number in sample). Use a standard normal table to find Z and divide by 3 to find Cpk or use the abridged Z/Cpk table below to define the Cpk.

Method 2 – Used if no defects are observed in the sample for methods 1 above.

Calculate the estimated proportion defective, p(d) = 1/(n+2) where n is the number of characteristics inspected, convert this number to a Z score using the one sided Z table and divide by 3 to obtain Cpk or use the abridged Z/Cpk table below.

|  |  |  |
| --- | --- | --- |
| Probability of defective = p(d) | Estimated Z | Cpk |
| Greater than 0.16 or > 16% | < 1 | < 0.33 |
| 0.16 to 0.023 or 16% to 2.3% | 1 to 2 | < 0.67 |
| 0.023 to 0.00135 | 2 to 3 | < 1.0 |
| 0.00135 to 0.000032 | 3 to 4 | > 1.0 but < 1.33 |
| Less than 0.000032 | > 4.0 | > 1.33 |

Method 3 - Used to estimate if Cpk > 1.0 with small samples. Cannot provide Cpk > 1.33 estimates. Given a very small sample (3 to 6 points) assessing process capability via direct calculation is not feasible. The approach outlined here assures that the process is well centered and has acceptable variation.

Variation test, calculate range of your sample of characteristics measurements

- Provide assurance that standard deviation of process is <= 16.6% of tolerance (that is, Z = 3)

Centering test, calculate mean of your sample

* Make sure that xbar is at least one standard deviation away from either tolerance limit with 99% assurance.

If your data satisfies both the mean and the range test as detailed in table below, you can estimate Cpk as > 1.0 (but not as high as 1.33).

|  |  |  |
| --- | --- | --- |
| # of Pieces | X Bar vs. Target | Range vs. Tolerance |
| 2 | +/- 3% | 31% |
| 3 | +/- 9% | 42% |
| 4 | +/- 12% | 49% |
| 5 | +/- 14% | 54% |
| 6 | +/- 16% | 57% |

C5. Typical Variable Data Charting/SPC Signs of process drift or instability

The following warning rules are commonly used in conjunction with the zones shown in Figure below:

* One Point falls beyond Zone A.
* Two out of three consecutive points on one side of the centerline fall in Zone A or beyond.
* Four out of five consecutive points on one side of the centerline fall in Zone B or beyond.
* Eight consecutive points fall anywhere on one side of the centerline.
* Six consecutive points steadily increasing or decreasing.
* Eight consecutive points on both sides of the centerline with none in Zone C.
* Fourteen (14) consecutive points alternating up and down.
* Fifteen (15) consecutive points on both sides of the centerline all in Zone C. This is a warning that the data may be too consistent. There could be a gage problem or some other event that has caused a shift in the output and should be investigated.

Upper Control Limit

+ 3 Sigma

+ 2 Sigma

+ 1 Sigma

- 1 Sigma

- 2 Sigma

- 3 Sigma

Lower Control Limit

Centerline

Zone A

Zone B

Zone A

Zone B

Zone C

Zone C

Table 2: CHARACTERISTIC ACCEPTANCE PLAN MATRIX (Ref. Appendix A, Table 1)

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Characteristic Type | PROCESS CONTROL CAPABILITY EVALUATION PLAN | MEASUREMENT DATA TYPE | PROCESS PERFORMANCE | PLANS ALLOWED |
| Supplier Defined KC:  Key Characteristics  OR  GE-AJA DRAWING:  - Critical  - Major  - Key Characteristic  See ([4]) | Process is IN CONTROL and CAPABLE for characteristic  “Control by Means other than 100% Characteristic Evaluation” permitted. Required to monitor by Control Chart | Variable | Cpk >= 1.33 | Variable Data:  Table 1 - #1 thru #8 |
| Attribute  [1] | ZERO  non-conformances  for an adequate [2] process period | Attribute Data:  Table 1 - All  except #4 |
| Process is NOT IN CONTROL and/or NOT CAPABLE for characteristic  “Control by Means other than 100% Characteristic Evaluation” is NOT permitted | Variable | Cpk<1.33 | Variable Data:  Table 1 - #1 |
| Attribute [1] | More than ZERO non-conformances for defined process [3] | Attribute Data:  Table 1- #1 |
| Non-Key Characteristics    Except characteristics described by [4] below | Process is IN CONTROL and CAPABLE for characteristic  “Control by Means other than 100% Characteristic Evaluation” permitted. Required to monitor by Control chart or Verification Plan | Variable | Cpk >= 1.0 | Variable Data:  Table 1 - #1  thru #8 |
| Attribute  [1] | <1 nonconformance per 750 evaluations over an adequate [2] process period | Attribute Data:  Table 1 - All  except #4 |
| Process is NOT IN CONTROL and/or NOT CAPABLE for characteristic  “Control by Means other than 100% Characteristic Evaluation” is NOT permitted | Variable | Cpk <1.0 | Variable Data:  Table 1 - #1 |
| Attribute  [1] | >1 nonconformance per 750 evaluations [3] | Variable Data:  Table 1 - #1 |
| [4] Inaccessible or other characteristics controlled by a special plan | Requires GE-AJA approved Acceptance Plan; such plans must include process control provisions | Variable or Attribute | Generic process capability Study or other capability assessment | Requires special plan plus Table 1 - #2 thru #8 |

[\*] Supplier establishes/explains a notation (other than above Drawing notations) e.g., [KC]. This notation included in the CLASS column of form GT1007-3.

[1] Characteristics required by Drawing and evaluated by “Attribute” gages which accept/reject to specific limits. Does not include characteristics which are not generated intentionally (e.g. scratches, dents, tool marks). 100% characteristic evaluation must be performed for NDE (Nondestructive Evaluation), when required on the drawing or in a referenced specification.

[2] “Adequate” period considers all common causes for process variation. A written rationale is defined by supplier and accepted by GE-AJA Quality Representative.

[3] A process which produced nonconformances may be re-evaluated after introduction of corrective action.

[4] Certain characteristics (e.g. certain cast dimensions) may require or justify special GE-AJA approved Acceptance Plans.

# Appendix D: OPERATOR Acceptance plan

D1. Purpose

To establish minimum requirements for an Operator Acceptance Plan, hereafter referred to as OAP. This plan will allow certified operators to verify characteristics at the point of generation. All elements of the plan are subject to purchaser disapproval.

D2. Minimum Requirements

D2.1 The supplier’s OAP shall identify provisions for training, certification, work station audits, disqualification, records and retention.

D2.2 Only certified operators or inspectors shall perform final verification of product characteristics.

D2.3 Characteristics generated by non-certified operators shall be verified by a certified operator or inspector.

D2.4 Traceability of measured characteristics to the inspector/certified operator shall be maintained to the part/lot.

D2.5 Recertification requirements shall be identified

D3. Training

D3.1 The Supplier’s OAP shall provide a process for training all operators on the procedures and work instructions that pertain to their immediate job function. Each operator shall be trained on the following, as applicable:

D3.1.1 Measurement and test equipment

D3.1.2 Engineering drawings

D3.1.3 Router/Op sheets/Work Instructions, usage and documentation

D3.1.4 Non-conforming hardware

D3.1.5 Safety and part handling

D3.1.6 Visual inspection techniques (e.g. tin soldier inspection)

D3.1.7 Geometric Tolerancing

D3.2 Consideration shall be given to the following when developing individual operator’s training.

D3.2.1 Previous related experience

D3.2.2 Performance reviews

D3.2.3 Job safety analysis results

D3.2.4 Non-conformance data

D3.2.5 Customer complaints/returns

D3.2.6 Internal workstation audit results

D3.2.7 Difficulty and criticality of the operation

D4. Certification

Each candidate shall be evaluated to assure their understanding of the training material and their ability to perform and document the required measurements. See example in Exhibit A

D5. OAP Workstation Audits

D5.1 Each certified operator shall be re-evaluated to an established audit plan. Audits shall be performed at least once per year, using a workstation audit form (see Exhibit B for an example)

D5.2 Satisfactory workstation audit completion shall result in continued certification. The records shall be updated, and the next audit date shall be established.

D5.3 If an operator fails the workstation audit, the Supplier will determine if re-training and/or increased audit frequency is necessary.

D5.4 Upon failed audit, the Supplier shall have a process to determine root cause, recommend a corrective action, and investigate whether non-conforming material was shipped to The Purchaser.

D6. Operator Disqualification - The Supplier shall develop a system for operator disqualification. Consideration shall be given to the following:

D6.1 Failure to follow documented work instructions

D6.2 Failure to pass an OAP workstation audit

D6.3 Inability to repeat/correlate measurements

D6.4 Change in job function or classification

D7. Record Retention

D7.1 Records pertaining to the Operator Acceptance Planning shall include, but are not limited to:

D7.1.1 Evaluation and training (initial training and any re-training)

D7.1.2 Certification test results

D7.1.3 Audit results

D7.2 Certification records shall be maintained for the entire duration of the operator’s employment, and audit results retained as an administrative quality record in accordance with S-1005.

**OAP EVALUATION FORM**

**Exhibit A**

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| OPERATOR/INSPECTOR NAME: | |  | | | BADGE: | | | |  |
| CLASSIFICATION: |  | | UNIT: |  | | SHIFT: |  | | |
| REQUIREMENTS | | | | | | | | | |
| Understands what and when to check and record, when required | | | | | | | |  | |
| Understands how to properly correct errors | | | | | | | |  | |
| Understands responsibility for all dimensions generated and checked | | | | | | | |  | |
| Understands what gages and fixtures to use | | | | | | | |  | |
| Understands that only gages**/**fixtures/instruments in cycle may be used | | | | | | | |  | |
| Understands method of documenting accept/reject decisions | | | | | | | |  | |
| Understands how to document a nonconformance | | | | | | | |  | |
| Understands no substitute gages, fixtures or tooling are allowed without prior written authorization | | | | | | | |  | |
| Understands what to do if a gage, fixture, instrumentor tool is damaged, or has overdue calibration | | | | | | | |  | |
| Understands responsibility of reviewing part paperwork, including special instructions and CTP (Continue to Process) dispositions prior to starting | | | | | | | |  | |
| OPERATOR/INSPECTOR HAS BEEN FULLY INSTRUCTED ON | | | | | | | | | |
| Identifying/measuring machined features such as Radii, Surface Finish, tool marks, etc. | | | | | | | |  | |
| Assuring that the previous operation is properly completed on the part paperwork and nonconformances are cleared for CTP (Continue to Process) disposition. | | | | | | | |  | |
| How to properly identify and document non-conformances | | | | | | | |  | |
| The need to follow work instructions as written, using the most current work instruction document | | | | | | | |  | |
| Use of proper Personal Protective Equipment (safety glasses, gloves, etc.) | | | | | | | |  | |
| How to use gauging/fixtures/instruments | | | | | | | |  | |
| Use of approved markers for temporary part marking | | | | | | | |  | |
| **DURING THE COMPLETION OF THE OPERATION** | | | | | | | | | |
| Does the employee follow Operation Sheets? | | | | | | | |  | |
| Does the employee use proper shop practices, tools, fixtures, etc.? | | | | | | | |  | |
| When required, does the employee make the proper checks and document properly? | | | | | | | |  | |
| **AFTER COMPLETION OF THE OPERATION** | | | | | | | | | |
| Is the part paperwork correctly completed? | | | | | | | |  | |

**DIMENSIONAL VERIFICATION** (5 MEASUREMENTS MINIMUM)

PART NUMBER:\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_ OPERATION NUMBER: \_\_\_\_\_\_\_\_\_\_\_\_\_\_\_

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| SERIAL NO. | OPERATOR VALUE | VERIFIED VALUE | DIFFERENCE | GAGE | Meets FAI gage requirements | Correlation? |
|  |  |  |  |  |  |  |
|  |  |  |  |  |  |  |
|  |  |  |  |  |  |  |
|  |  |  |  |  |  |  |
|  |  |  |  |  |  |  |

Satisfactory completion of this form certifies the employee to perform all operations in their classification in this unit unless noted below:

**OAP WORKSTATION AUDIT FORM**

**Exhibit B**

|  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- |
| OPERATOR/INSPECTOR NAME: | |  | | | BADGE: | |  | |
| CLASSIFICATION: |  | | Dept: |  | | SHIFT: | |  |

|  |  |
| --- | --- |
| Does part have matching router? |  |
| Does Router match the work instruction? |  |
| Is part marked with part no. and serial no? |  |
| Are work instructions being used? |  |
| Are gages at workstation within calibration cycle? |  |
| Are the correct gages being used? |  |
| When required, are all items from the planning recorded on the Log Sheet? |  |
| Are all nonconformances identified and documented? |  |
| Are approved markers/ink being used to mark on part? |  |

DIMENSIONAL VERIFICATION (3 MEASUREMENTS MINIMUM)

PART NUMBER:\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_ OPERATION NUMBER: \_\_\_\_\_\_\_\_\_\_\_\_\_\_\_

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| SERIAL NO. | OPERATOR VALUE | VERIFIED VALUE | DIFFERENCE | GAGE | Correlation? |
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| AUDIT PERFORMED BY: |  |

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| OPERATOR/INSPECTOR SIGNATURE: |  |

|  |  |  |
| --- | --- | --- |
| CONTINUE OAP CERTIFICATION? | YES | NO |

|  |  |  |
| --- | --- | --- |
| RETRAINING REQUIRED? | YES | NO |

|  |  |  |  |  |  |
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|  | |  |  |  |  |
| QUALITY SIGNATURE | | DATE |  | \* SUPERVISOR SIGNATURE | DATE |
|  |
| NEXT AUDIT DATE |

**Appendix E**

**FAI Applicability Guidance**

Is the part procured to a GE part number with a GE production release drawing or GE specification?

No

FAI not required

Yes

Has the GE part revision changed since the manufacturer last made the parts?

Has the manufacturing company and location made the GE part number (independent of revision) in the past 24 months?

Is this a unique single run production order? (not intended for ongoing production)

Yes

No

YES

Yes

Yes

Has there been a change in the manufacturing process or a change in sub-tier manufacturer or a change in the product acceptance plan?

Partial (Delta) FAI or Full FAI Required

Does the manufacturer have an approved FAI on file for the GE part number?

Full FAI Required

No

No

Yes

No

Yes

No