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MIL-I-45208 Inspection System

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#### APPLICATION NOTES (delete prior to release):

This inspection system is based upon MIL-I-45208 and is subject to Customer evaluation and verification. 3.13

The paragraph numbers in this quality manual do not correspond to the paragraph numbers in the MIL-I standard. This quality manual displays superscript numbers to establish the relationship between the standard and content in this quality manual. Superscript numbers correspond to paragraph numbers from MIL-I-45208A.

Paragraph numbers 1 and 2 and 4 through 6 in MIL-I-45208A only provide guidance (except 2.1) and do not require reference in the quality manual.

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#### **1.0SCOPE**<sup>3.1</sup>

It is a policy of the Company to perform all activities in a manner that reflects a total commitment to quality. This means maintaining the highest standards of quality in all products and services and a dedication to the principle of maintaining the highest levels of quality and integrity in communicating with people inside and outside of the Company. It is also a policy of the Company to prevent production and distribution of products that would pose unreasonable risks to health, safety, or the environment. It is a goal of the Company to encourage all employees to strive for individual excellence in their work and in their association with other people inside and outside of the workplace. The Company strives to motivate employees to achieve this excellence by providing leadership, training, proper materials, facilities and a cooperative environment.

Managers are responsible for developing organizations and systems that accommodate the goal of achieving Customer satisfaction. Managers must recognize and support employees charged with the responsibility of interacting with Customers. Employees who are authorized to work with Customers are responsible for carefully listening and fully understanding their requirements and expectations. These employees should be as responsive as possible to those needs within the province and spirit of good business practices. Managers must monitor Customer satisfaction on a continuing basis, making appropriate adjustments and corrections if problems occur. This Quality Manual is produced to provide guidance to achieve the policies and goals of the Company. This manual of policies and procedures are subject to review by the Customer. The Company's Mission is to continually improve products and services.

#### 2.0 ORGANIZATION

#### 2.1 Quality Responsibility and Authority<sup>3,2,3</sup>

The quality manager has the responsibility and authority to resolve matters relative to quality in products, processes, and services from internal and external sources. Quality may suspend internal and external processes and services that do not meet requirements until appropriate corrective and preventive action is implemented on an expedited, high priority basis. In addition, Quality may withhold internal and external shipments of products that do not meet requirements until appropriate corrective and preventive action is implemented on an expedited, high priority basis. The quality manager reports directly to the President. Quality supervisors, inspectors, and auditors report directly to the quality manager.

#### 2.1.1 Problem Resolution

Quality problems resulting from a variance to a program requirement are resolved by the organizational Group assigned the specific responsibility. Decisions affecting Quality, Cost, or Schedule are recorded using documented correspondence. Company correspondence is distributed and retained. Each organizational Group has the authority, responsibility, and freedom to initiate, recommend or provide solutions for programmatic problems; however, each Group is expected to fulfill this inspection system at all levels and protect the quality effort of other Groups upon which they have an influence.

#### **2.2 Initial Quality Planning**<sup>3.11.1, 3.11.2</sup>

The Quality Group is responsible for review of new and pending work based on the receipt of a Request for Proposal (RFP), the receipt of a new contract or potential contract or the activation

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of a Company-funded program to integrate special or unusual contract requirements into quality plans and procedures.

#### **2.3** Inspection and Testing Documentation<sup>3.2.1</sup>

#### 2.3.1 Preparation

All work affecting quality is described by clear and complete documented instructions of a type appropriate to the circumstance. Preparation, maintenance, reviews and compliance with instructions is accomplished in 'real-time' or as a result of the initial quality planning function.

2.3.2 Inspection Instructions<sup>3.9</sup>

The Quality Group prepares an inspection instruction sheet for all inspection work by performing tasks that may include, but are not limited to:

Prepare Inspection Instruction Sheet, (IIS). The IIS may include, but is not limited to:

IIS#	Specification number(s) and revision letter(s)
_Title of IIS	Mfg/QA Traveler/Planner# supported by the IIS
	5
1	
· 	

After approval the IIS is released for use where specified. The IIS is exempt from issue control; however,

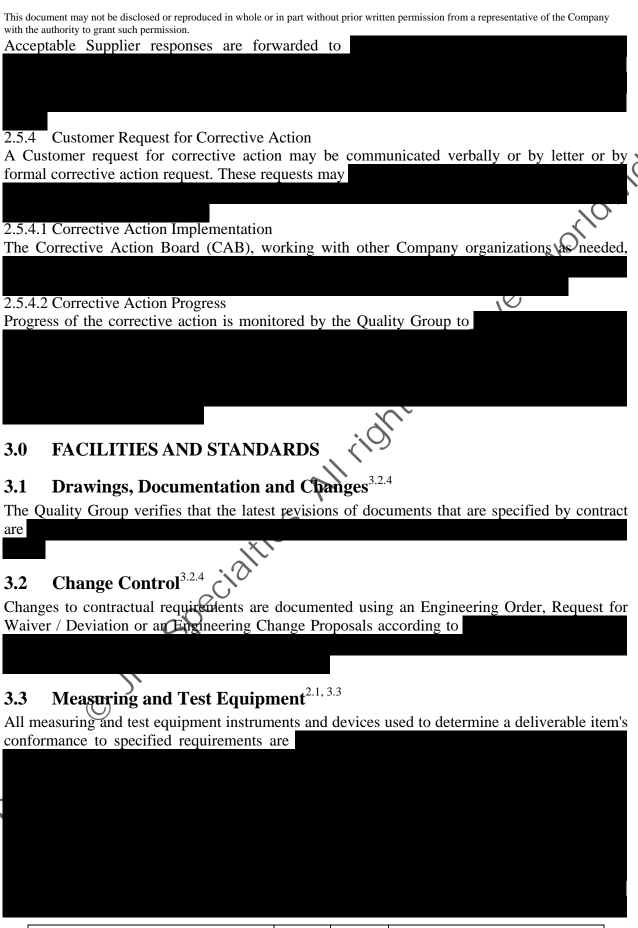
 $\textbf{Records}^{3.2.2}$ 

2.4.1 General

Data to be recorded includes

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2.4.2				
Reco	rds are examined for			
2.4.3	Record Maintenance			
		d to main	tain reco	rds as directed by the contract or for
	(7) years if not specified by the con			
50 / 01	(1) yours if not specified by the con	10000		e principal de orde de d
				70
2.4.4			11.	
		ned in the	e quality	area handling the inspection system.
Reco	rds are removed			
2.4.4	.1 Objective Evidence			· (C)
	rds are collected or produced to t	the extent	necessa	ry to
	3 2 2			is
2.5	<b>Corrective Action</b> <sup>3.2.3</sup>		X	
2.5.1	Internal Corrective Action Reques			)
		a Reques	for Cor	rective Action (RFCA) is initiated as
prom	ptly as practicable to			
2.5.2	Corrective Action Implementation	n by the N	IRB	
The 1	MRB forwards the CAR of RFCA t			
252	.1 Corrective Action Monitoring			
		nroveme	nte and c	orrections and the monitoring of the
	tiveness of actions taken are	iprovenie	ns and c	offections and the monitoring of the
CITOC	aveness of actions taken are			
2.5,3	Supplier Corrective Action			<u> </u>
				Purchasing Group or a Customer.
	<u> </u>			ipleted as specified by the Customer,
the N	IRB or by the Quality Group. The	CAR/RFO	CA form	is
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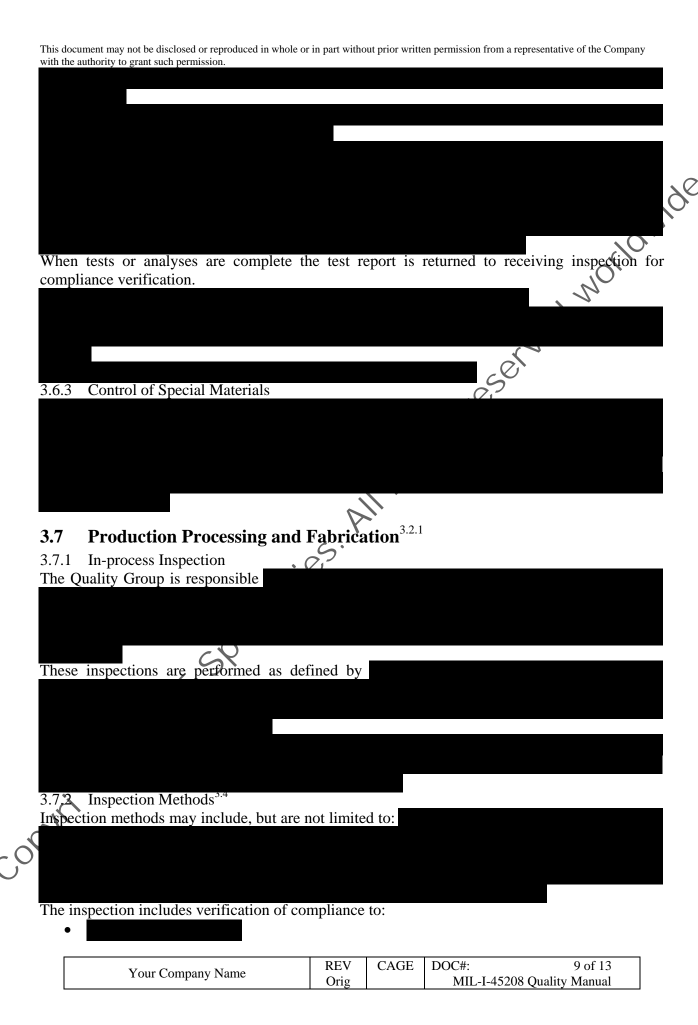
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Tools that are used for inspection purposes are calibrated prior to	1150
The environment where measuring and test equipment instruments and devices are	
calibrated and used is controlled to the extent necessary to assure required accuracy,	
consideration given to temperature, humidity, vibration, cleanliness and other controll factors.	abie
<b>(</b> 0	
3.4 Use of Contractor's Inspection Equipment <sup>3,3</sup>	
3.4.1 Availability Company owned gauges, inspection devices and test equipment are made available for use	e hv
Customers when	J Uy
<b>3.5</b> Control of Purchases <sup>3.11,3,111,3,11,3</sup>	
3.5.1 Procurement Document Requirements Review  The Quality Group reviews procurement documents to	
The Quality Group reviews procurement documents to	
The Supplier is directed to provide some or all of the following:	
•	
•	
•	
•	
3.6 Materials and Material Control <sup>3.9, 3.12</sup>	
3.6.1 Receiving Inspection	
All materials are evaluated by receiving inspection to the extent necessary to	

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with the authority to grant such permission.
Three levels of inspection sampling can be used:
Sampling to permit defects is not permitted.  When an item drawing is revised and/or when an item is purchased to a revision level that differs
from parts in stores, the early revision parts
Nom parts in stores, the early revision parts
Parts that have been sent out for special processing are
The count has made violation and the solid and the 1000/ improved in the value of the solid and the
The acceptable material from a lot subjected to 100% inspection may be released to production upon completion of appropriate documentation.
upon completion of appropriate documentation.
· C A
V 1
Receiving inspection personnel observe the following document order of precedence in the event
of conflict, a(16)guity or contradiction:  1.
2.
ર. દુ.
4.
5.
The Company's specifications do not take precedence
3.6.2 Raw Material Inspection The Purchasing Group specifies physical and/or chemical characteristics and properties on
purchase orders for raw materials. The purchase order requires the Supplier to provide
The same and the s

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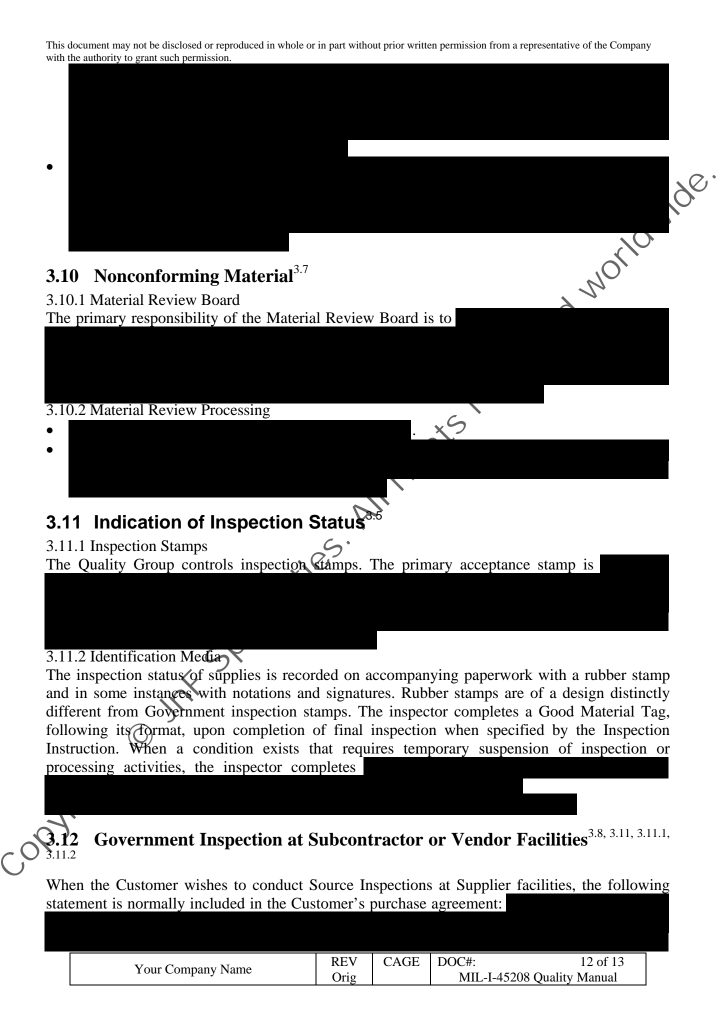
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•	
• W/b∘	n abveiced inspection of anneaged symples is impressible on disadvente seems indicate
	n physical inspection of processed supplies is impossible or disadvantageous, indirected of product quality is accomplished by
372	.1 Calculated Risk Release
In th Data limit	e event materials, components or assemblies are needed prior to receipt of Certified Test, Certificate of Compliance or Analysis, approved Request for Deviation or Waiver or other ed risk condition, cognizant MRB members of the Products and Quality Group may release rticles on a Calculated Risk.
An o	pen CRR prevents delivery of supplies unless waived by the Customer.
3.7.3	Identification <sup>3.5</sup>
Parts	or assemblies found to be in compliance with inspection requirements are identified as
3.7.4	
A M	laterial Report is initiated by process or inspection personnel for each failure detected ding those discovered during
3.7.5	
	production tools such as jigs, fixtures and templates used for producing deliverable supplies
are	
3.8	Completed Item Inspection and Testing <sup>3.2.1, 3.5</sup>
3.8.1	Final Physical and Visual Inspection
	inished goods are inspected as specified on the applicable inspection instruction or Traveler
	specified by the Quality Group. Parts and assemblies are processed only after all operations ified on applicable process documentation are identified as complete and accepted
	ections are made using
5	

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When the Customer contract is accepted, the Source Inspection statement is

## **3.13** Government Property<sup>3.6</sup>

Government and Customer property is controlled according to contractual requirements and applicable property and/or facility agreements, including, but not limited to:

- e.
  •
- 3.13.1 Bailed Property

Bailed property is controlled according to contractual requirements and applicable property and/or facility agreements.

Referenced Documents: (delete prior to release of manual)

ASTM D 3951 – Commercial Preservation Packageng, Packing and Marking QC-101 – Inspection Instruction Form QC-102 – Request for Corrective Action QC-103 – Material Report Form QC-104 – Calculated Risk Release Form QC-105 – Good Material Tag QC-106 – Withhold Tag QC-106 – Withhold Tag QC-107 – Routing Ticket QC-108 – R&I Inspection Record QC-109 – Engineering Order QC-110 – Request for Waiver / Deviation

QC-112 – Inspection Record
QC-113 Bad Material Tag
QC-114 – R&I Inspection Instructions
QC-115 – Property Control
QC-116 – Calibration System and Forms
QC-117 – Supplier Quality Requirements
QC-118 – Basic Contract Review
QC-119 – Traveler
QC-120 – Purchase Order
QC-121 – Purchase Order Review Instructions
QC-122 – Dimensional Analysis Record
QC-123 – Data List
QC-124 – Inspection System Survey

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SUPPLIER INFORMATION:		<b>CAGE CODE</b> :					
Supplier Name:		Supplier Code:					
Address:							
(Street)	(City)	(State)	(Zip)				
Quality Manager:	Phone:	Fax:	-inle				
SURVEY BACKGROUND INFORMAT	ION:		10,				
Reason for Survey: New Supplier	Recertification	Corrective Action Follow-Up					
Survey Date:	Ap	proval Date:					
Approval Method: Survey	History	o No					
(If History, attach summary) History summar	•	Yes No No					
Special Process Codes (if known)		XS (O					
SUPPLIER BACKGROUND INFORMA	ATION:						
Housekeeping is adequate Ye		)					
Floor Space is adequate Ye	es No						
Government Source Inspection is on Site Ye	es No						
GSI is Itinerant	os No						
Number of shifts 1	2 3						
Number of Employees	per in Quality	Years in Business _					
Delegated active Material Review Board Ye	es No 🗌	Delegated by					
% Government Business							
SURVEY RESULTS:	APPROVAL STAT	TUS: (A, C, or D)					
A = Approved	C = Conditional	D = Disapproved					
Survey Expiration Date (if required by applicable sp	ecification)						
Survey Follow-Up Required: Yes No	Additional Com	ments are attached: Yes No	о				
This survey was performed by							
Surveyor's Office Phone Number:		Survey was requested by:					
Signature:		Date:					

Audit and HDBK 51 Question Number	MIL-I-45208 Paragraphs	Acceptable	In space under 'Acceptable' enter 'Y' for conformance, 'N' for Nonconformance, or 'X' for Not Applicable. Under comments enter statement describing nature of nonconformance for each 'N' entry. Comments are mandatory for all objective evidence observed.
	1.0 Scope		ble)
	1.1 Scope (Not applicable)		
	1.2 Applicability (Not applicable)	12 1	
	<ul><li>1.2.2 Relation to other contract requirements (Not ap 1.2.3 Options (Not applicable)</li></ul>	риса	Die)
	2.0 Applicable Documents (Self explanatory)		CO
	2.1 General (Self explanatory)		2/6
	2.2 Amendments and revisions (Self explanatory)		
	2.3 Ordering Government Documents (Self explanate	ory)	
	3. Requirements		.07
	3.1 Contractor Responsibilities		(0
1. (1)	Does the inspection system cover all supplies and		*5
2 (2)	services offered to the Government for acceptance?		
2. (2)	Does the inspection system cover all supplies and services procured from subcontractors or vendors?	. 6	
3. (3)	Does the inspection system assure that all supplies and	1	2
3. (3)	services submitted to the Government for acceptance		
	conform to contract requirements?		
4. (4)			
. ,	5.		
	3.2 Documentation, Records and Corrective Action		
9. (1)	Are all inspection instructions clear, complete and up to		
	date?		
10. (2)	Are all required instructions available and current?		
11. (3)			
C 04	3.2.2 Records		
13. (1)	Does the contractor maintain adequate records of all		
	examinations and tests?		
14. (2)	Do the records indicate the nature and number of		
	observations made?		
15. (3)			

Audit and HDBK 51 Question Number	MIL-I-45208 Paragraphs	Acceptable	In space under 'Acceptable' enter 'Y' for conformance, 'N' for Nonconformance, or 'X' for Not Applicable. Under comments enter statement describing nature of nonconformance for each 'N' entry.  Comments are mandatory for all objective evidence observed.
17. (5)	3.2.3 Corrective Action		10,
18. (1)	Is action taken promptly to correct all conditions that		
	cause defects to be submitted for Government		. 1/0
10 (2)	acceptance?		\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \
19. (2)			100
			, eserve
	3.2.4 Drawings and Changes	I .	
21. (1)	Does the contractor's inspection system provide		nis
	procedures that assure that only the latest applicable	_	
	drawings, specifications and instructions, including all approved changes, are used for fabrication, examination	1	D)
	and testing?	1	
22. (2)			
	3.3 Measuring and Test Equipment		
23. (1)	Are the gauges, testing and measuring equipment that are necessary to assure that products meet technical requirements available and are procedures established for their use?		
24. (2)	Is the test and measuring equipment properly		
25. (3)	maintained?		
25. (5)			
~ OX			
32. (10)	Does the contractor make inspection equipment and		
	facilities available to the Government representative for verification of the contractor's results where required?		
33. (11)	vermeation of the contractor's results where required?		
23. (11)			

Audit and HDBK 51 Question Number	MIL-I-45208 Paragraphs  3.4 Process Controls	Acceptable	In space under 'Acceptable' enter 'Y' for conformance, 'N' for Nonconformance, or 'X' for Not Applicable. Under comments enter statement describing nature of nonconformance for each 'N' entry.  Comments are mandatory for all objective evidence observed.
34. (1)	Are there contract or specification requirements for		,0
3 (1)	control of any specific manufacturing processes or operations?		NOT.
35. (2)			8
	3.5 Indication of Inspection Status	1	10
36. (1)	Does the contractor have an effective system for identifying the inspection status of products?		
37. (2)			(6)
	3.6 Government Furnished Material		
	3.6.1 Damaged Government Furnished Material (GF	'M)	×S
38. (1)	Does the contractor examine GFM upon receipt for damage, quantity, completeness and type?	٠.	
39. (2)			2
42. (5)	Does the contractor record and report to the Government any damage, malfunction or deterioration of GRM prior to, during and after installation?		
43. (6)			
	3.7 Nonconforming Material	ı	
44. (1)	Does the contractor have an effective system for controlling nonconforming material?		
45. (2)			
	<ul><li>3.8 Qualified Products (Not applicable)</li><li>3.9 Sampling Inspection</li></ul>		

Audit and HDBK 51 Question Number	MIL-I-45208 Paragraphs	Acceptable	In space under 'Acceptable' enter 'Y' for conformance, 'N' for Nonconformance, or 'X' for Not Applicable. Under comments enter statement describing nature of nonconformance for each 'N' entry. Comments are mandatory for all objective evidence observed.
49. (1)	Do required sampling procedures conform to the applicable specification or other procurement documents?		orld
50. (2)			, N
	3.10 Inspection Provisions	ı	
51. (1)	Has the contractor elected to use any inspection equipment or procedures other than those specified or referenced in the contract?		, eserved
52. (2)			(es
	3.11 Government Inspection at Subcontractor or Ver 3.11.1 Government Inspection Requirements 3.11.2 Purchasing Documents 3.11.3 Referenced Data	ndor l	Facilities
56. (1)	Do contractor purchasing documents require Government source inspection of supplies only when the Government so requested?		
57. (2)			
	3.12 Receiving Inspection		
59. (1)	Is all received material inspected as necessary to assure conformance with contractual requirements?		
60.63	Is the Government representative notified of all defects found in material subjected to Government procurement quality assurance actions at source?		
61. (3)			
	3.13 Government Evaluation	•	
62. (1)	Does the contractor permit the Government representative to evaluate the inspection system and the		

Audit and HDBK 51 Question Number	MIL-I-45208 Paragraphs	Acceptable	In space under 'Acceptable' enter 'Y' for conformance, 'N' for Nonconformance, or 'X' for Not Applicable.  Under comments enter statement describing nature of nonconformance for each 'N' entry.  Comments are mandatory for all objective evidence observed.
	supplies it generates?		190
63. (2)			ed work
	NOT	ES	arve
			6
		<u>``</u>	2),
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	°C,		
	- 58		
	<u> </u>		
Survey	red by: Ti	tle:	
Site:	Ci	ty:	State:
Teleph	one No		

# **Quality Systems Cross Reference Matrix**

<b>Quality System Elements</b>	MIL-I- 45208A	MIL-Q- 9858	ISO 9001:94	ISO 9001:2008	ISO 9001:2015
Management Responsibility:	3.1	1.3, 3.1	4.1	5.1, 5.3, 5.4.1, 5.5.1, 5.5.2, 5.6, 6.1, 6.2.1, 8.5.1	
Quality System, Initial Quality Planning:	1.1	1.3, 3.2	4.2	4.1, 4.2.1, 4.2.2, 5.4.2, 7.1	
Contract Review:	1.2	3.2, 1.4	4.3	5.2, 7.2.1, 7.2.2, 7.2.3	
Design Control:	N/A	4.1	4.4	7.2.1, 7.3	2
Document and Data Control:	3.2	4.1	4.5	4.2.3	0
Purchasing:	N/A	5	4.6	7.4.1, 7.4.2, 7.4.3	
Control of Customer Supplied Product:	3.6	7.2	4.7	7.5.4	
Product Identification and Traceability:	N/A	6.1	4.8	7.5:3	
Process Control:	3.4	6.2	4.9	63, 6.4, 7.5.1, 7.5.2	
Inspection and Testing:	3.1, 3.2.1, 3.12	6.1, 6.2, 6.3	4.10	7.1, 7.4.3, 7.5.3, 8.1, 8.2.4	
Control of Inspection, Measuring and Test Equipment:	3.3	4.2-8.5	4.11	7.6	
Inspection and Test Status:	3.5	06.7	4.12	7.5.3	
Control of Nonconforming Product:	30	6.5	4.13	8.3	
Corrective Action:	3.2.3	1.3, 3.5	4.14	8.5.2, 8.5.3	
Handling, Storage, Packaging, Preservation, and Delivery:	3.6	6.4	4.15	7.5.1, 7.5.5	
Control of Quality Records:	3.2.2	3.4	4.16	4.2.4	
Internal Quality Audits:	N/A	N/A	4.17	8.2.2, 8.2.3	
Training:	N/A	N/A	4.18	6.2.2	
Servicing:	N/A	1.3	4.19	7.5.1	
Statistical Techniques:	N/A	6.6	4.20	8.1, 8.2.3, 8.2.4, 8.4	

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37 T	Special Instructions:	
Your Logo		

REQUEST FOR CORRECTIVE ACTION

1	RFCA#:	Date:	MR#:
2	Internal		External
3	To:	Return To: Your (Attention: Address:	Co.
4	Classification of Defect  Critical Major Minor  Required Response(Working Days)  Days 15Days 30Days  Implement Next Purchase Order	Lot Qty:	Report#: Spec#: Reject Qty:
5			
11			

Your Logo QC-102 (mo/yr)

	Nonconfor	mance	e Rep	ort	: <b>Dispos</b>	sition I	Process Rev:
			(mo	o/yr	Allid	is les	
	· · · · · · · · · · · · · · · · · · ·	Revisio	ons				Rev:
Letter	E.O. Number	Desci	ription				Date
	SQ						
	X						
	711.						
	X O						
Used On	Contract#:						
66,					You	r Comp	any Name
Prepared By:		Date				r	
Your Dept:		Date					
Your Dept:		Date					OGRAM
Your Dept:		Date	G: 1		1	our Proc	
Your Dept:		Date	Size:	A	CAGE:		Your Form # (mo/yr) 1 of 1

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Reporting Agent	When a nonconformance, continuous improvement or calculated risk condition occurs in manufacturing, testing or inspection, record the condition on the top-half of a Material Report, QC-103-2, following its format. Do not
Reporting Agent	Forward completed MR to Document Control (DCC).
DCC	Enter MR into the routing database, copy the MR, stamp DCC on the form and forward original to the Quality Mgr.
Quality Mgr.	
1st MRB Reviewer	
IF	THEN
MRB Staff	
	Reporting Agent DCC  Quality Mgr.  1st MRB Reviewer  IF Engineering Order (EO) or Request for Waiver (RFW) is the normal course of action 1st MRB Reviewer

Your Company Name	REV	CAGE	DOC#:	2 of 2
			Your Procedure #	

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ermission.	IF	THEN
	MRB Member	TIILIV
	Disagrees with	
	recorded	
	disposition	
	disposition	
5.1	MRB Staff	Perform actions required to maintain the disposition status on the MR Form, e.g., re-sign MR Form A/R to keep it current through each disposition event; hand-carry for completion, caucus for consensus, etc.
6.0	Quality Mgr.	
	IF	THEN
	Customer	Forward MR to Configuration and Discrepancy Mgr. for
	Required	retrieval of Customer concurrence of disposition or signature when required by contract (RFW or ECP A/R).
6.1	Quality Mgr.	Upon completion of the MRB, forward the completed MR to the Configuration and Discrepancy Mgr.
7.0	Configuration and	
	Discrepancy Mgr.	

Your Company Name	REV	CAGE	DOC#:	3 of 3
1 3			You	ır Procedure #

# MATERIAL REPORT

CONTRACTOR:		DATE RECEIVED:	
MR#:		SHEET	OF
raveler#:	Op#:	Quantity Received: Job Number:	
Item Name:		Description: ID S/B Spec#, Para#, & IS Condition w/Quantity &Dimension Affected	# Discrep
Dwg/Spec:			
Part#:			
Part# Rev:			
Lot or S/N:			
P.O.#:			

## CALCULATED RISK RELEASE

Date:		

Your Logo QC-104 (mo/yr)

Green = Good, Yellow = Withhold, Red = Bad

Use standard, colored card stock – size approximately 3.5" tall by 5.75" wide or use stock size

GOOD M	MATERIAL TAG		Your Logo				
P/N:	PO #:		Date:				
Dwg #:	Rev:		Lot #:				

QC-105 (mo/yr)

GOOL	MATERIAL TAG	Your Logo		
P/N:	PO #:		Date:	
Dwg #:	Rev:		Your Lot #:	

QC-105-1 (mo/yr)

WITHHOLD TAG	Your Logo
(mo/yr)	

Your Logo				
Item Name:				
Item P/N:				
	Item Name:			

QC-113 (mo/yr)

			**							
GOO	OD MATERIAL TAG		Yo	our Logo		GO	OD MATERIAL TAG			Your Logo
P/N:		Rev:	I	Date:		P/N:		Rev:		Date:
PO#:		Le	ot#:			PO#:		I	.ot#:	
MR#:		Qty	Ok:			MR#:		Qty	Ok:	
Ready F	or:					Ready F	or:			
Initia	als:					Initia	als:			
				QC-105-2 (mo/yr)						QC-105-2 (mo/yr)
GO	OD MATERIAL TAG		Yo	our Logo		GO	OD MATERIAL TAG			Your Logo
P/N:		Rev:	I	Date:		P/N:		Rev:	S	Date:
PO#:		Lo	ot#:			PO#:		N	ot#:	
MR#:		Qty	Ok:			MR#:		Qty	Ok:	
Ready F	or:					Ready F	or:	5		
Initia	als:					Initia	VI			
				QC-105-2 (mo/yr)			,0,5			QC-105-2 (mo/yr)
GO	OD MATERIAL TAG		Yo	our Logo		GO	OD MATERIAL TAG			Your Logo
P/N:		Rev:	I	Date:		P/N:		Rev:		Date:
PO#:		Lo	ot#:			PO#:	9)	I	.ot#:	
MR#:		Qty	Ok:		-	MR#:		Qty	Ok:	
Ready F	or:					Ready F	or:			
Initia	als:					Initia	als:			
				QC-105-2 (mo/yr)						QC-105-2 (mo/yr)
GO	OD MATERIAL TAG		Yo	our Logo		GO	OD MATERIAL TAG			Your Logo
P/N:		Rev:	I	Date:		P/N:		Rev:		Date:
PO#:			ot#:			PO#:		I	.ot#:	
MR#:		Qty	OK.			MR#:		Qty	Ok:	
Ready F	or:	, 6	7			Ready F	or:	•		
Initia	als:	<b>(</b>				Initi	als:			
	- 7			QC-105-2 (mo/yr)	l		1			QC-105-2 (mo/yr)
GO	OD MATERIAL TAG		Yo	our Logo		GO	OD MATERIAL TAG			Your Logo
P/N:	· dh	Rev:	I	Date:		P/N:		Rev:		Date:
PO#:	1/1/2	Le	ot#:			PO#:		I	ot#:	
MR#:	01	Qty	Ok:			MR#:		Qty	Ok:	
Ready F	or:					Ready F	or:			
Initia	als:					Initi	als:			
				QC-105-2 (mo/yr)						QC-105-2 (mo/yr)

WITH	HOLD TAG	Your Logo					
Date:		Item Name:					
PO #:		Item Part Number:					
Lot #:		Material Report #:					
S/N:		Initials:					
Reason for Withholding:							

QC-106-1 (mo/yr)

WITH	HOLD TAG	Your Logo	
Date:		Item Name:	
PO #:		Item Part Number:	
Lot #:		Material Report #:	
S/N:		Initials:	

Reason for Withholding:

OC-106-1 (mo/v)

QC-106-1 (mo/yr)

WITH	HOLD TAG	Yo	our Logo	
Date:		Item Name:	)×	
PO #:		Item Part Number:		
Lot #:	*	Material Report #:		
S/N:	:00	Initials:		
Reason for Withholding:				
	Oz.			

WITH	HOLD TAG	Your Logo		
Date:		Item Name:		
PO #:		Item Part Number:		
Lot #:		Material Report #:		
S/N:		Initials:	76.	
Reason	for Withholding	:	ANII	
			7/0	

WITH	HOLD TAG	el Yo	our Logo
Date:	, (O.	Item Name:	
PO #:	Wis	Item Part Number:	
Lot #	(2)	Material Report #:	
S/N:		Initials:	
* Reason f	for Withholding	:	

QC-106-1 (mo/yr)

WITH	HOLD TAG	Your Logo	
Date:		Item Name:	
PO #:		Item Part Number:	
Lot #:		Material Report #:	
S/N:		Initials:	

QC-106-1 (mo/yr)

#### Helpful Hint:

Purchase green "presentation" paper for the Good Material Tag and yellow "presentation" paper for the Withhold Tag, then print and cut whenever you need...

ACCEPTED MATERIAI	_	Y	Your Logo	
THIS MATERIAL HAS BEE	THIS MATERIAL HAS BEEN INSPECTED AND ACCEPTED			
P/N:	Rev:		Date:	
PO#:	I	.ot#:		
MR#:	Qty	Ok:		
Initials:	•			

ACCE	ACCEPTED MATERIAL			Your Logo
THIS	MATERIAL HAS BEEN I	NSPEC	TED A	AND ACCEPTED
P/N:		Rev:		Date:
PO#:		I	Lot#:	
MR#:		Qty	Ok:	
Initia	als:			

QC-105-3 (mo/yr)

ACCEI	PTI	ED MATERIAL	Your Logo			A	
THIS	MA'	TERIAL HAS BEEN I	NSPEC	TED A	AND ACCEPTED		
P/N:			Rev:		Date:	7	
PO#:			I	Lot#:		] _ '	~
MR#:			Qty	Ok:	•	Ó.	N
Initia	ls:						
					QC-105-3 (mo/yt)	_	
ACCEI	ACCEPTED MATERIAL		Your Logo			A	
THIS	MA'	TERIAL HAS BEEN I	NSPEC	TED	AND ACCEPTED	1	

	PTED MATERIAL	Your Logo
THIS	MATERIAL HAS BEEN I	INSPECTED AND ACCEPTED
P/N:		Rey Date:
PO#:		Lot#:
MR#:		Qty Ok:
Initia	als:	

QC-105-3 (mo/yr)

ACCE	PTED MATERIAL	Your Logo			
THIS	THIS MATERIAL HAS BEEN INSPECTED AND ACCEPTED				
P/N!		Rev:		Date:	
PO#:		I	Lot#:		
MR#:		Qty	Ok:		
Initia	als:				

QC-105-3 (mo/yr)

	PTED MATERIAL	Your Logo VINSPECTED AND ACCEPTED		
P/N:		Rev:		Date:
PO#:		I	Lot#:	
MR#:		Qty Ok:		
Initia	als:			

	PTED MATERIAL	<b>\</b>	Your Logo
THIS	MATERIAL HAS BEEN I	INSPECTED A	AND ACCEPTED
P/N:		Rev:	Date:
PO#:		Lot#:	
MR#:	.76	Qty Ok:	
Initia	als:		

QC-105-3 (mo/yr)

	ACCEPTED MATERIAL			Your Logo			
L	THIS	HIS MATERIAL HAS BEEN INSPECTED AND ACCEPTED					
-	P/N:			Rev:		Date:	
1	PO#:		Lot#:				
Ī	MR#:			Qty Ok:			
	Initia	als:					

QC-105-3 (mo/yr)

ACCE	ACCEPTED MATERIAL			Your Logo		
THIS	THIS MATERIAL HAS BEEN INSPECTED AND ACCEPTED					
P/N:			Rev:		Date:	
PO#:			I	Lot#:		
MR#:			Qty	Ok:		
Initials:						

QC-105-3 (mo/yr)

ACCE	ACCEPTED MATERIAL			Your Logo		
THIS	THIS MATERIAL HAS BEEN INSPECTED AND ACCEPTED					
P/N:			Rev:		Date:	
PO#:			I	Lot#:		
MR#:			Qty	Ok:		
Initials:						

QC-105-3 (mo/yr)

FINAL INSPECTION	FINAL INSPECTION			
PERFORMED BY	PERFORMED BY			
YOUR LOGO	YOUR LOGO			
QC-105-4 (mo/yr)	QC-105-4 (mo/yr)			
FINAL INSPECTION	FINAL INSPECTION			
PERFORMED BY	PERFORMED BY			
YOUR LOGO	YOUR LOGO			
QC-105-4 (mo/yr)	QC-105-4 (mo/yr)			
FINAL INSPECTION	FINAL INSPECTION			
PERFORMED BY	PERFORMED BY			
YOUR LOGO	YOUR LOGO			
QC-105-4 (mo/yr)	QC-105-4 (mo/yr)			
FINAL INSPECTION	FINAL INSPECTION			
PERFORMED BY	PERFORMED BY			
YOUR LOGO	YOUR LOGO			
OC-I05-4 (mo/yr)	QC-105-4 (mo/yr)			
FINAL INSPECTION	FINAL INSPECTION			
PERFORMED BY	PERFORMED BY			
GOUR LOGO	YOUR LOGO			
QC-105-4 (mo/yr)	QC-105-4 (mo/yr)			
FINAL INSPECTION	FINAL INSPECTION			
PERFORMED BY	PERFORMED BY			
YOUR LOGO	YOUR LOGO			
QC-105-4 (mo/yr)	QC-105-4 (mo/yr)			

## Helpful Hints:

Purchase "presentation" paper in your choice of color and then print and cut labels whenever you need.

Purchase peel-and-stick labels of the correct size and then print whenever you need.

## Your Logo

## ROUTING TICKET

ACCOUNT#:

	necounti.				
Operator:		Date:			
XXX Lot#:		XXX Lot#:			
Prepare Routing Ticket for each QC-107 (					

Drawing No:	RECEIVING INSPECTION REPORT			
Item Name:	Your Co			
Sampling Plan:				
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3				
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34 35				
35				

Your Logo QC-108 (mo/yr)

<b>EO NUMBER:</b>	DATE:	MR#:	
<b>ENGINEERING</b>	CLASS	PERSON REQUEST ENGINEERING ORI	DER:
	I II	PERSON WRIT	
ORDER Page of	CUSTOMER APPROVAL	CUSTOMER CONCURRENCE	EXISTING PARTS AFFECTED
HOLD PO'S PENDING APPROVAL  YES NO	YES NO	YES NO	YES NO
Your Logo			OC-109 (mo/yr)

Your Logo	REQUEST FOR DEVIATION / WAIVER							DEVIATION / WAIVER						
1. NAME AND ADDRESS		2. CAGE CO			3. RDW NO.									
Your Co		4. PURCHAS	SE ORDER NO.	:	5. DATE									
		4a. PURCHAS	SE ORDER LINE NO	O. (	6. DEVIATION	WAIVER								
7.														
12.														
						QC-110 (mo/yr)								

Drawing No:									IN	ISPE	ECTIO		ORD								Q	C-112 (mo/yr)
Item Name:				(Your Co)								Front										
	_				(Description of Inspection Process)																	
																	$\Box$			+		
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34																						

			Receiving Inspection Instru	ctions	QC-114	(mo/yr) Page 1 of 1
	** -		Special Instructions:	Specification:	-	
	Your L	ogo	ANSI Z 1.4; Level I reduced, AQL 1.0 Die-controlled = 5/lot	Specification:		
			Commercial or items >50Lbs = 1/Lot	Approval:		
Oper	Qty		on of Inspection Operation	l	Gage	Comment
R&I		Op 1				
		Op 2:				
		O 4. V	if the Compliant Little Line (Learning Little Line)	N 11 T . ! 4		
		Op 4: Vei	rify the Supplier is listed in the approved S	Supplier List		
		Ор				
		Op 9:				
		Op 10: V	erify			
		Op 11: V	erify			
		Op 11. V	City			
		Op 12: V	erify			
		On 13. A	ffix a Good Material Tag to acceptable sur	onlies. For supplies that ex	xhihit	
		a lot num		opines, i or supplies that ex		
					_	
		On 14. Pr	repare a Material Report for nonconforming	o sunnlies		
			omplete inspection record QC-108 and rec			
		number(s	) in the Remarks field			
		Op 16:			_	
					l	

## LLC. All rights reserved worldwide. **Property Control**

Rev: E.O. Number Description Letter Date Contract#: **Your Company Name** Prepared By: Date Your Dept: Date **YOUR PROGRAM** Your Dept: Date Your Dept: Your Procedure # Date Your Dept: CAGE: Date Size: 1 of 10 Your Form # (mo/yr)

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15.0 WORKMANSHIP	

Vour Company Name	REV	CAGE	DOC#	2 of 10
Your Company Name	ILL V	CHOL	DOCIII.	2 01 10
				Your Procedure #

### 1.0 PURPOSE

To prescribe the minimum procedures for the control of Customer Property according to the regulations outlined in the Federal Acquisition Regulation, Part 45.

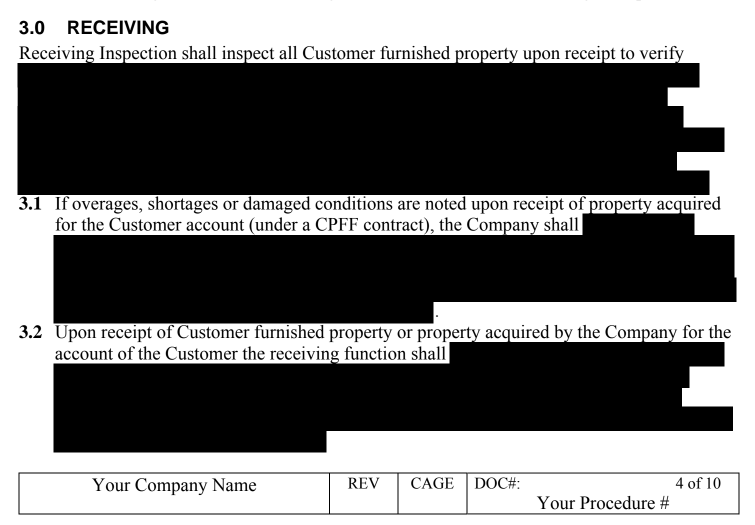
### 2.0 SCOPE

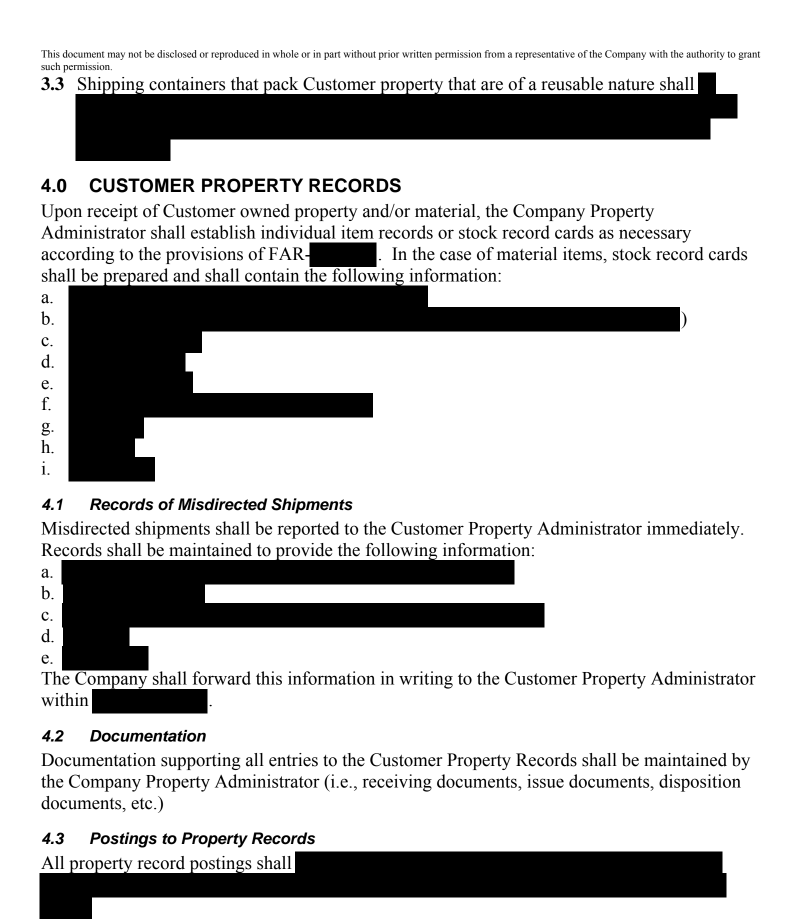
This procedure shall cover all property furnished to or acquired for use on contracts.

- a. Property Administrator means the individual duly designed by appropriate authority to administer the contract requirements and obligations relative to property. The person is an authorized representative of the Contracting Officer.
- b. Property means all property owned by or leased or acquired by the Customer under the terms of a contract. Property and contractor acquired property is defined as:
- 1. Property in the possession of or acquired directly by the Customer and subsequently delivered or otherwise made available to the contractor.
- 2. Contractor acquired property is property procured or otherwise provided by the contractor for the performance of a contract, title to which is vested in the Customer.
- c. Customer material is property that may be incorporated into or attached to an end item to be delivered under a contract or which may be consumed in the performance of a contract. It includes, but is not limited to, raw and processed material, parts, components, assemblies and small tools and supplies.
- d. Special Tooling means all dies, jigs, fixtures, molds, patterns, taps, gauges, other equipment and manufacturing aids and replacements thereof, acquired or manufactured by the contractor for use in the performance of a contract, which are of such specialized nature that, without substantial modification or alteration, their use is limited to the development or production of particular services (does not include consumable property, special test equipment or buildings, non-serviceable structures, general or special machine tools or similar capital items).
- e. Plant Equipment means personal property of a capital nature (equipment, vehicles, machine tools, test equipment, furniture and accessory and auxiliary items, excluding Special Tools and Special Test Equipment) used or capable of use in the manufacture of supplies or in the performance of services or for any administrative or general plant purpose.
- f. Scrap means property that has no reasonable prospect of being sold except for recovering value of its basic material content.
- g. Salvage means property recoverable for further use which because of its worn, damaged or deteriorated, incomplete condition or specialized nature, has no reasonable prospect of sale or use as serviceable property without major repairs or alterations, but which has some value in excess of scrap.
- h. Custodial Records means written memoranda or identifying method of any description or the type used to control items issued from Tool Cribs, Tool Rooms, Stockrooms, etc., such as requisitions, issue hand receipts, took checks, stock record cards or books and the like.

Your Company Name	REV	CAGE	DOC#:	3 of 10
				Your Procedure #

- i. Individual Item Record means a separate card, form or document, used to account for one item of property.
- j. Stock Record means a perpetual inventory form for recording quantities and types of items received, in stock and issued to requesters against a specific contract. The form serves as a posting reference and records the balance of stock items on hand and their unit prices.
- k. Discrepancies Incident to Shipment means all deficiencies incident to the shipment of Customer property to or from a contractor or vendor's facility, Customer depot or like source wherein differences exist between the property said to have been shipped and the property actually received are identified as discrepancies incident to shipment. These deficiencies include, but are not limited to, loss, damage, destruction, improper status and condition coding, error in documentation, i.e., identity or classification and improper status and consignment or unit not furnished.
- 1. Work-in-Process is the definition used for the purpose of financial reporting and covers material which has been released to the production element.
- m. CPFF Material, Contractor procured CPFF material is property purchased by the contractor, acting as agent for the Customer, for use in connection with a specific cost-plus type contract. This material becomes Customer property upon receipt and acceptance by the contractor.
- n. Bonded Storage means a secure storage area with access limited to designated personnel.





Your Company Name	REV	CAGE	DOC#:	5 of 10
				Your Procedure #

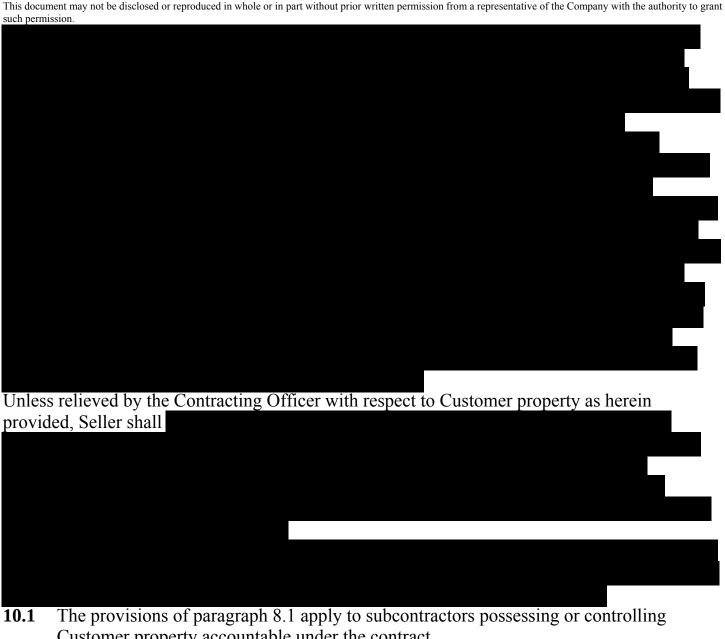
This document may not be disclosed or reproduced in whole or in part without prior written permission from a representative of the Company with the authority to grant such permission.  5.0 MATERIAL REQUISITION/ISSUE
5.0 MATERIAL REQUISITION/ISSUE  After receipt of Customer furnished material and preparation by the Company Property  Administrator of the required stock record cards the material shall be
<ul><li>5.1 Sensitive material issued according to 5.0 shall be maintained in a secure area with access limited to authorized personnel.</li></ul>
6.0 UTILIZATION
It is the responsibility of the Company Property Administrator to assure
7.0 MAINTENANCE
The Company Property Administrator shall insure
8.0 PHYSICAL INVENTORIES
Inventory, as used in this procedure, consists of sighting, tagging or marking (when considered necessary), describing, recording and reporting the property concerned and reconciling the property recorded and reported with the property records.
The personnel who perform the physical inventory shall not be the same individuals who
maintain the property records or have custody of the property.

Your Company Name

REV CAGE DOC#: 6 of 10

Your Procedure #

This document may not be disclosed or reproduced in whole or in part w such permission.	rithout prior writ	ten permission fi	rom a representative of the Company with the authority to grant
<b>8.1</b> The Company shall investigate and rall cases of loss, damage or destruction completed/	•		1 0
The report shall contain at a minimum: A. B. C. D. E. F. G. H.			
9.0 DISPOSITION			
At the completion of a contract under whe shall perform an inventory to determine is remaining. If residual items or scrap rem	f any resid	dual items	s of Customer property or scrap are
10.0 SUBCONTRACT CONTROL  The Company purchasing function shall is subcontracts or vendor purchase orders we furnished to the subcontractor or vendor: Responsibility for Property			•
Your Company Name	REV	CAGE	DOC#: 7 of 10 Your Procedure #



Customer property accountable under the contract.

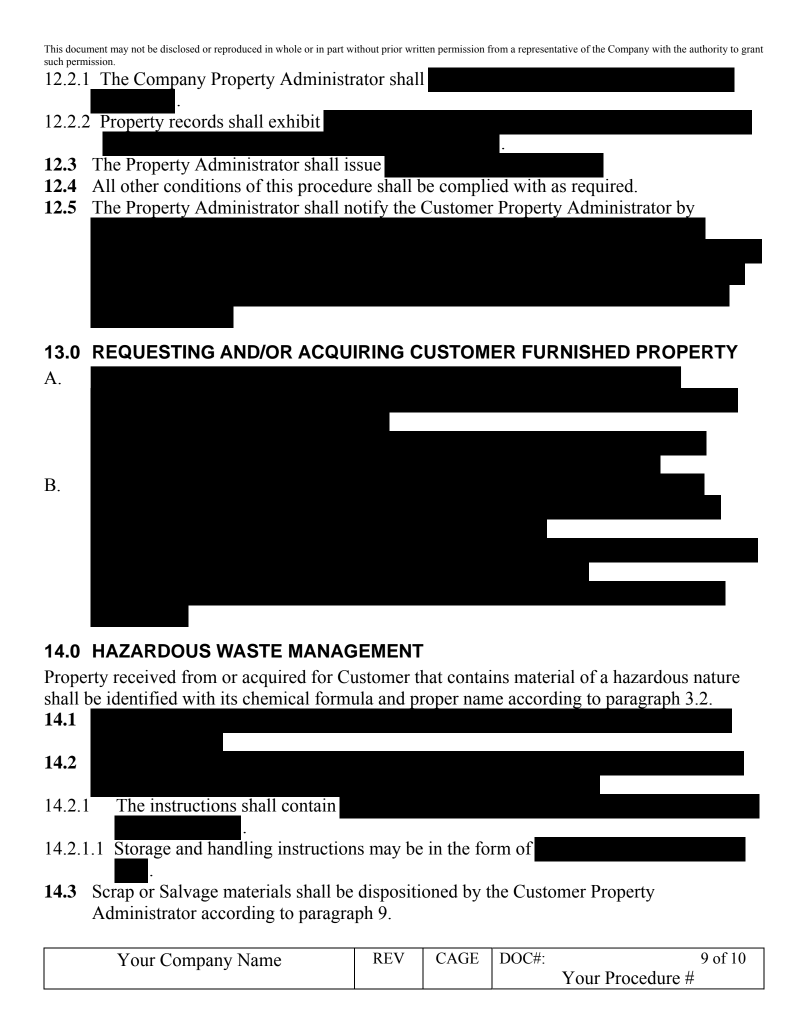
### 11.0 REPORTS

Reports shall be prepared by the Property Administrator according to the terms of individual Customer contracts.

### 12.0 PRECIOUS METALS,

- 12.1 Immediately upon receipt Receiving Inspection (R&I) shall inspect material according to para. 3 - 3.2.
- 12.1.1 Sensitive material shall be stored in bonded storage immediately upon acceptance or rejection by R & I.
- 12.2 The Company's Property Administrator, upon taking possession of accepted sensitive material, shall

Your Company Name	REV	CAGE	DOC#:	8 of 10
			Yo	our Procedure #



### 15.0 WORKMANSHIP

Adherence to applicable federal, state, local and environmental, health and safety requirements is mandatory.

Your Company Name	REV	CAGE	DOC#:	10 of 10
The state of the s				Your Procedure #

Metrology Recall Card

		1,1	$\frac{1}{2}$				
Des	scription:			Calib Fr	equency:		
Type:			Model:		S/N:		
Prop	erty ID#:						

QC-116-1 (mo/yr)

Instrument and Case Identification Tag (shrink to fit)

Tool #:	Tech:

QC-116-2 (mo/yr)

Instrument Deviation Tag (shrink to fit)

Tool#:	
Tool	Standard
Value	Value
Tech:	
	QC-116-3 (mo/yr)

**Measuring and Test Equipment Calibration Report** 

Department: Equipment: Size-Range:  Mfg-Model:  March and March an	In Toleran	ce as Received	Out-of-To	olerance as Recv'd	PO# for M&TE:	
Equipment: Size-Range: Mfg-Model:  Mfg-Mod		10001100	0 4.0 01 1		1 0 11 11 11 11 11 11 11 11 11 11 11 11	
Size-Range:  Mig-Model:  Mig-M	Equipment:			Location:		
	Size-Range:			Mfg-Model:		
Remarks:						
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QC-116-4 (mo/yr) Front

### IMPACT ANALYSIS REPORT

Number of parts that may be out-of-spec – List Model # and projected quantities for each type
that
± tolerance range for each dimension checked with the out-of-spec equipment – list by P/N
toronate range for each annichable through the control of the cont
<u>.</u>

QC-116-4 (mo/yr) Back

Your Pro Rev: (mo	cedure # /yr)		[Title] Calibration Instruction Sheet	QC-116-5 (mo/yr) Page 1 of
Spe	cial Instru	actions:		
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### LLC. All rights reserved worldwide. **Calibration System Policies and Procedures**

	Revisions					Rev:	
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### 1.0 Scope

These procedures comply with the re-	equirements of	Measuring instruments
are calibrated, at a temperature of	and	relative humidity, in the
F	For cases where calibration	must be conducted in the
production area, stabilization time is	also allowed.	

### 2.0 Definitions

- a) Gages are precision devices that compare the characteristics of an item to specified requirements.
- b) Recall All gages require recertification at established intervals. Recall dates are identified by a month/year designation. Certification is performed no later than the last day of the month/year designation except as otherwise provided. All gages may be used for acceptance/rejection of product during the month/year recall interval.
- c) M&TE Measurement and test equipment
- d) Standards Accepted values of natural physical constants or values traceable to National or International Standards.
- e) Procurement of Gages Gages are procured from a qualified source and are inspected by Gage Inspection before use. A newly acquired measuring or test device that has been certified as calibrated, and whose certification indicates an NIST reference number, may be issued to the user activity after a calibration interval and records have been established.
- f) Special Equipment (Your Co) standards, instruments, chemicals, and tools for which a measurement standard is not available on-site to perform calibrations.
- g) Significantly out-of-tolerance An instrument's accuracy that exceeds the manufacturer's published limits.
- h) Adequacy Adequacy, range, resolution and stability of M&TE and standards is determined by quality characteristic measurement requirements on an individual basis.
- I) Accuracy Ratio 10:1 for linear, weight, current, and voltage transfer standards.

### 3.0 Procedures

### 3.1 Identification

When a gage does not provide its own serial number then a number is issued. The numbers run consecutively for each gage size and may be further identified under a type-coding system. This number is etched or otherwise imprinted upon the gage.

### 3.2 Storage of Gages

All company owned gages are kept clean and are stored in cabinets and bins in the inspection department, tool crib or other storage areas when not in use.

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3.3 Recall
A rotating card file system is maintained on all instruments. The form used is QC-116-1.
The rotating card file provides the means for implementation of recall for any gage that has
expired its certification period.
Portable gages are physically removed from service and recertified during the recall interval as
time permits. Permanent gages are
3.4 Working Record
In addition to the card file system, a working record sheet, QC-116-4, is kept on each company-
owned gage/standard. The purpose of this record is to
3.5 Calibration Frequency
Calibration intervals are based on the following criteria:
·
Calibration intervals are established in terms of
and the schedule of Table I.
Tools that are identified as "Spares" in the calibration database are calibrated based upon usage rather than time and a usage tag is exhibited on the tool or its case. A "Spare" tool is calibrated
after it

Your Company Name	REV	CAGE	DOC#:	4 of 9
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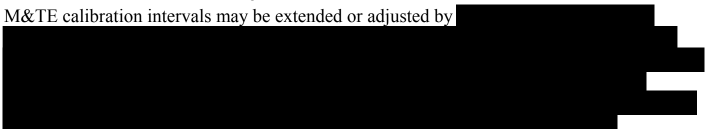
### **TABLE I, Calibration Intervals**

Calibration Cycle	Recalibration Cycles to Qualify for New Calibration Cycle	New Calibration Cycle
Annual		
Bi-Annual		
3 - 4 Years		
5 Years	N/A	

### 3.6 Interval Adjustment

M&TE whose calibration error is recorded as being greater than the last recorded calibration error, but not significantly out of tolerance, reverts to

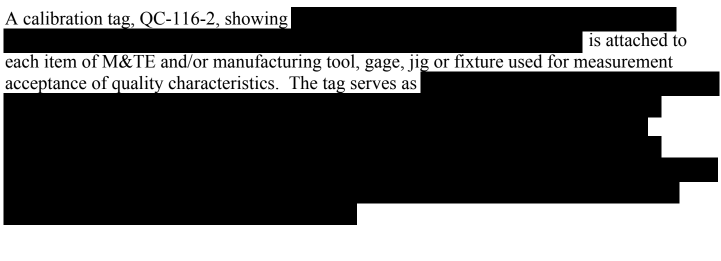
### 3.7 Interval Extension / Adjustment



### 3.8 Calibration Overdue

Overdue items are prevented from use as practicable. A calibration overdue notice in the form of an inter-office memo or other format may be used to facilitate recall of portable gages.

### 3.9 Calibration Identification

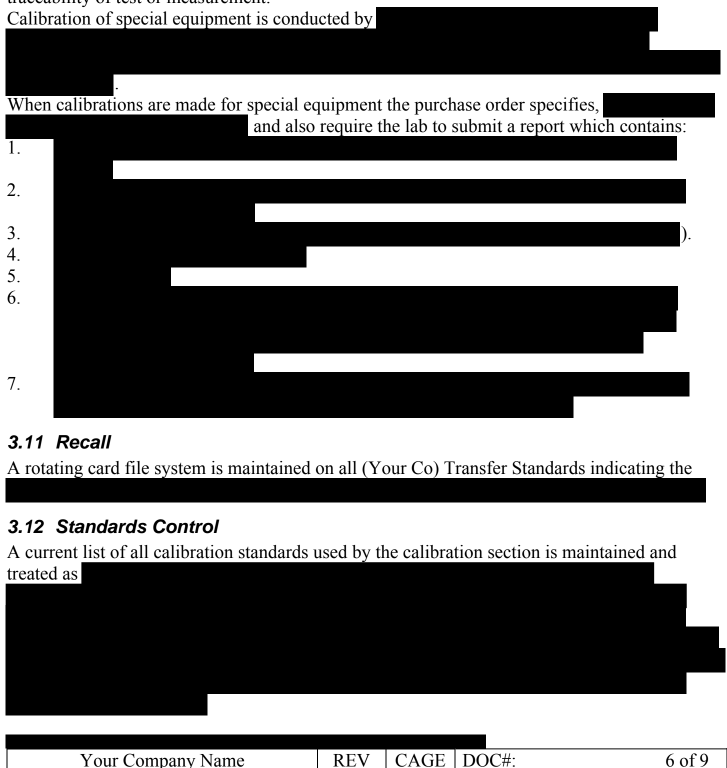


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### 3.10 Calibration Standards/Special Equipment

It is the position of the National Conference of Standards Laboratories (NCSL) that:

"Test report numbers issued by the NIST are intended to be used solely for administrative purposes. Although they are often used to uniquely identify documents which bear evidence of traceability, test report numbers should not be used nor be required as proof of adequacy or traceability of test or measurement."



Your Procedure #

Equipment and tooling found to be significantly out of tolerance, damaged, inoperative, erratic or exhibiting some other form of anomalous condition should be

M&TE found significantly out of tolerance at recalibration for from use (except as otherwise provided) by

All out of tolerance data is utilized in an evaluation to determine the adequacy of the M&TE for the intended use and to determine the effectiveness of the calibration procedure and measuring or test procedure. A notice is prepared and

### **3.15** Provision for Use of Out-of-Tolerance Equipment (apply sparingly)

An instrument whose calibration error is significantly out-of-tolerance (over a short portion of a specified range) is returned to service only when

### 3.16 Suspected Product Nonconformance

Any product certified with M&TE subsequently found to be out-of-tolerance is immediately reported to the Customer. "The impact on quality of products examined or tested by equipment found to be out-of-tolerance during calibration will

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### 3.17 Traceability

Inspection instruction sheets and manufacturing travelers specify measurement and test equipment utilized for product conformance inspection. The M&TE number is recorded on

### 3.18 Production Tooling Used as Media of Inspection

Any production tooling which is used to accept attributes of a part, sub-assembly or assembly is verified for accuracy prior to its use and

### 3.19 Employee Owned Tools

Personal Tooling or gages owned as personal property by employees of (Your Co) are

### 3.20 Subcontractor Calibration

The quality requirements outlined in Supplier Quality Requirements QC-117 are imposed to the level required by the (Your Co) Quality Group. Criteria for the selection of the inspection level are based on

### 3.21 Storage and Handling of M&TE

M&TE is handled during movement using the manufacturers recommendations or handling practices that prevent exposure to

- except that which is normally encountered during movement -- and

M&TE requiring transportation to a calibration laboratory is packaged to

3.21.1 Calibration Prior to Archive / Long-Term Storage

An instrument does not require accuracy verification prior to archive / long-term storage if it was

### 3.22 Setting / Selecting a Reference Standard

Rule: The measurement range of a device being checked for accuracy must be less than the maximum measurement range of the reference standard – see the following examples.

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### **VOLTMETER:**

A voltmeter that is required to be calibrated shall be verified for accuracy within an equivalent range on the reference standard, e.g.,

The voltmeter reference standard has scales that range from 2-20V, 20-200V, etc. – the voltmeter being checked for accuracy must be set to the same range as the reference standard – the reference standard must be set to a range that brackets the same range as the voltmeter being checked for accuracy, i.e., if the voltmeter being checked is set to 2-20V then the standard must be set to the same range – do not use the 20-200V range on the standard to check the 2-20V range on the voltmeter being checked for accuracy.

CURRENT SHUNT: The measurement range of a reference standard shunt must not be greater th measurement range of the shunt being checked for accuracy, e.g., a 100A cureference standard can be used to calibrate a but the standard to calibrate a	
OTHER MEASUREMENT DEVICES: Any reference standard whose maximum measurement range is the same as checked for accuracy must be at least more accurate than the device e.g., a device being checked has a 1% tolerance then the reference standard is tolerance.	e being checked,

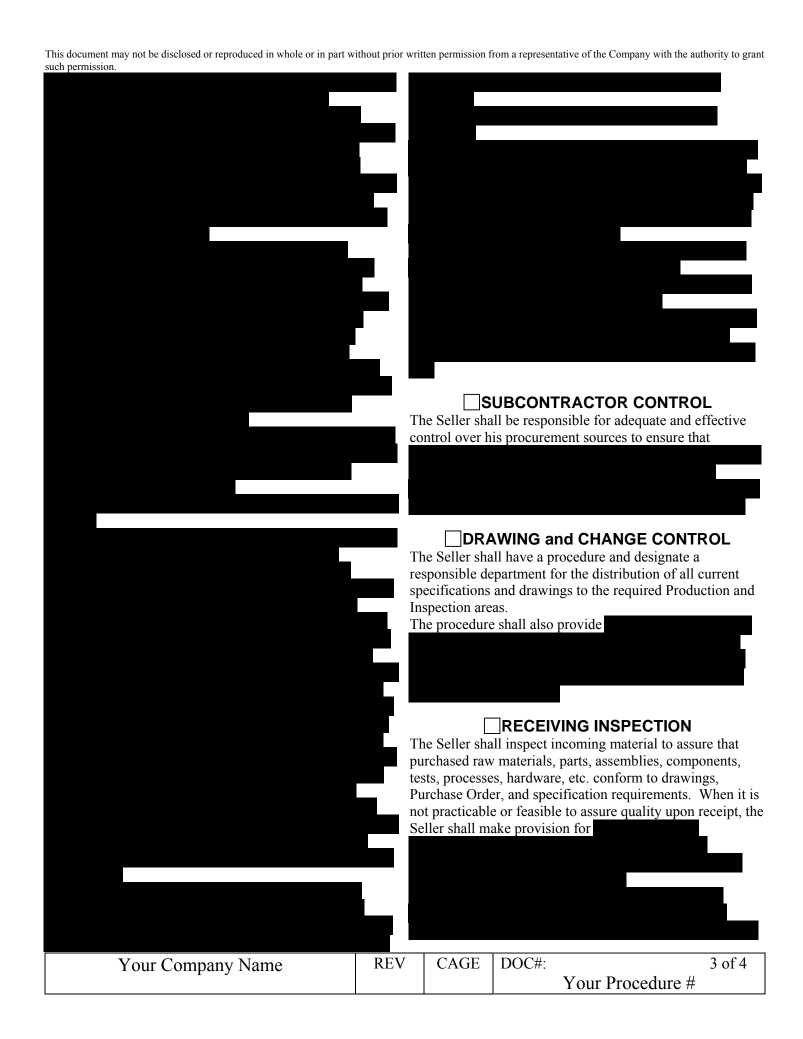
Your Company Name	REV	CAGE	DOC#:	9 of 9
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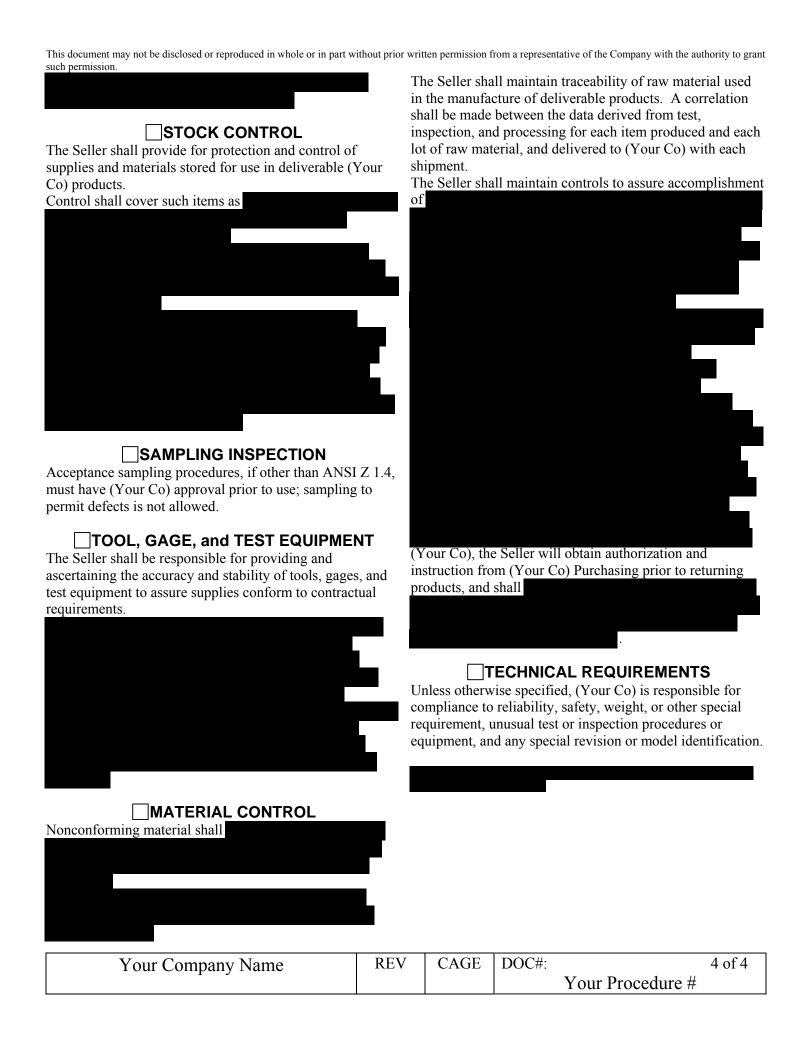
# Supplier Quality Requirements (No/Yr Mo/Yr All rights reserved worldwide)

Rev: E.O. Number Description Letter Date Contract#: **Your Company Name** Prepared By: Date Your Dept: Date **YOUR PROGRAM** Your Dept: Date Your Dept: Your Procedure # Date Your Dept: CAGE: Date Size: 1 of 4 Your Form # (mo/yr)

This document may not be disclosed or reproduced in whole or in part without prior such permission.	written permission from a representative of the Company with the authority to grant
To establish the minimum requirements for supplier Quality Systems necessary to ensure that materials, parts, components, and services meet the requirements of the Contract. Procedures used to implement the provisions of this requirement shall be subject to (Your Co) approval upon request.  APPLICABILITY  These requirements shall apply to all supplies and services when referenced on the Purchase Order and amendments thereto.  When (Your Co)'s Purchase Order includes Seller's Inspection System QC-117 Level I, as a requirement, Seller's contractual commitment for an Inspection System shall be defined by all paragraphs of this specification.  When (Your Co)'s Purchase Order indicates Level II as a requirement then the Seller's contractual commitment for an Inspection System shall be defined only by those paragraphs of this specification which are checked-off.	PROPRIETARY INFORMATION  The Seller must identify in writing the intended use in performance of the Purchase Order of an item, material, component or process with respect to which access by (Your Co) or (Your Co) Customer representatives for purpose of Quality Assurance by inspection, test or process surveillance is proposed to be restricted.  The absence of such written identification is a representation by Seller that
DEFINITIONS and ABBREVIATIONS  A. The term 'Buyer' or '(Your Co)' means (Your Co).  B. The term 'Seller' means the legal entity that is the contracting party with the Buyer with respect to the Purchase Order.  C. 'IAW' means in accordance with.  D. 'MRB' means Material Review Board  SELLER'S QUALITY SYSTEM, GENERAL  The Seller shall maintain an effective Quality System planned and developed in conjunction with his other functions to comply with contractual requirements. In order that the Quality System will be effective, it shall provide	PROCESS CONTROL  The Seller shall provide for complete review of contract requirements at the earliest practical phase of contract performance to
Records shall be kept available for  NEGOTIATIONS  It is not the intent of this specification to restrict the Seller in his mode of operation; therefore, it is possible that certain items herein may be subject to negotiation. Until such time as the subject of the negotiation is resolved, the Seller is	The Seller shall develop an Inspection/Test Plan specific in nature and related directly to the hardware produced.  The Plan shall

Your Company Name	REV	CAGE	DOC#:		2 of 4
				Your Procedure #	





### **BASIC CONTRACT REVIEW**

Program Name: Program Source:		
Program Source:		
RFP#:	Contract Type:	
	, <u>, , , , , , , , , , , , , , , , , , </u>	
	<u> </u>	
Da	ate:	Date:

QC-118 (mo/yr)

Program:   Rev mo/yr   Customer P/N-Rev:   P.O.# & Rev:	,	Your Company Name			MFG/QA TRAVELER Your Title					QC-119 (mo/yr) Page 1 of 3	
Program: Account#: Customer: SPECIAL INSTRUCTIONS:  FO APPROVAL:   FO#   Data List# & Rev.   ECP#;   DPROVAL:   FO#   Perform ata List# Vo# & Rev.   Perform test operation is transfer of testing the Vo# & Rev.   ECP#;   DPROVAL:   FO#   Perform test List# Vo# & Perform List*   Data List#   Data List#   Data List#   DPROVAL:   FO#   Perform test closed to Alta Vo#   Perform List*	Your #	<u>‡</u>	Rev mo	o/vr		ev.		<u> </u>			
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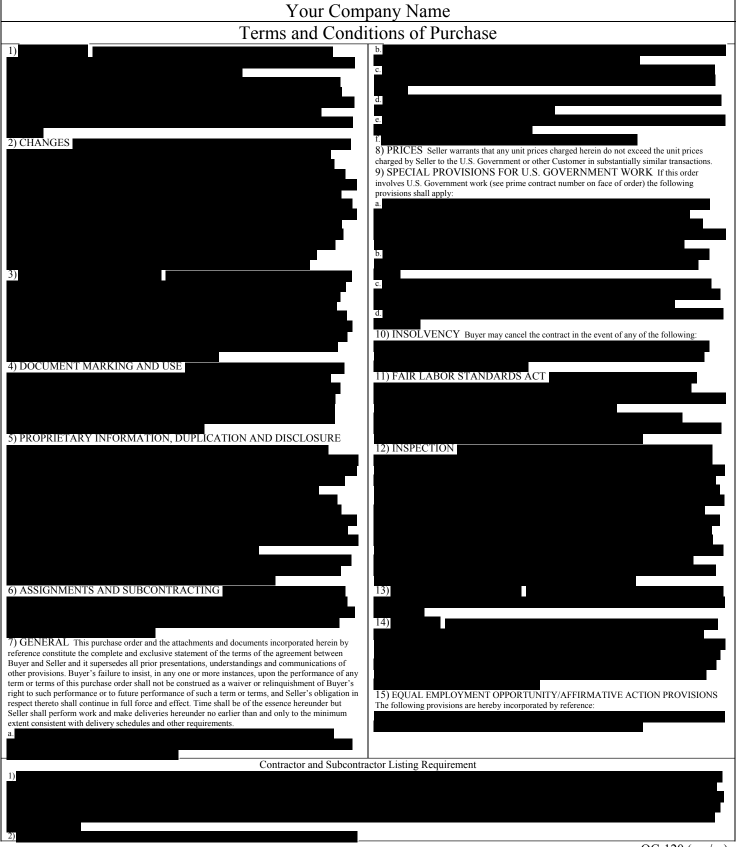
DEFINITIONS: P/S=Packing Slip PR=Product Report PS=Product Specification
IAW=In accordance with A/R=As required CEI=Contract End Item EIDP=End Item Data Package
BOL=Beginning of Life PPP&M=Preservation, Packaging, Packing and Marking

Company Name MFG/QA TRAVELER			QC-119 (mo/yr) PAGE 2 of 3				
			Your #			IAC	JE 2 01 3
OPER	DEPT	Description of Task		SIGN	MR – ECP - ACN	Date	Gage
115	QC	Disposition rework IA accurately recorded.	W Your #. Verify rework observations are				
120	ENG &TECH	Select item for DPA to operation.	est & record S/N in accepted column of this				
122	TECH		one sample from approximately 50 items IAW				
			e assembly to temperature in excess of 75°F.				
		Forward components t					. •
125	QC	Review DPA test data documentation to trave	provided by engineering & attach supporting eler.			:106	)
127	QC	Prior to ATP: Ve	erify test tolerances reflect CEI BOL		7.	71,	
			ipment tolerance, and a test flow		70		
		_	nted for each test. Verify Customer		.01,		
		-	rocedure is in contract file-Consult		dworld		
		QAM	PART III		01		
128	TECH	Computer program: N		- 3		1	
128.1	TECH	Computer program: N	mechanical connection to each item for	250			
120.1	ILCII		ly check each mechanical connection from the	25			
			ta acquisition equipment for tightness, i.e., TC's,	0			
			ads, torqued threads, etc.				
130	TECH	Perform test IAW You	ır#.				
135	QC	Verify test IAW Your					
140	TECH	Perform test IAW You	ır#.				
		Copyright © JnF Specialties plan.com/copyright.htm	s, LLC. All rights reserved worldwide, www.quality-control-				
145	QC	Verify test IAW Your	#.				
150	TECH	Perform test IAW You					
155	QC	Verify test IAW Your	#.				
160	TECH	Perform test IAW Your	*				
165	QC	Verify test IAW Your					
170	TECH	Perform test IAW You					
175	QC	Verify test IAW Your					
180	TECH	Perform test IAW You					
185 190	QC TECH	Verify test IAW Your Perform test IAW You					
195	QC		op 190 and compliance to Notes supplied with				
173	QC		by contract - consult QAM.				
197	QC		ection for ATP data review A/R.				
199	CUST		ection for ATP data IAW contract				
		requirements.					
		* (A)	PART IV				
201	QC	Verify tests have been notify Engineering for	completed prior to next Op. If not completed,				
203	QG		& record results on Your #.				
205	QC		ource for Monitoring.			1	
207	CUST		ections IAW Contract directives.			1	
220	TECH	Weld IAW Your #. R	ework A/R IAW Your # & record accurate				
)		disposition on this trav					
225	QC		ur #. Test weld IAW <b>Op 25</b> ; record results on				]
			k was performed IAW Your # & disposition was				
		accurately recorded or	i mis traveier.				
COM	MENTS:						

Company Name			MFG/QA TRAVELER			QC-119 (mo/yr)	
			Your #			PAGE 3 of 3	
OPER	DEPT	Description of Tasl	(	SIGN	MR – ECP - ACN	Date	Gage
230	TECH	If radiographs were no	ot produced prior to Op Your # then radiograph				
			red by contract - consult QAM.				
235	QC	Perform dimensional i					
		Record dimensions on					
237	LAB		diographs IAW Your #unless Level II				
			d by contract; forward radiographs to Supplier				
247	00	A/R for certified exam					
247	QC Perform final test on items IAW <b>Op 25</b> , Your #, & record results on Your #.						
250	DDOD						
250 255	PROD QC	Clean items IAW Your # (provide QC with first article for marking).  Perform final visual inspection IAW Your #.					
260	PROD	Prepare items & package IAW Your # & contract unique PPP&M					
200	FROD		vable special handling tag may be required for				
		each unit pack - cons					
265	QC		packaging IAW Your # & PPP&M instructions.				
203	Q C		n or damaged at packaging.				
267	QC		data package: (Blank lines are N/G on forms)				
			ce & Cert. of Authenticity.				
		2. Pre-ATP & ATP t					
		3. Item weights.					
		4. Flowcharts from a					
			ckage IAW Customer Format - consult QAM				
			of the EIDP specifies the CEI name, P/N &				
			listing when appropriate)				
			AW Customer Format - consult QAM				
		Format consult	le data package and shipping list IAW Customer				
			have been resolved prior to final acceptance.				
271	QC		deliver with end item data package.				
2/1	QC		gbook for all materials, components, and sub-				
			opy of traceability logbook for the final logbook.				
273	QC		ource Inspection for FINAL ACCEPTANCE.				
275	CUST		ection IAW contract requirements.				
		Acceptance is defined	d as approval of the end item data package				
		and the contract end					
280	PROD		packages, & mark carton & crate IAW Your # &				
		contract unique PPP&					
285	QC	, i	mine all cartons & crates for marking IAW Your				
			ons. Verify 2 copies of the data package are in	1			
			e need for crate handling procedure inclusion	1			
			be maintained at 32° to 68°F if items are stored in	1			
			f items are to be stored more than 2 weeks they t 32° to 40°F - <b>consult QAM.</b>				
295	QC		for 20 years contact DCC for assistance.				
493	<u> </u>	1 repare and store data	101 20 years contact DCC 101 assistance.	<u> </u>	1	L	

COMMENTS:		

PURCHASE ORDER	Date:		
Your Company Name	Purchase Order #:		
Phone: xxx-xxx-xxxx Fax: xxx-xxx-xxxx			
Address, City, State, Zip Code			
If a Prime Contract # is entered hereon, this procurement is			
Cumpliane	~		
Supplier:	Ship To:		
Phone#:	DDAC D 4 1		
	DPAS Rated:		
	Purchase Order	Amount:	
			and/or
			C-120 (mo/yr)



# LLC. All rights reserved worldwide. **Purchase Order Review**

Rev: E.O. Number Description Letter Date Contract#: **Your Company Name** Prepared By: Date Your Dept: Date **YOUR PROGRAM** Your Dept: Date Your Dept: Your Procedure # Date Your Dept: CAGE: Date Size: 1 of 1 Your Form # (mo/yr)

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1	Quality Group	<ul> <li>The reviewer determines the need for, and if justified, imposes the requirements of QC-117, Supplier Quality Requirements, to the Requisition or P.O.</li> <li>Complete the Used-On and Contract# sections on the cover page of QC-117         Used-On = J/N or Program Acronym; Contract# = P.O.#     </li> <li>Check-off applicable requirement boxes on Requisition</li> </ul>
2	Quality Group	Add known QA requirements to the requisition for entry on the PO; such as letter survey to Suppliers to determine their conformance to MIL-STD-45662 or ANSI/NCSL Z540-1;
	IF	THEN
2.1	Older Revision Supply Required	ITIEN
2.2	Requisition is marked "Under Revision"	
2.3	A Raw Material Requirement <b>is not</b> Specified	 
2.4	Deviation to drawing is noted on Requisition such as "Less Note" Deviation to drawing is	

Your Company Name	REV	CAGE	DOC#:	2 of 2
1 3			You	ır Procedure #

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ussion.		
	noted on Requisition such as "Less Note"	
2.5	Order is for production but doesn't reference engineering drawing #	Copy the PO to Drafting with comment to produce drawing A/R; This provision is not applicable to commercially available supplies
3	Quality Group	Add provisions for any one or combination of the following to the Requisition or P.O. when justified:
		]
		Certification that the delivered goods conform to the procurement
		document requirements

Your Company Name	REV	CAGE	DOC#:	3 of 2
1 3			,	Your Procedure #

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ermission.		control
4	Quality Group	Relative to the procurement of software, the reviewer determines the need
-	Quality Group	for, and if justified, adds to the procurement document provisions for any
		one or combination of the following:
5	Discrepancy in	
	Requisition or P.O.	
5.1	Supplier Quality	
	Requirements applies	Copy to R&I
5.2	P.O. requires additional	Record supplier related add-on text to Requisition or P.O.
	conditions related to	
	supplier	
	IF III	THEN
5.2.1		Record add-on text to Requisition or P.O. and forward to User
	conditions related to in- house processing	
5.2.2	Requisition or P.O. Ok	When R&I QC is required, sign and forward <i>PO's in numerical order</i> to
3.2.2	requisition of 1.0. Ok	R&I (Procurement Technician must be cognizant of all purchases)
-	Ovality Cassa	Engrand Subsenting to Evaluation Overticans in to the Sumulian and the
6	Quality Group ISO 9001 Applies	Forward Subcontractor Evaluation Questionnaire to the Supplier; perform required follow-up routines (Your #).
L	190 Just Whites	required follow-up routines (10th #).

Your Company Name	REV	CAGE	DOC#:	4 of 3
1 3				Your Procedure #

#### (Your Company Name) Dimensional Analysis Record

Item Name:	<b>Customer:</b>	
<b>Drawing Number:</b>	Inspector:	
	_	
Key Parameter	Key Parameter	

Your Logo QC-122 (mo/yr)

(Your Co Name) DATA LIST	DATA LIST REVISION:		PROJECT:		Page 1 of
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#### FEDERAL, MILITARY and SOCIETY SPECIFICATIONS

SPECIFICATION NUMBER	REV	DESCRIPTION

Use latest revision at the time of contract, or as specified by contract

A/D = As Designed; A/B = As Built; or use A/T = As Tested

\* An asterisk placed in the revision column indicates a tabulated drawing. Use the latest revision of the tabulated drawing at the time of contract.

#### SUMMARY OF DATA LIST REVISIONS

D/L REV	DOCUMENT AFFECTED	E.O.#	E.O. DATE	D/L REV	DOCUMENT AFFECTED	E.O.#	E.O. DATE

QC-123 (mo/yr)

Ref:		Your Co	ompany Name					
Page 1 / of /		SURVEY REPORT						
Project:		Place:						
1 Toject.		Tidee.						
Subsystem:		Date:						
Satisfactory:								
Satisfactory v	with							
Satisfactory	with							
Rejected:								
Writer:								
Participant:	Manufacturer	Contractor	Prime	Customer				

Page 2 / of / SURVEY REPORT Continuation			Your Company Name
Page 2 / of / Continuation			SURVEY REPORT
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As Designed / As Built.	As Designed / As Built.		
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Ref:	Your Company Name	
	SURVEY REPORT	
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Page 3 / of /		
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#### **CERTIFICATE OF COMPLIANCE**

From:	
To:	NOTICE
	NOTICE
Attention: Receiving Inspection	
PO#:	THIS CERTIFICATE
	OF COMPLIANCE
Customer P/N:	
	MUST
Your Co P/N:	

Form Rev: Orig

Your Logo



# GENERAL REQUIREMENTS

Origination Date: XXXX

Document Identifier:	Name, Number, Unique ID
Date:	Latest Revision Date
Project:	Customer, Unique ID, Part Number
Document Status:	Draft, Redline, Released, Obsolete
Document Link:	Location on Server (if used)

#### Abstract:

This document describes general manufacturing and interpretation requirements.



#### Your Company Name

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#### **REVISION LOG**

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#### 1. SCOPE

This document describes general requirements and methods of interpreting engineering requirements specified in specifications and/or drawings.

#### 2. THEORY

The space available on product drawings sometimes limits the opportunity for a clear description of engineering requirements. Details that the engineer would like to include on the drawing are sometimes left off with the assumption that the User will understand what is meant within the notation(s). These assumptions are valid only when the User is the engineer; otherwise, concise training of personnel is required.

#### 3. REFERENCES

ANSI B46.1, Surface Roughness ASME B1.1, Unified Inch Screw Threads ASME B18.2.2, Square and Hex Nuts ASTM E 29, Significant Digits FED-STD-H28, Screw Thread Standards for Federal Services

#### 4. REQUIREMENTS

#### 4.1 Order of Precedence

In the event of conflicting requirements the following order of precedence governs: The Customer's requirements always supersede Company requirements unless approved by the

The Customer's requirements always supersede Company requirements unless approved by the CCB.



Military or Society Procedures or Standards

#### 4.2 Significant Digits

Calculations may be performed with a greater number of significant digits than shown on the applicable drawing; however, <u>measurements and calculations</u> must be *reported* to the same number of significant figures as specified by the applicable drawing. For instance,

Your	Logo

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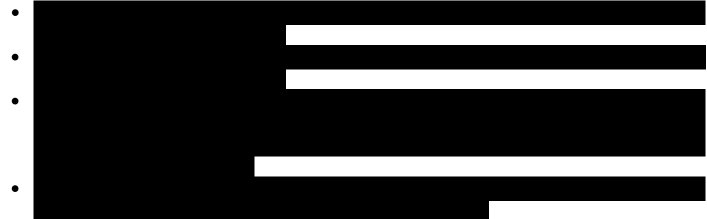
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however, the drawing specification requires reporting the calculated value to no more significant digits than shown.

## 4.3 Determining Conformance; Absolute Method or Rounding Method (ASTM E 29)

Unless otherwise specified by the CCB, the 'rounding method' is used for determining compliance of test data to product specifications according to ASTM E 29: (some contracts specify "NO ROUNDING — TOLERANCES ARE ABSOLUTE") Rounding is performed as follows: [quoted from ASTM E 29]



#### Example — 0.0<u>21</u>"±0.001"

0.0211" to 0.0214" *must* be rounded down to for 2 significant digit specifications; 0.0215" to 0.0219" *must* be rounded up to for 2 significant digit specifications; (unless the last retained digit from the rounding method is even, e.g., 0.0225" is rounded down to 0.022" since the last digit retained is even, while 0.0226" is rounded up to

#### Example — 0.0<u>215</u>"±0.0015"

0.0211" to 0.0219" must be applied as observed - no rounding is possible if the measurement equipment can only read to the 4th decimal place. If the equipment is capable of reading beyond the 4th decimal place then round to the last digit retained as described herein paying particular attention to whether the retained digit is odd or even when followed by the numeral 5.

#### **Example** — **550**±**50**

499.1 to 499.4 *must* be rounded down to 499.5 to 499.9 *must* be rounded up to 600.1 to 600.5 *must* be rounded down to 600.6 to 600.9 *must* be rounded up to

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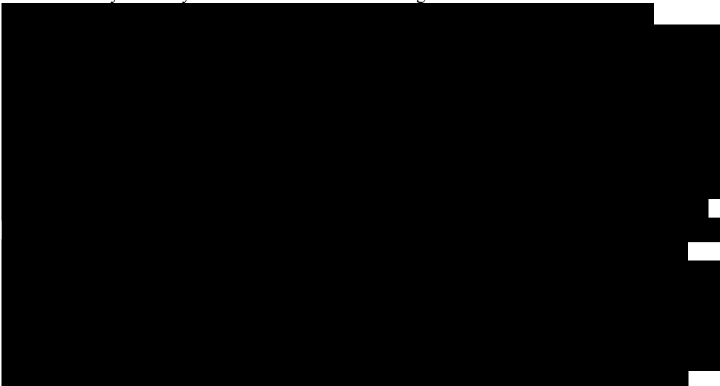
4.3.1 Equipment Tolerance

for 2 significant digit specifications; 0.0211" or 0.0214" **must** be rounded down to however, for 3 significant digit specifications, where instrument accuracy is specified as  $\pm 0.0001$ ", the original measurement figure of 0.0211" may actually be 0.0210" or 0.0212".

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The instrument tolerance can be used to affect 'rounding' by an amount equal to the plus or

minus accuracy stated by the instrument's calibration tag or record.



#### 4.4 Target, Goal and Should-Be Specification

Target, Goal and Should-Be specifications are suggested specifications, they are not fixed and compliance is more a judgment than a rule.

4.4.1 Application of the Drawing or Procedure Specification

Monitor data for compliance to the target value. When the product or process does not match the target value specified by the product drawing or process procedure,

#### 4.4.2 Target and Range Specification

When Range values are specified in addition to a Target value then product compliance to the Range values is

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#### 4.5 Potting and Encapsulation

#### 4.5.1 Engineering Drawing Note(s)

Potting and encapsulation operations may or may not be defined by a drawing note that references a manufacturing procedure that defines the lot formation and use of an epoxy log book for every mix.

#### 4.5.1.1 Application of Drawing Specification

Unless otherwise specified, potting and encapsulation materials whose shelf-life has expired must not be used on deliverable products unless authorized in writing by the MRB or CCB.

All potting and encapsulation materials are identified with a shelf-life expiration date.

The expiration date is sometimes modified by an annotation that applies an additional expiration date after the container is opened, e.g., 6 month shelf-life while un-opened and properly stored and then a 3 month shelf-life after opening the container. Prior to using any potting or encapsulation material, determine its shelf-life expiration date by

#### 4.6 Dimensional Requirements and Allowances

#### 4.6.1 Surface Flaws

Surface flaws include

Acceptance of parts having surface flaws shall be at the discretion of the REA and shall be based upon the function of the part.

#### 4.6.2 Free State Variation

If material flexibility or normal stresses can be expected to cause parts to be out of tolerance, appropriate inspection procedures shall be obtained from the REA prior to inspection of parts.

#### 4.6.3 Blind Holes

The drill point shall

#### 4.6.4 Gaging Hole Diameters

The diameter of a hole is within required limits when accepted by "GO" and "NOT GO" plug gages of appropriate size without reasonable evidence during plug gaging that the hole is out-



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of-round in excess of the diameter limits. Bell-mouthed holes are

#### 4.6.5 Hole Quality

The walls of holes shall be clean cut and shall present a uniform machined surface.

Hole edges shall be free from burrs and shall not

These requirements are subject to visual inspection only and are to be evaluated in terms that are

#### 4.6.6 Removing Burrs and Sharp Edges

All burrs and sharp edges shall be removed to the extent that material fragments are not visible and sharpness cannot be felt by using either a chamfer or radius. If it is necessary to break sharp edges or to deburr after application of chemical surface treatment, the bared metal shall be touched-up according to section herein named "Correcting Defects in Coating". Flash on molded plastic parts that does not cause the part to exceed maximum dimensional limits need not be removed. These requirements do not apply to rough and semi-finished

#### 4.6.7 Correction of Manufacturing Defects

#### 4.6.7.1 Permissible Corrections

Correction is permissible if

#### 4.6.7.2 Non-permissible Corrections

Corrective methods that add material to the product or that employ techniques abnormal to the production process are

#### 4.6.8 Correcting Defects in Coating

Defects in chemical organic and metallic coatings may be corrected by

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#### 4.6.9 Flat Surfaces

Where no parallelism tolerance is specified, flat surfaces of a part shown as parallel on a drawing shall be parallel within their limits of size. Flat surfaces of a part shown as perpendicular on a drawing shall be perpendicular within

#### 4.6.10 Thread Form

All threads shall be free from

Where no thread form is specified, threads shall comply with

Cold forming thread tools such as Besly "X-Press" may be used in lieu of metal cutting tools. A slight groove may appear along the thread crest as a result of the metal flowing action of a Besly tool and is acceptable if the overall thread crest height conforms to limits specified by and applicable specification sheet.

#### 4.6.11 Thread Gaging

Thread wires and measurement indicators may be used to accept thread dimensions.

When "GO" gages are used, the product shall allow the "GO" gage to enter or to be entered the specified full length or depth of the thread; however, the thread must be functional.

When "NOT GO" plug or ring gages are used, the product is acceptable when it does not enter the gage or there is

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4.6.12 Surface Roughness

When the surface rou	ghness specified	is less than	32 microinches,	measurement s	hall be
performed according	to			When the	surface
roughness specified is 3	32 microinches or	greater, visua	al and tactile comp	parison of the act	ual par
surface with commercia	al roughness com	parison specir	nens is acceptable	in lieu of	

## 4.7 Requirements for Cleaning, Protection and Identification of Raw Material, Parts and Assemblies

#### 4.7.1 Protection

All parts	and assemb	olies shall	be	adequately pro	tected fi	rom accumulati	on of	foreign	matte	er
corrosion,	physical	damage	or	deterioration.	These	requirements	shall	apply	to	

#### 4.7.2 Cleanup of Parts and Assemblies

All finished parts and subassemblies shall be adequately cleaned before final assembly. Final assembly and necessary subassembly shall be performed in an environment appropriate to the type of product. All parts and assemblies shall be thoroughly cleaned to remove foreign and manufacturing waste material such as:



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#### **DEFINITION OF TERMS**

Accuracy:	
CCB	Configuration Control Board
Goal Posts	A range that defines a minimum and maximum specification
GR&R	Gage Reproducibility and Repeatability
IAW:	In accordance with
Independent	Results obtained in a manner not influenced by any previous result on
Test Results:	the same or similar test object
MRB:	Material Review Board
Precision:	
REA:	Responsible Engineering Authority
Repeatability	
Conditions:	
D 4 1 '1'4	
Repeatability:	
Reproducibility	
Conditions:	
Reproducibility:	
	Population standard deviation is known
S:	Population standard deviation is estimated
Trueness:	

### **INSPECTOR STAMP LOG**

Signature	Password Signature	Initials	Date	

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# CONTROL OF NONCONFORMANCES

Origination Date: XXXX

Document Identifier:	Control of Nonconformances
Date:	Latest Revision Date
Project:	Customer, Unique ID, Part Number
Document Status:	Draft, Redline, Released, Obsolete
Document Link:	Location on Server (if used)

#### Abstract:

This document describes procedures for control of nonconformances.

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	Your Company Name	Rev: Orig

#### **REVISION LOG**

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#### 1.0 PURPOSE

This document defines and makes reference to the procedures necessary for the control of nonconformances.

#### 2.0 THEORY

Work that has failed inspections or tests or that in any way does not meet requirements are considered "nonconformances". Such work must be controlled to ensure it is not accidentally delivered or used. The Company's system ensures that nonconformances are identified when found and are segregated, investigated and dispositioned. Corrective and/or preventive actions are taken to ensure nonconformances do not reoccur.

#### 3.0 GENERAL PROCEDURE

- 3.1 "Nonconformance" is any work or raw material used by the Company or listed as a Customer complaint, such as:
- Acceptable inspection limits
- Acceptable test results
- Customer requirements (prints, specs, etc.)
- Design requirements (prints, specs, etc.)
- Material shelf life limits
- Statutory or regulatory requirements (safety, packaging, etc.)
- 3.2 Nonconformances must be withheld pending disposition by a completed Request for Support (RFS) or by direction from Quality. A Calculated Risk Release may also be used for disposition; however, the Calc-Risk must be closed before Customer acceptance.
- 3.3 All employees are empowered to engage this procedure when they discover nonconformances. No employee may work on yellow-tagged nonconformances.
- 3.4 Upon discovery of a nonconformance, an employee may make an attempt to perform immediate rework if such rework is within that employee's ability. For example, if an item requires sanding and the nonconformance appears to be insufficient sanding, the employee may continue to sand the item to bring it into conformance without any further action.
- 3.5 When an employee cannot bring the work into conformance through immediate rework, the employee
- 3.6 If an employee or supervisor cannot
- 3.7 The employee completes the top portion of the Request for Support form, filling in all pertinent spaces. The employee then submits the Request for Support (RFS) to Quality.

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report number on the tag. A yellow-tag. Whenever possible the work should be as 3.9 Upon receipt of the Request 3.10 Quality will then assign the Result and Corrective Action Request the Request for Support. For more of 3.12 Quality will also indicate on the configuration change is required, etc. 3.13 The RFS is submitted to MRB actions that affect configuration.	nonconforming work with a yellow no ag may be used without a Request for be physically segregated from other we for Support, the Quality representative eport to an appropriate authority for recertained or estimated to be the fault (CAR) to the Supplier. In such cases a the CAR system see the Corrective and the Request for Support form if a document of the support form if a docu	nconformance tag and indicates the Support for temporary identification. ork.  We will  esolution. This includes  of a Supplier, Quality may elect to a the CAR number is referenced on and Preventive Action Procedure.  ument supplement is required or if a RB) for review and disposition.
decision.	or ous decision, the Senior Manager act reporting of delivered work that may	_
PROPRIETARY INFORMATION	This document expires 30 days after printing unle	ss marked "Released".  Date Printed:  Form Rev: Orig

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	Your Company Name	Rev: Orig
4.1.1 Major: This classification ap	oplies to	
4.1.2 Minor: This classification ap	plies to	
4.1.3 None: This classification ap	plies to	
4.2 MRB dispositions may includ	le but are not limited to:	
4.2.1 Clarification	o, but allo not immed to.	
The MRB may determine that a	Request for Support (RFS) was	prepared because of ambiguity or
•	and as such may disposition the RFS	•
must not be classified as	mprovement Opportunity at the top of	erpretation of the requirement on the
at the discretion of the MRB. This M		and retining a dispersional design to
4.2.2 Conditional Acceptance		
Nonconforming supplies or work n	nay be dispositioned 'conditional acc	ept' if they do not adversely affect
A 'conditional accept' disposition is	a qualified acceptance of work classifi	ed as
7. Conditional accept disposition to		disposition is subject to
4.2.3 Non-Deliverable Work	od Mars Deliverschiel odere design	and the second terms of the second se
conditions for individuals or their	ed 'Non-Deliverable' when their use use could adversely a	could result in nazardous or unsate
The state of the s		sposition is not subject to
4.2.4 Notification	rovement Onnewtypity may not evict fo	
	rovement Opportunity may not exist for cord of evaluation for historical reten	
party. This MRB disposition is not su		
4.2.5 Precautionary	·	
	RFS was prepared because of an ur	defined condition and as such may
disposition the RFS as 'Precautional The condition is not be classified a		s the condition and indicates on the
	hen checks Continuous Improvemen	
Further action is at the discretion of	the MRB. This MRB disposition is no	subject to
4.2.6 Repair (Non-Standard and S	tandard)	
	ible, repair action may be authorized	The MRB completes the RFS and
	prepared for their review. The MRE	
		This MADD
disposition is subject to		. This MRB
	This document expires 30 days after printing unl	ess marked "Released". Form Pour Orig
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Waiver or Request for Deviation of Subject to  4.2.8 Return to Supplier (Receiving When supplies deviate from requirgers and forwards it to Quality for Supplies received that a Supplies received that suppli	se' by the MRB but departs from specificacy be submitted to the Customer for a large Inspection).  Ing Inspection)  The ements but are considered useable the processing. This MRB disposition is substant are obviously unfit for use may  Standard)  Standard)  The according to a standard rework work appletion of a rework, the responsible per per per per per per per per per pe	en receiving personnel prepares an abject to  instruction or a Customer approved ersonnel sign the RFS and forwards ruction and if found acceptable the
<ul><li>5.1 Major: A Waiver/Deviation of</li><li>5.2 RTV and Scrap dispositions</li></ul>	are and Non-Standard Rework/Repair dis	positions are

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# CORRECTIVE AND PREVENTIVE ACTION

Origination Date: XXXX

Document Identifier:	Corrective and Preventive Action
Date:	Latest Revision Date
Project:	Customer, Unique ID, Part Number
Document Status:	Draft, Redline, Released, Obsolete
Document Link:	Location on Server (if used)

#### Abstract:

This document describes the procedures used to correct and prevent nonconformities.

Your Logo	Vous Company Nama	Corrective and Preventive Action
	Your Company Name	Rev: Orig

#### **REVISION LOG**

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	Your Company Name	Rev: Orig

#### 1.0 PURPOSE

This document provides details and procedures for the process governing the discovery, reporting, resolution and recording of actions taken to correct or prevent nonconformities.

#### 2.0 THEORY

Corrective action is taken to correct nonconformities, which could be work defects found during production, errors found in documents, equipment problems or problems related to how the Company performs functions in its processes. "Corrective action" is simply the "fix" that corrects the problem.

Whenever we take corrective action we also attempt to prevent the problem from recurring, which is known as "preventive action". There are times when preventive action is a standalone activity, specifically when reporting a problem that does not exist at the moment but could exist if something isn't done.

Having a formal system to record and resolve both existing and potential problems ensures that these problems do not occur or reoccur, thereby improving our work, processes and work environment.

#### 3.0 PROCEDURE: INTERNAL REPORTS

- 3.1 The Company utilizes a Request for Support (RFS) form to record both nonconformances related to its work, processes and quality system as well as compliments or positive feedback. The form and system are used for both potential problems (corrective action) and possible problems (preventive action.) In all cases such problems or compliments may be reported internally, reported by Customers or other external parties. A Bulletin form should be used to clarify management instructions for activities that do not strictly fall within MRB or CCB disposition.
- 3.2 ALL employees are empowered with the ability to report sources of problems and nonconformances.
- 3.3 No disciplinary action may be attached to the submission of RFS's.
- 3.4 The Project Inspector has been assigned the role of RFS Administrator.
- 3.5 For the processing and routing of RFS's see Process Map.
- 3.6 If the responsible manager determines they are not responsible for the issue involved, they must return the RFS to the RFS Administrator for re-routing.
- 3.7 Actions taken are to the degree appropriate to the problem, as deemed by management.
- 3.8 The Project Inspector monitors the RFS Log to determine overdue RFS's and takes appropriate action to see that such RFS's are resolved.
- 3.9 In addition to corrective action efforts, management utilizes

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- 3.10 The management review process ensures
- 3.11 Where work is suspected of a nonconformance, the Company takes preventive action that includes notification

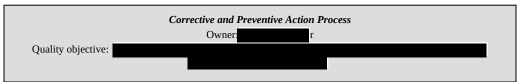
### 4.0 PROCEDURE: CORRECTIVE ACTION REQUEST (CAR)

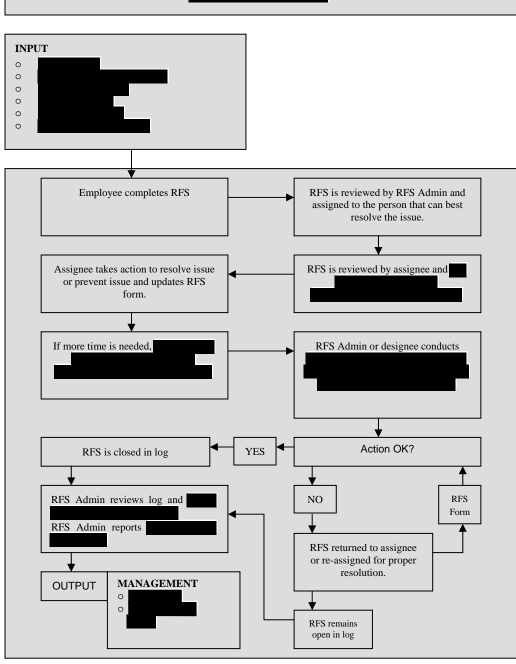
- 4.1 Any purchasing agent may submit a Corrective Action Request (CAR) to a Supplier that has shown
- 4.2 CAR's are processed through the same steps as the RFS but are routed to the Supplier for root cause analysis and action planning. CAR's are logged separately.
- 4.3 Failure of a Supplier to respond to a CAR or to respond with an insufficient action plan may mean

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Corrective and Preventive Action
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### 5.0 PROCESS MAP





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# DOCUMENT CONTROL

Origination Date: XXXX

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Date:	Latest Revision Date
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Document Link:	Location on Server (if used)

### Abstract:

This document describes procedures for controlling documents.

Your Logo	Vous Company Nama	Document Control
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### 1.0 PURPOSE

This procedure defines the requirements for the control of documents within the quality program. The scope of this procedure is to control documents specifically defined in section 3.0. The Document Control Center ensures that documents are controlled so that information on them is accessible, legible and suitably maintained and obsolete documents are stamped "Superseded".

The following documents are not subject to this procedure:



### 2.0 THEORY

Documents must be controlled so that only reviewed and approved information is released and used by employees. This ensures that no mistakes are made due to the usage of obsolete information.

### 3.0 DOCUMENT TYPES

- 3.1. Quality Program: this document provides the primary Governing Policies. It also defines top-level requirements for the quality program and defines how the Company meets the requirements of Colorado Resolution 35 for non-residential structures.
- 3.2. QMS Procedures: these documents provide additional detail for certain procedures where such detail is required. The Quality Program includes references in bold-italic font to the applicable QMS procedures.
- 3.3. General Work Instructions: these documents provide machine-level or task-level details on what is required to perform specific work. These are typically specific to a department or work step. These do not include job-specific work instructions that are made part of the engineering documents and controlled via other procedures (see 1.0 above.)
- 3.4. Inspection Instructions: these documents are developed by or under the supervision of the Project Inspector using requirements from the applicable engineering drawings and/or technical documentation.
- 3.5. Forms: these documents are produced by a streamlined creation and control process. Any department manager or area supervisor may develop a new form for use in their area.
- 3.6. Records that are created for temporary retention of miscellaneous information are not required to be maintained or controlled, such as personal notes written on a scratch pad, post-it note or form identified with a watermark or the term "Note Pad".

### 4.0 QUALITY PROGRAM

4.1. Creating the Quality Program

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The Quality Program has been devel	loped by top management of the Com	pany.
4.2. Review and Approval The Quality Program is reviewed an	nd approved by top management	
,	ectronically through the Company's in retain older hardcopies or softcopies	
In some cases, a hardcopy of the O If the document is needed for more t	Quality Program may be given to an han thirty (30) days it is	employee, department or Customer.
Request for Change or an Engineeri	e to the Quality Program. Requests fing Order (EO) form and submitting it management. All changes to the Qu	to top management or to the Project
<ul><li>5.0 QUALITY PROGRA</li><li>5.1. Creating New QMS Procedure</li><li>QMS procedures should be created</li></ul>		
	approved by top management. At lead document should be responsible for	
	ctronically through the Company's in retain older hardcopies or softcopies	
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	Your Company Name	Rev: Orig

#### 5.4. Change Control

Changes to QMS procedures are performed in the same manner as the Quality Program.

#### 6.0 GENERAL WORK INSTRUCTIONS

#### 6.1. Creating New Work Instructions

Where necessary, work affecting quality is described by clear and complete documented work instructions that define what is required to perform specific work functions. Typically, new work instructions are developed by or under the supervision of an area manager or subject matter expert. Work instructions should be created as

#### NOTE REGARDING JOB SPECIFIC WORK INSTRUCTIONS:

Engineering may develop work instructions that are specific to a given job, which are released and controlled as part of the technical documentation. Their format may be different from general work instructions.

#### 6.2. Review and Approval

Work instructions must be reviewed and approved by the CCB. At least one member of the CCB responsible for reviewing the document should be responsible for the area affected by the document. Approval is indicated by

#### 6.3. Distribution

General work instructions are distributed electronically through the Company's internet server and/or via the intranet. The Document Control Center may retain older hardcopies or softcopies for

#### 6.4. Change Control

Changes to general work instructions are performed in the same manner as the Quality Program. When general work instructions are changed,

### 7.0 INSPECTION INSTRUCTIONS

#### 7.1. Creating New Inspection Instructions

New inspection instructions are developed by or under the supervision of the Project Inspector using requirements from the applicable engineering drawings and/or technical documentation. Inspection instructions should be created as

#### NOTE REGARDING JOB SPECIFIC INSPECTION INSTRUCTIONS:

Engineering may develop inspection instructions that are specific to a given job, which are released and controlled as part of the technical documentation and subject to the controls of the Configuration Management Procedure - not this procedure. Their format may be different from general work instructions.

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	Your Company Name	Rev: Orig

#### 7.2. Review and Approval

Approval is indicated by

#### 7.3. Distribution

Inspection instructions are distributed electronically through the Company's internet server and/or intranet. The Document Control Center may retain older hardcopies or softcopies for

#### 7.4. Change Control

Any employee may request a change to inspection instructions by completing a Request for Change form and submitting it to the Project Inspector. All changes to inspection instructions go through the same review and approval as the original release. When changes are approved the revision indicator is

### 8.0 FORMS

#### 8.1. Creating New Forms

Forms undergo a streamlined creation and control process. Any department manager or area supervisor may develop a new form for use in their area. The form should be created in software format (MS word, Excel, etc) and then submitted to the appropriate department manager for review and approval. Forms are a special kind of document that may be photocopied as needed; furthermore, forms do not require an approval signature. Department managers are responsible for creating and using forms in their areas.

#### 8.2. Review and Approval

Forms may be reviewed and approved by the manager of the department or area primarily affected by the form. Forms do not require a signature approval; instead, the manager approving the form shall

#### 8.3. Distribution

Forms are made available through the Company's internet server, intranet or Document Control Center. These may be printed and photocopied as needed. When hardcopies run out, a new copy is to be printed for photocopying. Photocopying from a previously photocopied form is

### 8.4. Change Control

Any employee may submit a request for change to the appropriate area manager responsible for the form and the manager will determine if the form should be revised. Revised forms go through the same review and

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Your Logo	Vous Company Nama	Document Control
	Your Company Name	Rev: Orig

approval as originals but must

### 9.0 EXTERNAL DOCUMENTS

9.1. Some external (third party) standards or specifications may be maintained on file without control provided that the revision indicator is evident somewhere in the document. This is necessary because Unless otherwise specified, if the revision level is not shown on documents, then

9.2. Third party specifications and engineering drawings, including those of the Customer are controlled. Where control of an external document is deemed necessary, they shall be made available by the Document Control Center, which shall

### 10.0 PERIODIC RE-EVALUATION OF DOCUMENTS

The entire set of quality documentation is subject to continuous improvement. Change control documents are

Your Logo Your Company Name

# PRODUCTION PROCEDURE

Origination Date: XXXX

Document Identifier:	Production Procedure
Date:	Latest Revision Date
Project:	Client, Unique ID, Part Number
Document Status:	Draft, Redline, Released, Obsolete
Document Link:	Location on Server (if used)

### Abstract:

This document describes the production process.

Vour Logo	Vous Company Nama	Production Procedure
Your Logo	Your Company Name	Rev: Orig

### **REVISION LOG**

Issue	Date	Comment	Author
Orig			

### DOCUMENT CHANGE RECORD

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Form Rev: Orig

Vour Logo	Vous Company Nama	Production Procedure
Your Logo	Your Company Name	Rev: Orig

### 1.0 PURPOSE

This document defines the overall Production process and includes or makes reference to the procedures necessary for the process. The Production process includes all required inspections and tests.

### 2.0 THEORY

Production operations or tasks are conducted under controlled conditions to ensure Production quality. By this we mean:

- Ensuring Operators have a good work environment and training
- Ensuring Operators have good equipment and tools
- Properly handling and preserving raw materials
- Supplying adequate work instructions, drawings, etc., where needed

### 3.0 PROBLEM RESOLUTION

All employees are instructed to immediately notify a Responsible Authority (RA) whenever a process or Production related problem occurs that cannot be corrected according to established process controls and could affect or actually affects the quality of the Production.

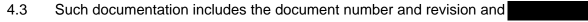
It is understood that the appropriate responsible authority will occasionally not be available for support; in that event, contact each of the following personnel in the order listed until an appropriate authority can make a decision to resolve the problem. No disciplinary action may be attached to an employee's attempt to resolve a problem.

For instance (replace with your responsible authority):



### 4.0 DOCUMENTATION

- 4.1 All revision controlled documents are available at the point of use.
- 4.2 In addition to this procedure, additional documentation may be required according to



4.4 Records that are created for temporary retention of miscellaneous information are not required to be maintained or controlled, such as

PROPRIETARY INFORMATION	This document expires 30 days after printing unless marked "Released".  Date Printed:	Form Rev: Orig

	Your Logo	Your Company Name	Production Procedure
	Tour Logo	Tour Company Ivame	Rev: Orig
5.0	IDENTIFICATION		
5.1	Production materials/work-zo	nes are identified by any of the followi	ing methods:
•			
5.2 staff re		s maintained on the appropriate pap determine the requirements for seria	
	Nonconforming Production thements is <b>ontrol of Nonconformances</b>	at has failed an inspection or test and <b>Procedure</b> .	d cannot be reworked to comply with
5.4	Any materials or Production	n not marked with a tag are cons	idered
5.5	Identification of Transfer Che	mical Containers	
5.5.1 contair	Whenever a portion of chemer, the	mical is transferred from its origina	l container to a smaller temporary
5.5.2 contail	Whenever a portion of cherner, the	mical is transferred from its original	container to a smaller permanent
6.0	<b>MATERIAL HANDL</b>	ING	
6.1	Work instructions and/or train	ing instructs Operators on the proper	and safe handling of materials.
6.2	In all cases, Operators handle	e materials	
		able safety and personal protection e lired to wear or use such equipmen	

### 7.0 PRESERVATION

<ul> <li>7.2 Operators employ</li> <li>7.3 Operators employ</li> <li>7.4 Operators employ</li> </ul>	7.1	Operators (	employ			
	7.2	Operators	employ			
7.4 Operators employ	7.3	Operators				
	7.4	Operators	employ			

	Vouglass	Vous Company Name	Production Procedure
	Your Logo	Your Company Name	Rev: Orig
7.5	Work instructions and training	ng methods ensure	
7.0	Work mondonone and training	ig mornous enound	
7.6	Work instructions and training		
7.7	Work instructions and training	g methods ensure	ls.
<b>8.0</b>	<b>CLIENT AND GOVE</b>	ERNMENT PROPERTY C	ONTROL
8.1 the C includ	lient and Government or acqu	erty (C&G Property) means all hardwired by the Client and Government u	
•			
•			
•			
8.2 <b>Proce</b>		urnished property is inspected upon shortages are communicated to the Cl	
8.3	C&G Property is identified	with	
8.4	Sensitive material as defined	by the Client or Government is	
8.5	C&G Property is only used a	as instructed or required by the Client	t or Government contract and is not
8.6	C&G provided equipment is	subject to	
8.7	The Company investigates a	and reports to the Client or Governn	nent any cases of
8.8	Requirements for the control	of C&G property is	
9.0	SHELF LIFE EXTER	NSION	
9.1 expira	Items that are subject to e	xpiration may be repeatedly extend	
Produ	ction conditions; for instance:	or b	y verification of performance under
	·		
0.2	Chamicals that are purchase	d or propored by the laboratory are	
9.2	Onemicais mai are purchased	d or prepared by the laboratory are	
		The state of the s	1 100 1 10 1
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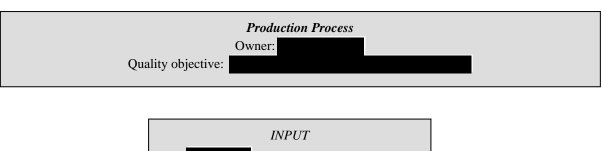
Your Logo Your Company Name Production Procedure

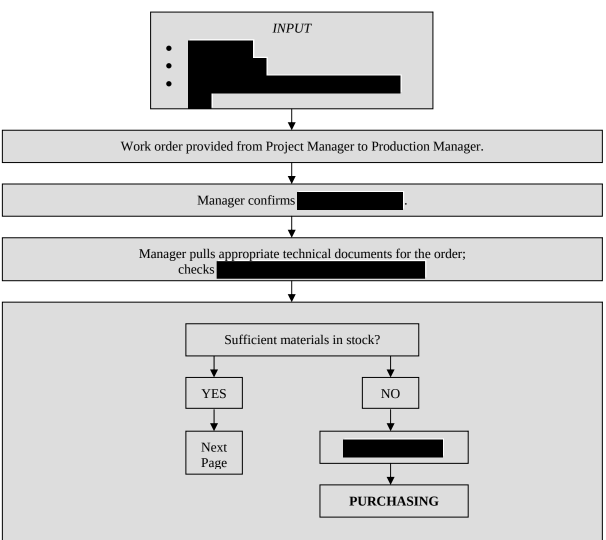
Rev: Orig

9.3 Raw material components whose shelf life has been extended displays the

### **10.0 PROCESS MAP**

continued next page...



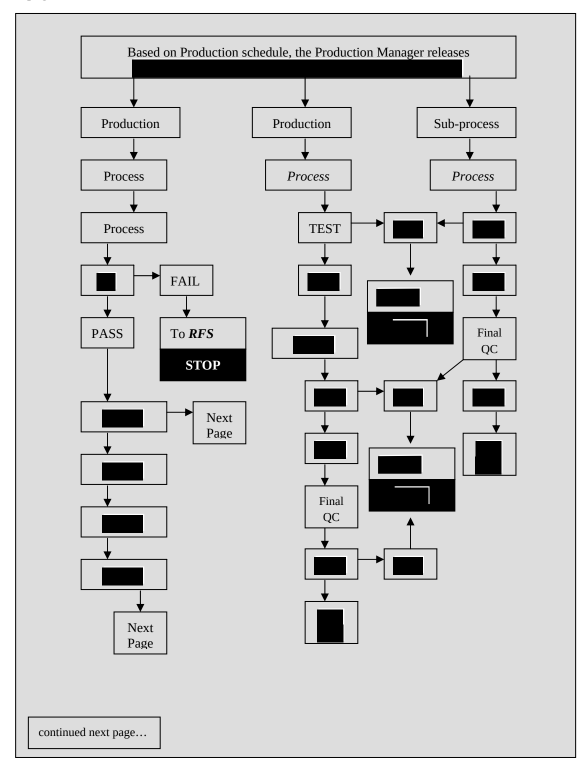


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Your Logo Your Company Name Production Procedure

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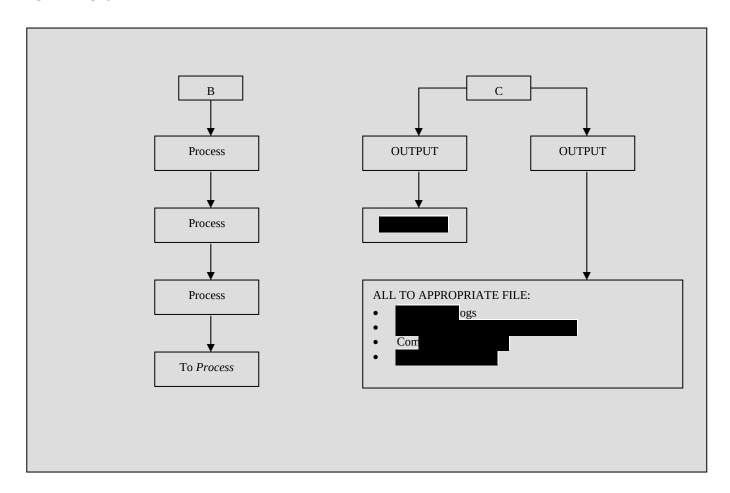
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Your Logo Your Company Name Production Procedure

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Your Logo Your Company Name

# **PURCHASING**

Origination Date: XXXX

Document Identifier:	Purchasing Procedure	
Date:	Latest Revision Date	
Project:	Customer, Unique ID, Part Number	
Document Status:	Draft, Redline, Released, Obsolete	
Document Link:	Location on Server (if used)	

### Abstract:

This document describes the purchasing process.

Your Logo	Vous Company Nama	Purchasing Procedure
	Your Company Name	Rev: Orig

### **REVISION LOG**

Issue	Date	Comment	Author
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### DOCUMENT CHANGE RECORD

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	Your Company Name	Rev: Orig

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5.0	OTHER PURCHASING RULES	6
6.0	PROCESS MAP	8

Your Logo	Purchasing Procedure	
	Your Company Name	Rev: Orig

### 1.0 PURPOSE

This document defines the Purchasing process including or making reference to procedures for the various activities within the process.

Note: this procedure applies to suppliers of products or providers of services that directly affects the quality of our products or services. Suppliers that provide office and maintenance supplies, furniture, grounds keeping services, etc. are not subject to the controls of this procedure.

### 2.0 THEORY

The purchase of materials that go into our products or services that help us produce products affects everything we make. As a result, it is important to monitor and control the quality of both products and services that we receive as well as the suppliers of such products and services.

### 3.0 PROCEDURE: SUPPLIER EVALUATION AND SELECTION

- 3.1 All suppliers of product related materials or services must be evaluated unless these suppliers are:
- 3.2 Supplier evaluation is conducted by following the format on the Supplier Evaluation Form.
- 3.3 The Supplier Evaluation Form ensures
- 3.4 Once approved through the Supplier Evaluation Form, the Project Inspector will update the Approved Supplier List.
- 3.5 The following ratings apply to suppliers:
- RESTRICTED:
- CONDITIONAL:
- UNRESTRICTED:
- DOCK-TO-STOCK:
- 3.6 Once entered into the Approved Supplier List, suppliers are rated as
- 3.7 Using incoming (receiving) inspection results for product suppliers and employee feedback on service providers, the Project Inspector will

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	Your Logo	V C N	Purchasing Procedure	
		Your Company Name	Rev:	Orig
3.8 will	Using the results from combine	nation of the following functions for pro	oduct suppliers, the Project Inspe	ctor
VVIII				
3.9 Perfor		duct, incoming inspection results a which calculates the Supplier's cur		
receiv	ed and parts accepted. A ne		may be upgrade	
UNRE	ESTRICTED.			
3.10	If a new Supplier rates			
	и пом обррно пакоо			
3.11	If any Supplier rates			
	, , , ,			
3.12	If itoms are returned to any	Supplier using a Material Shipper, th	o Project Inspector will	
5.12	in items are returned to any	Supplier using a Material Shipper, th	le i roject inspector wiii	
3.13	Any Supplier may be de-rate	d to		
3.14	Management may override			
3.15	During management review,	the entire Approved Supplier List is		
4.0	PROCESSING REC	UISITIONS AND PURCH	ASE ORDERS	
4.1	During review of each requis	ition, the Quality Group will		
4.2	When appropriate, the purch	ase order defines		
4.3	As applicable, purchase orde	r information includes:		
a)				
b)				
c)				
d) rea	uirements relative to:			
-	anomente relative to.			
	DD ODDIETA DV INFORMATION	This document expires 30 days after printing unle	ess marked "Released".	
	PROPRIETARY INFORMATION		Date Printed: Form Rev: Orig	

	Your Logo	Your Company Name	Purchasing Procedure
		Tour Company Ivame	Rev: Orig
- e)			
f)			
g)			
4.4	The requirements for delegat	ion are defined when the Company	
		. ,	
4.5	When the Company or its C	ustomer needs to perform verification	n activities at a Supplier facility, the
1.0	When the company of its of	actomer medac to perform vermeation	Tractivities at a supplier lacinty, the
16	Soo the process man herein		
4.6	See the process map herein.		
4.7			ize the shift foreman and/or the
maint	enance foreman emergency p	urchase authority for	
5.0	OTHER PURCHASI	NG RUI FS	
5.1	In all instances, the Purchasir		
	,		
5.2	Any employee of the Purch:	asing Denartment that has any fina	ncial or other interest in a supplier
_		any member of his/her immediate fa	
5.3	The acceptance by purchasin	g personnel of gifts or gratuities from	euppliore is
5.5	The acceptance by purchasin	g personner or girts or gratuities from	suppliers is
5.4		ended for the purpose of advertise	ment and bearing the name of the
Suppl	ier is		
5.5			I activities and will participate where
	sted in all necessary meetings in all necess	with Customers. Customers wishing	to visit or contact suppliers regarding
mater	iale on order may		
5.6	The Purchasing department	will not, in any way,	
	PROPRIETARY INFORMATION	This document expires 30 days after printing unle	ess marked "Released".  Date Printed:  Form Rev: Orig

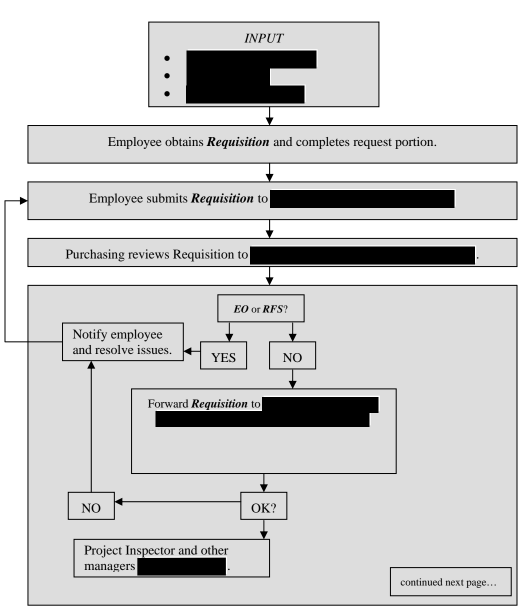
Your Logo	Vous Company Nama	Purchasing Procedure
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5.7 The Company will abide by all Government clauses or other statutory or regulatory requirements as referenced by the order, contract or other requirements document.

Your Logo
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Purchasing Procedure
Rev: Orig

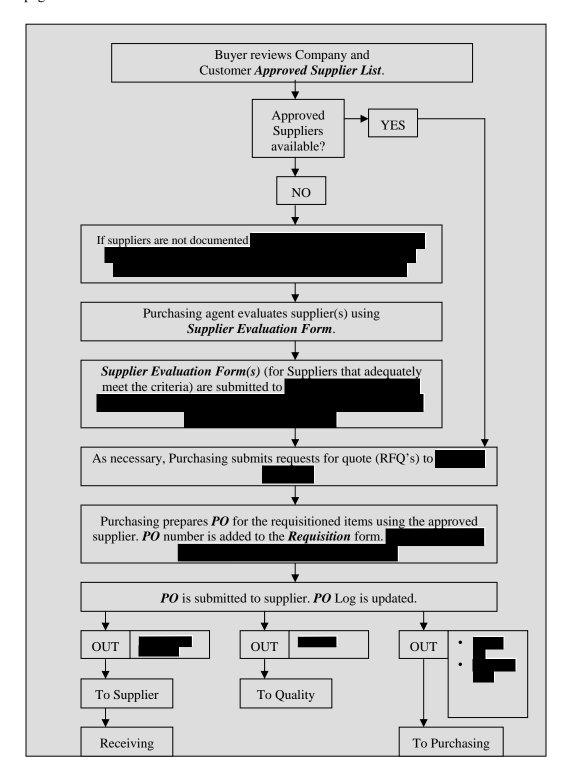
### 6.0 PROCESS MAP





Your Logo	Your Company Name	Purchasing Procedure
		Rev: Orig

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Your Logo Your Company Name

## RECEIVING INSPECTION

Origination Date: XXXX

Document Identifier:	Receiving Inspection	
Date:	Latest Revision Date	
Project:	Customer, Unique ID, Part Number	
Document Status:	Draft, Redline, Released, Obsolete	
Document Link:	Location on Server (if used)	

### Abstract:

This document describes the receiving and inspection process.

Your Logo	Your Company Name	Receiving Inspection
		Rev: Orig

### **REVISION LOG**

Issue	Date	Comment	Author
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### DOCUMENT CHANGE RECORD

Issue	Item	Reason for Change

Your Logo	Vous Company Nama	Receiving Inspection		
Your Company Name	Rev: Orig			

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PROPRIETARY INFORMATION

Your Logo	Vous Company Nama	Receiving Inspection
Tour Company Name	Your Company Name	Rev: Orig

### 1.0 PURPOSE

This document defines the Receiving process including receiving inspection activities and includes or makes reference to procedures for the various activities within the process.

### 2.0 THEORY

Receiving is the first line of defense to prevent sub-standard supplies from affecting the Company's process or product quality; however, sampling and 100% incoming inspection is only a part of the collection of applications that are necessary to assure the release of conforming supplies to stock. Receiving inspection cannot provide 100% assurance of product or process quality because the affect that the supplies have on production-level activities cannot be assessed or verified - only by 100% use of supplies will defects be detected in product or process quality.

As a result of teaming and intelligent design, the Company ensures that deliverable supplies meet Customer requirements prior to packaging and shipping.

### 3.0 PROCEDURE: RECEIVING

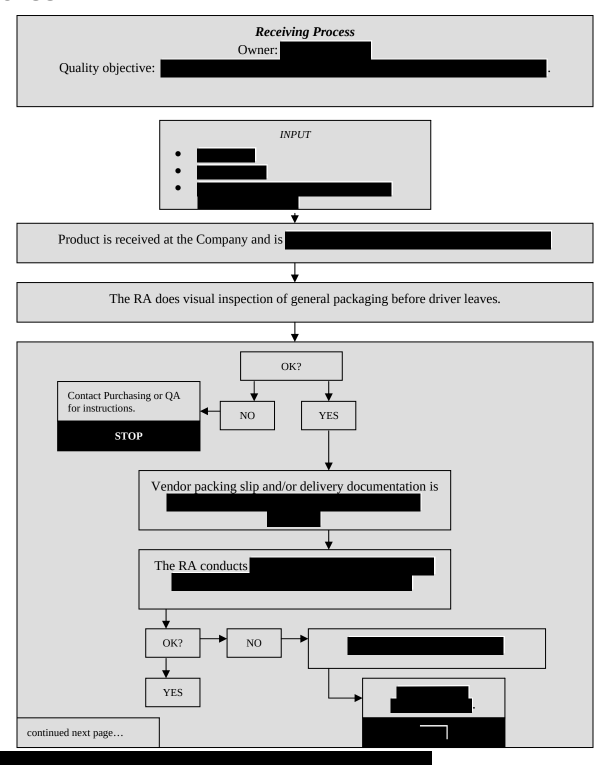


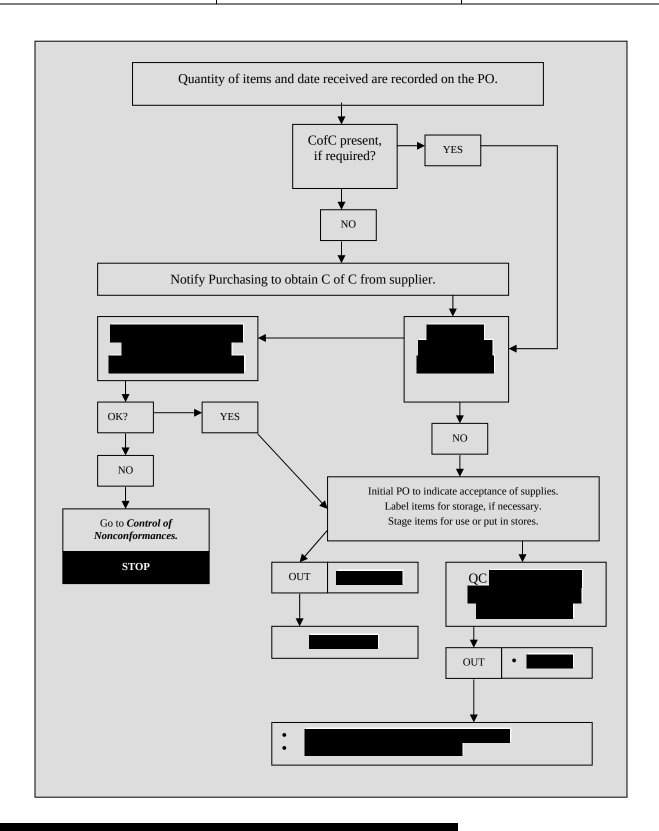
### 4.0 PROCEDURE: RECEIVING INSPECTION

- 4.1 The inspector will receive the items and original paperwork from the RA and acquire the applicable PO. (see the Purchasing Procedure)
- 4.2 Inspections are performed according to Appendix A or as required by work instruction, Customer requirements or other documentation. The results are recorded on the applicable forms and the purchase order is processed according to Appendix B.

Your Logo
Your Company Name
Receiving Inspection
Rev: Orig

### **PROCESS MAP**





Your Logo	Vous Company Nama	Receiving Inspection
Tour Compa	Your Company Name	Rev: Orig

## **APPENDIX A - RECEIVING INSPECTION WORK INSTRUCTIONS**

Op 1:	Acquire copy of purchase orde	r. Perform <r< th=""><th>ough Order&gt;</th><th>verifica</th><th>tion</th><th></th><th></th><th></th></r<>	ough Order>	verifica	tion			
Op 2:	Verify supply							
Op 3:								
On 4	Verify the Supplier is							
οр 4.	voiny the Supplier is							
Op 5:	If the supply is a <catalog co<="" td=""><td>ommercial&gt; it</td><td>em.</td><td></td><td></td><td></td><td></td><td></td></catalog>	ommercial> it	em.					
Op 6:	Perform First Piece Mechan	ical/Visual in	spection on	a new	production	part numb	per to	
Op 7:	SAMPLING PLAN:							
			41					
Op 8:			then					
	then							
Op 9:		. t	hen					
Op 10	: Verify conformance to	,						
	<ul> <li>: When raw material is accepved Supplier List for item critica</li> </ul>					ınalysis, re	eview the	current
	itical item:							
For no	n-critical item:							
	PROPRIETARY INFORMATION	This documen	nt expires 30 days a	fter printing		eleased". Printed:	Form Rev: Orig	;

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Your Company Name
Receiving Inspection
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Op 12: Verify lot traceability is
Op 13: If the Supplier is a distributor
Op 14: Affix a Good Material Tag to accepted supplies. For supplies that exhibit a lot number for traceability use the
Op 15: If supplies are nonconforming or their conformance cannot be determined within 30 days of receip prepare
Op 16: Complete inspection record and record the Op 17: Complete shelf life expiration log for supplies that have an expiration date Op 18: Record the the Purchase Order according to Appendix B
Op 19: If the Supplier's
Op 20: Inspect Customer/Government furnished property upon receipt to

Your Logo	Vour Company Nama	Receiving Inspection
	Your Company Name	Rev: Orig

## **APPENDIX B - PURCHASE ORDER PROCESSING**

Step	IF	THEN
1	Supply is not the Last Item on PO	Produce a copy of the PO - attach packing slip to the copy of PO and
2	Supply is the last Item on PO	Attach the Supplier's packing slip to the original PO - produce a copy of the PO - forward
2.1	Supply is the last Item on PO	Optional:

Your Logo Your Company Name

# **RECORDS CONTROL**

Origination Date: XXXX

Document Identifier:	Records Control	
Date:	Latest Revision Date	
Project:	Customer, Unique ID, Part Number	
Document Status:	Draft, Redline, Released, Obsolete	
Document Link:	Location on Server (if used)	

#### Abstract:

This document describes the procedure for control of records.

Your Logo	Voya Compony Nome	Records Control
	Your Company Name	Rev: Orig

## **REVISION LOG**

Issue	Date	Comment	Author
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## DOCUMENT CHANGE RECORD

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Your Logo	Vous Company Nama	Records Control
	Your Company Name	Rev: Orig

#### 1.0 PURPOSE

This procedure defines the requirements for the control of records within the quality program. The scope of this procedure is to control only the records referenced in this document; other records are not controlled.

#### 2.0 THEORY

A record is any written or electronic piece of evidence that may be needed later to provide evidence of conformity to requirements. Typically a blank "form" becomes a "record" when it is completed. Records must be controlled so that the information on them is accessible, legible and suitably maintained.

#### 3.0 RULES FOR CONTROL OF RECORDS

3.1 The controls for each type of record are defined in *Appendix A* of this procedure. 3.2 The listed "controller" must 3.3 Records for active contracts are maintained in the quality department handling the operations. The Document Control Center maintains archive files for records. Records shall be maintained a minimum of 3.4 Records that are discarded after retention shall 3.5 Hardcopy records are to be 3.6 3.7 Records are available for review by the Customer and copies of non-proprietary records are furnished to the Customer upon request. Non-disclosure agreements are required for non-Governmental entities. 3.8 Records are verified for The Company does not require vendors to maintain records for the Company; instead, 3.9 To ensure protection of records, electronic records are subject to 3.10 Local computer data that is stored on company computers must 3.11 3.12 When making corrections to written record entries, the error is Correction fluid or correction tape is 3.13

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Your Logo	Vous Commons Nome	Records Control
	Your Company Name	Rev: Orig

## **Appendix A: Records Matrix**

Required Record or Document Type	Company Record	Controller	Туре	Location	Minimum Retention
Calibration records	Calibration		Form		
Contract review records	Contract review		Form		
Control of Nonconformances	RFS		Form		

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Your Logo Your Company Name

# SHIPPING PROCESS

Origination Date: XXXX

Document Identifier:	Shipping	
Date:	Latest Revision Date	
Project:	Customer, Unique ID, Part Number	
Document Status:	Draft, Redline, Released, Obsolete	
Document Link:	Location on Server (if used)	

#### Abstract:

This document describes the shipping process.

Your Logo	Vous Company Nama	Shipping Procedure
	Your Company Name	Rev: Orig

## **REVISION LOG**

Issue	Date	Comment	Author
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## DOCUMENT CHANGE RECORD

Issue	Item	Reason for Change

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Your Logo	Vous Company Nama	Shipping Procedure
	Your Company Name	Rev: Orig

## 1.0 PURPOSE

This document defines the Shipping process including product packaging activities.

#### 2.0 THEORY

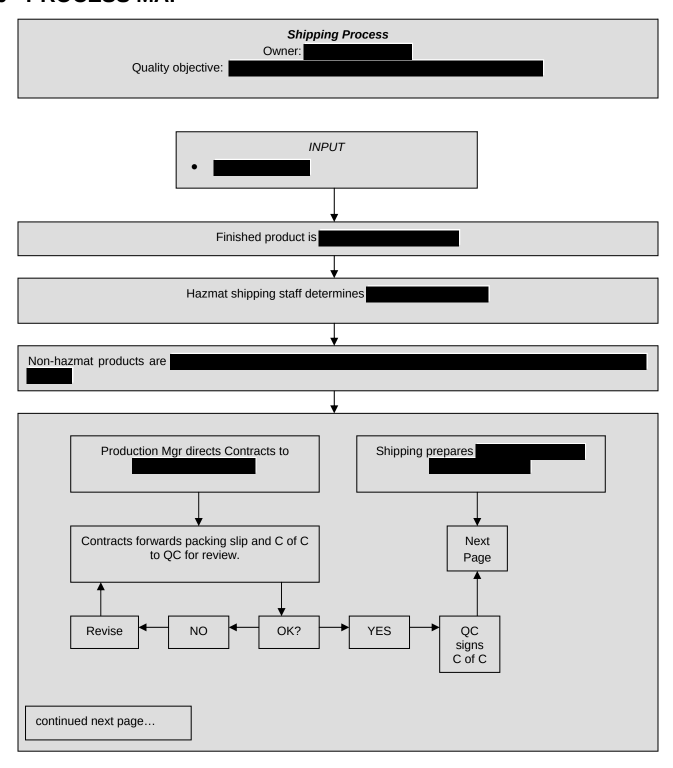
The final packaging and arrangement of shipping is

#### 3.0 PROCEDURE: PACKAGING AND SHIPPING

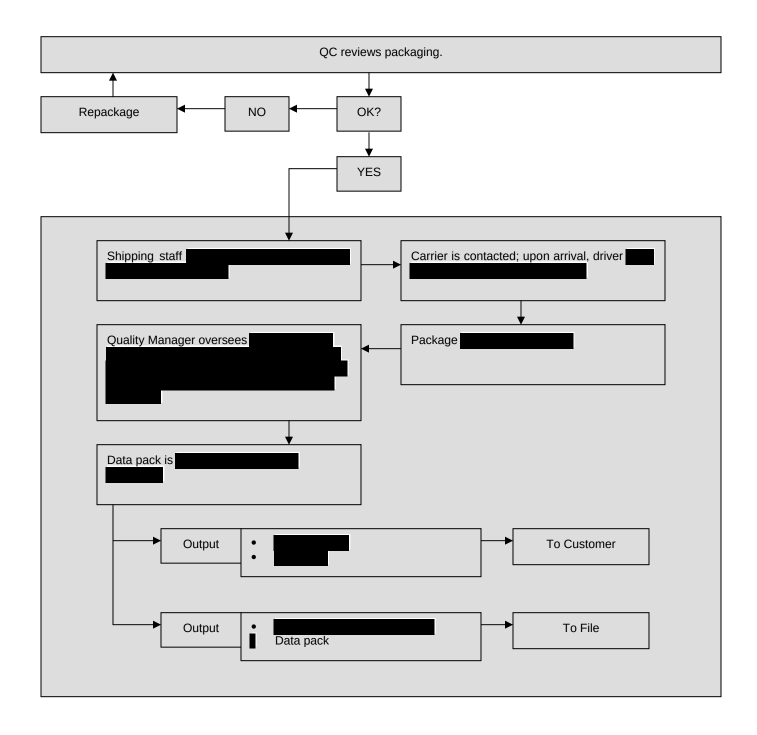
See process map.

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Your Company Name
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#### 4.0 PROCESS MAP



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# TRAINING PROGRAM

Origination Date: XXXX

Document Identifier:	Training	
Date:	Latest Revision Date	
Project:	Customer, Unique ID, Part Number	
Document Status:	Draft, Redline, Released, Obsolete	
Document Link:	Location on Server (if used)	

#### Abstract:

This document describes training program and requirements.

Your Logo	Vous Company Nama	Training Program
	Your Company Name	Rev: Orig

## **REVISION LOG**

Issue	Date	Comment	Author
0-0			

## DOCUMENT CHANGE RECORD

Item	Reason for Change
	Item

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2.0	THEORY	. 4
3.0	TRAINING PROCEDURE	. 4

Your Logo	Your Company Name	Training Program	
	Tour Company Name	Rev: Orig	

#### 1.0 PURPOSE

This document provides details on the Company's training program and requirements.

#### 2.0 THEORY

Employees can only perform their duties adequately when properly trained. The Company intends to ensure adequate employee performance through a robust training program that includes initial orientation, assessment of abilities and on-the-job training to enhance those abilities.

#### 3.0 TRAINING PROCEDURE

3.1 Hiring

Employees are hired on their basis to

To accomplish this, potential candidates are

3.2 Initial Indoctrination and Orientation

Once hired, new employees are

#### 3.3 On the Job Training

Once an employee has completed initial indoctrination they undergo on-the-job training relative to their position. This training is

#### 3.4 Additional Training

At the discretion of management, additional training may be conducted at any time.

This may be necessitated by

PROGRAM NAME:	DOCUMENTS AFFECTED:	
PROCESS AFFECTED:	PROJECT ENGINEER AFFECTED:	
PROCESS OPERATOR	SUPERVISOR	
AFFECTED:	AFFECTED:	
QUALITY OPERATOR AFFECTED:	PREPARED BY:	Date:
ATTECTED.		
This Bulletin is a protected doo	cument 'tab' to each field to input info	ormation.
To preserve the format of this in <a href="Forms">Forms</a> .	Bulletin please click on <tools> <prote< td=""><td>ect Document&gt;</td></prote<></tools>	ect Document>
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Delete the text is this area		
		r (list your exception here)
This Bulletin does <b>NOT</b> affect the	e Configuration Status of any Program	
PAGE 2 TEXT BLOCK: Insert page 2	text here	

BULLETIN

Your Logo

NUMBER:

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Your Logo

# INVESTIGATION AND CORRECTIVE ACTION REQUEST

ICA.	<b>R</b> Responsi	ble Supplie	er:			
Customer:	Part#	Applicabl	e Customer P.C	or Job #		
						<u></u>
						_
					_	
			_			
			Verified how?			
Similar par	rts affected: _					
6.						
9. Congratula	te the Team!					
					1	

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## REQUEST FOR SUPPORT

<b>■</b> Nonconforma	ance [	<b>Cont</b>	inuous Improve	ment O	pportunity	$\square$ Calculated Ris	k Release
SUBCONTRACTOR:					DATE RE	CEIVED:	
RFS#:						SHEET	OF
Punch #:	I	Bldg#:		Quantify:		Job Number:	
Item Name:	•		Description: ID S/B Spec		Condition w/Quant	ity &Dimension Affected	# Discrepant
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Form Rev: Orig

## REQUEST FOR CHANGE

Desired Change:		
Preparer Name:	Submittal Date:	

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#### Abstract:

This document describes xxxxxx.

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Issue	Date	Comment	Author
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## DOCUMENT CHANGE RECORD

Issue	Item	Reason for Change

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- 1.0 SCOPE
- 2.0 THEORY
- 3.0 REFERENCES
- 4.0 EQUIPMENT
- 5.0 MATERIALS
- 6.0 OPERATING PROCEDURES
- 7.0 WORKMANSHIP

Appli	cation		Revi	sions	
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## WORK INSTRUCTION NAME

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STEP	RESPONSIBILITY Operator	ACTION							
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	IF	THEN							
	Operator								
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	IF	THEN							
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- 1.0 PURPOSE
- 2.0 THEORY
- 3.0 PROCEDURE
- 4.0 PROCESS MAP

#### 5.0 REVISION HISTORY AND APPROVAL RECORD

Rev	Nature of changes	Eff. Date	Approved by
Orig			

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