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Abstract:

This document describes the Company's quality management system.

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Paragraph 5.7.3 is "value added" content.

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1.0 Purpose

The purpose of the Quality Management System is to

fabricate steel products.

2.0 Scope

The Company's quality management system

The Company's AISC Certification should

This Quality Management System includes

2.1 Exclusions

The Company cites no exclusions to the AISC standard. (revise as required)

3.0 References

The latest editions of the following documents and standards are required:



4.0 Definitions

See **QMS-16 Definitions, Abbreviations and Symbols Procedure** for more details. Subordinate or external documentation referenced herein is displayed in **Bold Italics**.

5.0 Management Responsibility



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5.1 Policy for Quality and Quality Goals

The Co	mpany's quality policy which includes	according to	
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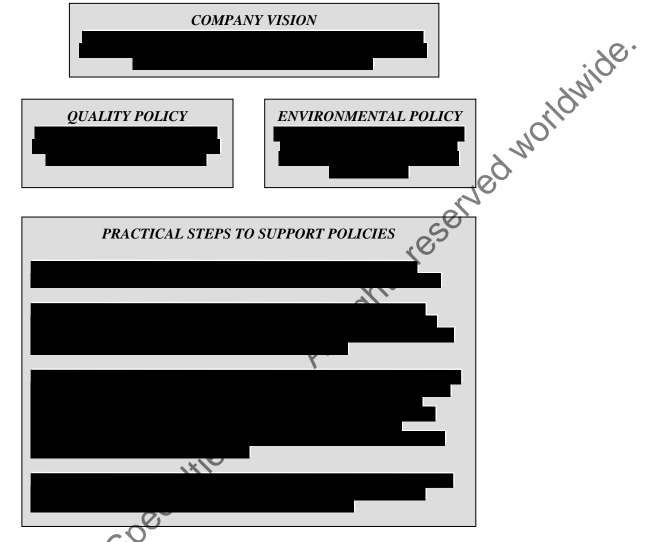
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QUALITY POLICY

ENVIRONMENTAL POLICY



Quality Management System renumbered from 5.5

The Company's quality management system is designed to satisfy AISC standard 207-23,

and includes

The Quality System ensures

This assures

Necessary records of activities are

The System is structured from top-down using this Quality Manual,

Supporting Documents, Work Instructions and Quality Records.

The Company maintains

All Managers are responsible for

The quality system

documentation is comprised of a hierarchy of documents that flow from this Quality Manual.

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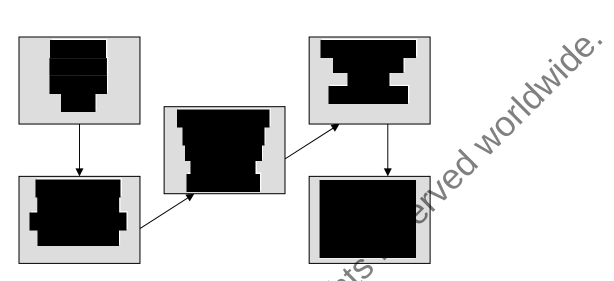
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5.3 Management Review

Review meetings are

Reviews are reported and records

are retained and maintained. The controls for management review are defined in the **QMS-04**Management Procedure, which defines

Management review meeting reports are

Internal quality audits are conducted according

to the QMS-12 Internal Auditing Procedure to

Records of the management review meetings and

internal audits are controlled according to the QMS-01 Control of Documented Information Procedure.

5.4 Responsible Quality Personnel

The individual designated as Quality Manager (QM)

The Quality Manager

however,

Although the Quality Manager

The Quality Manager has

The Quality Manager ensures

5.5 Resource Management

The Company has

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5.5.1 Personnel

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Personnel assigned to key positions and those performing field operations

according to the QMS-06

Training Procedure. Unless otherwise noted, personnel can

Specifically, individuals responsible for Quality Assurance and Quality Control management do idhits reserve not

The Company retains

5.5.2 Buildings, Workspace, Equipment and Associated Utilities

The facility consists of areas and buildings that

The areas and buildings are

The fabrication facility includes

Ambient conditions are

Equipment includes

The facility also provides

5.5.3 Fabrication Process Equipment (Hardware & Software)

The Company has under its control

Equipment is maintained to

Internal Communication

To ensure proper communication between and throughout all levels of employees within the Company,

which is documented in the QMS-04 Management Procedure.

Management periodically

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Employees are encouraged to

This system requires management to mide.

Quality Manual

The primary purpose of the Quality Manual and QMS Procedures is

The Quality Manual has been developed by

The quality manual is It is meant to be The quality manual is

Externally distributed copies are

Additional procedures and work

instructions have

For instance:

5.7.1 Organization

Review meetings are held by all managers

Reviews are

reported and records are retained and maintained. The controls for management review are defined in the QMS-04 Management Procedure.

The organizational chart

which are further defined in the QMS-05 Responsibilities and Authorities

Procedure.

The qualifications of key personnel and managers listed in paragraph 5.4.1 are maintained in records and/or job descriptions according to the training program that is defined in the QMS-06 Training Procedure.

- See applicable project facility plan/map for detailed description of facility.
- See applicable equipment list designated for projects.

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5.7.2 Approval

This manual is issued under the authority of referenced herein are

Management ensures the QMS is
Subsequent major changes that may affect the performance, quality or reliability of deliverable items are identified, reviewed and

5.7.3 Order of Precedence Value-Added

The order of precedence of order-specific documentation is as follows unless otherwise directed by Customer or government requirements:

1. er	
2.	
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5.	

6.0 Construction Document Review and Communication

The Company performs according to the **QMS-07 Proposal Development and Contract Review Procedure**. The review

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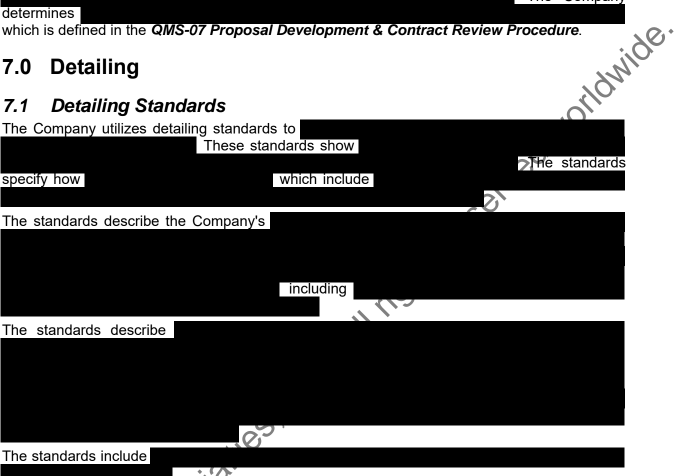
determines

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which is defined in the QMS-07 Proposal Development & Contract Review Procedure.

Detailing 7.0

Detailing Standards 7.1



7.1.1 Digital Document Production - Preparation of Fabrication and Erection Documents (also see 7.8)

The Company has prepared and implemented the QMS-17 Detailing Procedure for The procedure identifies The procedure describes The procedure also describes Detailing procedures are defined in the **QMS-17 Detailing Standard**.

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7.2 Checking

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The Company has prepared and implemented the **QMS-17 Detailing Standard** to provide for

The procedure describes

Detailing procedures are defined in the

QMS-17 Detailing Standard, which includes

7.3 Control of Approval Documents and Release for Fabrication

The **QMS-21 Control of Approval Documents Procedure** describes the method to The methods include

7.4 Shop Drawings/Documents Supplied by Others

Shop drawings/documents received from the Owner/Buyer are

7.5 Management of Detailing

Detailing Management Connection Consultation and other detailing functions may

Personnel performing Detailing Management are responsible for

Management personnel is qualified by one

or more of the following:

Experience includes

7.6 Detailing Functions

Personnel that detail and/or check shop drawings/documents have

including, but

not limited to,

A qualified Checker

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When applicable, Checkers have

The Company maintains the current references as a library. Detailing procedures are defined in the QMS-17 Detailing Standard.

7.6.2 Connection Consultation

Personnel directing Detailers performing connection detailing are qualified by one or more of the following:

Subcontract Services 7.7

In lieu of employed staff personnel, subcontract services may be used for the following functions:

however, the Company

The Company defines and documents the qualification and selection process for choosing subcontract detailers according to QMS-08 Purchasing Procedure.

Design Procedure 7.8

The Company's design and development process ensures design activities are conducted in a controlled manner that is defined in the QMS-17 Design and Development Procedure, which includes policies for:

A. В. C.

D

Design for Standard Components/Services 7.9

The controls for standard component/services are defined in the QMS-17 Design and Development Procedure.

7.10 Design for Non-standard Components/Services

The controls for non-standard component/services are defined in the QMS-17 Design and Development Procedure.

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8.0 Control of Management System Documents and Project Documents

8.1 Management System Documents

A method has been established and maintained showing the latest revisions and location of the Quality Manual. The controls are defined in the QMS-01 Control of Documented Information Procedure and QMS-02 Configuration Management Procedure.

8.1.1 Quality Management System Documents

											' A		
The	Quality	System	ensure	es									
						This a	ssures	that p	roduc	ced iter	ms		
												records	of
activ	rities are	retained	and r	naintained.	Quality	improve	ements						
											The	System	is
struc	ctured from	om top-d	own u	sing									
												<u> </u>	

8.1.2 Review and Approval of Quality Management System Documents

Internal documents that affect quality are
Revisions to the Quality Manual and other quality management system documents are

Management establishes the frequency and requirements for

Revision controls are defined in the **QMS-02 Configuration Management Procedure**.

8.1.3 Revision Control of Quality Management System Documents

Controlled management system documents that are

The Quality
Manual has a cover page showing the current revision date and the name and location of the
Company. The revision is clearly identifiable on

The Company has established a
method

Documents are controlled so that

The controls for document control and configuration management are defined in the QMS-01 Control of Documented Information Procedure and QMS-02 Configuration Management Procedure.

8.1.4 Access to Quality Management System Documents

Relevant and current procedures and policies pertinent to an area of operation or management are

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The controls are defined in the **QMS-10 Steel Work Procedure**.

8.1.5 Communication of Changes and Revisions to Quality Management System Documents

Changes and revisions to project and quality management system documents are according to the QMS-02 Configuration

Management Procedure and applicable Change Order(s).

8.2 Project Documents

A method has been established and maintained showing

The controls are defined in the QMS-01 Control of Documented Information Procedure and QMS-02 Configuration Management Procedure.

8.2.1 Tracking Project Documents

A *Transmittal Register* and *Contract Log* have

8.2.2 Revision Control of Project Documents

Controlled project documents that are

The revision level of project documents is
The Company has established a method to esign drawings/documents and referenced procedures are identified from the previous revision.

Documents are controlled so that

Documented procedures control

A process has been established to ensure

The controls for document control and configuration management are defined in the QMS-01 Control of Documented Information Procedure and QMS-02 Configuration Management Procedure.

8.2.3 Access to Project Documents

Relevant and current plans, procedures and policies pertinent to an area of operation or management are

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The controls are defined in the QMS-10

Steel Work Procedure.

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9.0 Maintenance of Quality Records

Records are retained and maintained to

Quality records are available

All quality records and final inspections are

All quality records are

Records that document quality typically include:

9.1 Retention of Quality Records

The control of records is defined in the QMS-01 Control of Documented Information Procedure.

9.2 Storage of Quality Records (

Records are controlled to provide

according to the QMS-01 Control of Documented Information

Procedure.

9.3 Retrieval of Quality Records

Proprietary records are

10.0 Purchasing

Purchasing is treated as a process within the Company's quality system according to the **QMS-08 Purchasing Procedure**. The Company accepts responsibility

The Company does not

10.1 Purchasing Data

Purchase documents clearly

including

Purchasing documents for

includes

Purchasing documents include requirements for:

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The supplier evaluation process is fully defined in the QMS-08 Purchasing Procedure.

10.2.1 **Fabrication Subcontractors**

When required by contract, the Company

10.2.2 **Detailing Subcontractors**

The Company performs initial and ongoing evaluation of Detailing Subcontractors according to the QMS-08 Purchasing Procedure.

10.3 Verification of Purchased Product, Materials and Services

The responsibility for quality Documented procedures are established and maintained to Purchased products are The methods used for verification of purchased items are defined in the QMS-09 Receiving Procedure.

Material Receipt Inspection

Materials received are The Receiver The Receiver

Deliveries are checked Defective supplies Nonconforming supplies

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Documented procedures are established and maintained for Deliverable items are

Test reports are

The methods for performing

receiving inspections are defined in the QMS-09 Receiving Procedure.

10.3.2 Customer Verification of Fabricated Product

If specified in the Customer's purchase contract, the Customer or nominated representative is

The methods used for the control of Customer verification are defined in the **QMS-08 Purchasing Procedure**.

10.4 Control of Customer-Furnished Work and Material

A negotiated agreement to verify, store and maintain supplied items is

A documented procedure has been established and maintained Verification includes

The methods for the control of supplied materials are defined in the QMS-10 Steel Work Procedure.

10.5 Purchasing Records

Purchasing documents,

for

are retained and maintained according to the *QMS-01 Control of Documented Information Procedure*. The methods used for verification of purchased items and periodic evaluations of Subcontractors and Suppliers are defined in the *QMS-09 Receiving Procedure*.

11.0 Material Identification

A documented procedure has been established and maintained for

The procedure provides for identification of material as stated in

Purchasing documents for

materials furnished

The filing and retention

Records are retained according to the **QMS-01 Control of Documented Information Procedure.** The methods for the control of supplied materials

are defined in the QMS-10 Steel Work Procedure.

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11.1 Traceability

The Company identifies

according to the QMS-10 Steel

Work Procedure.

12.0 Process Controls

Processes that create a condition where quality of deliverable items cannot be verified through normal methods are which may include

Corrective action is

Procedures and records are maintained that demonstrate

The procedures include

Effective implementation of the following documented procedures is required as a minimum:

- The methods for the control of the fabrication process are defined in the **QMS-10 Steel Work**

12.1 Welding

Procedure.

The Company's welding procedures address

and include:

- •

- •

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The methods used to develop and manage welding operations are defined in the QMS-10 Steel ildwide. Work Procedure.

12.2 Bolt Installation

The Company's bolting procedure QMS-25 Bolting Procedure is compliant with and includes:

The methods used to develop and manage bolt installation are defined in the QMS-10 Steel Work Procedure.

12.3 Material Preparation for Application of Coatings

The Company's prepares material for coating application according to

12.4 Coating Application

The Company applies and cures coatings according to

12.5 Equipment Maintenance

A documented preventive maintenance program QMS-24 Maintenance Procedure is implemented for otherwise, the Company Preventive maintenance activities are It is acceptable to

12.6 Laydown/Assembly

The Company's documented procedure for shop assembly of field connections includes the following:

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13.0 Inspection and Testing

•	J	
To ensure conformance to requirement	ents of deliverable items,	
	Checks occur	
Inspection consists of		
The methods for the	control of the inspection and	d testing process are defined
in the QMS-10 Steel Work Procedu	ire . Nonconforming items are	e controlled according to the
QMS-14 Control of Nonconformanc	es Procedure.	

13.1 Assignment of QC Inspections and Monitoring

QC inspectors are assigned on the basis of

QMS-06 Training Procedure

Non-QC personnel may be assigned to inspection duties under the following conditions:

13.2 Receipt Inspection

Materials received are
The Responsible Authority

The Responsible Authority

13.3 In-Process Inspection

The Company retains and maintains in-process inspection

All materials used

In-process inspections are performed and monitored for processes that include

In-process inspections may

Records of in-process

inspections are maintained and retained according to the QMS-01 Control of Documented Information Procedure.

tn-process inspections are

The following inspections are described in the *QMS-10 Steel Work Procedure*: (revise as required, here and in QMS-10)

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13.4 Final Inspection

Qualified inspectors perform final inspection

Inspection

Molyhide.

records

Records of final inspections are maintained and retained according to the QMS-01 Control of Documented Information Procedure.

13.5 Inspection Records

Inspection records provide

according to the

QMS-01 Control of Documented Information Procedure.

14.0 Calibration of Inspection, Measuring and Test Equipment

Company owned, rented or borrowed measuring and test equipment instruments and devices that are used to determine an item's conformance to specified requirements are

The controls for such equipment and calibration activities are defined in the **QMS-15 Calibration Procedure**.

15.0 Control of Nonconformances

Nonconformances are

Documented procedures are established and maintained for

The methods used to control nonconformances are defined in the **QMS-14 Control of Nonconformances Procedure**.

15.1 Nonconformance with Management Systems

The Company conducts

Nonconformances are also The Company assigns

Responsible Authorities to

according to the **QMS-04 Management Procedure**.

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15.2 Nonconforming Work

When a nonconformance occurs, including complaints, the Company

according to the QMS-13 Corrective Action Procedure and QMS-14 Control

of Nonconformances Procedure. The Company evaluates the need for action to

The Company implements

The Company ensures

The Company retains and maintains records regarding the nature of nonconformances, subsequent actions and results of corrective actions according to the **QMS-01 Control of Documented Information Procedure**.

16.0 Corrective Action

												-	* .				
The	Compa	any I	has	imp	leme	ented	and	m	ainta	ains							
											1.						
The	Compa	any	det	ermi	ines												

Corrective action is applied when

In addition to the preventive measures taken for corrective action requests used to
the corrective action process is used to
The corrective action process is

defined in the QMS-13 Corrective Action Procedure.

17.0 Handling, Storage and Delivery of Materials, Fabricated Work, and Components

According to contractual directives, instructions are

General rules are defined in the *QMS-10 Steel Work Procedure*. Material is

Material is

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Shipments by All Pre. subcontractors are The handling and shipping process is defined in the QMS-11 Shipping Procedure. 18.0 Training All Company personnel are Subsequent training is The Company has implemented a training program that: Appropriate records Management conducts The training program is defined in the QMS-06 Training Procedure. 19.0 Internal Audit 🚜 Internal quality audits are which is accomplished by Audit requirements include The internal audit process is defined in the QMS-12 Internal Auditing Procedure.

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QUALITY MANUAL for STRUCTURAL STEEL ERECTOR Origination Date: (month year)

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Company Location: (your address, city, state, zip)

Abstract:

This document describes the Company's quality management system for structural steel erector

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Paragraphs 5.7.1 through 5.7.3 are "value added" content.

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1.0 Purpose

The purpose of the Quality Management System is

2.0 Scope

The Company's quality management system applies to

This Quality Management System includes

2.1 Exclusions

The Company cites no exclusions to the AISC standard. (revise required)

3.0 References

The latest editions of the following documents and standards are required: (revise as required)



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3.1 Seismic Erection

For the erection of structures requiring the use of ANSI/AISC 341 Seismic Provisions for Structural Steel Buildings, the Company has

meet the requirements of:

•

QMC#:

•

3.2 Metal Deck Installation

When work includes the installation of metal deck, the Company has

Instructions for

metal deck installation are provided in the Erection Plan and the Safety Plan.

(a)

3.3 Bridge Erection

For the erection of bridges, the Company meet the requirements of:

- •
- •

3.4 Safety

Employees and others that perform work for the Company are

which also includes

4.0 Definitions

See **QMS-16 Definitions, Abbreviations and Symbols Procedure** for more details. Subordinate or external documentation is referenced in **Bold Italics**.

5.0 Management Responsibility

The Company is committed to

To ensure this, management

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Policy for Quality and Quality Goals **5.1**

The Company's quality policy defines and pays particular attention to Left blank intentionally

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Quality Management System renumbered from paragraph 5.5

The Company's quality management system is designed to The Quality System ensures This assures Necessary records of activities are retained and maintained. The System is

Company maintains

All Managers are responsible for

The quality system

documentation is comprised of

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Management Review 5.3

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Management review meetings	s are	
	riews are reported and records are retained and review are defined in the QMS-04 Managemen	
Management review meeting		
	e QMS-12 Internal Auditing Procedure to en	Records of
management review meetings QMS-01 Control of Docume	s and internal audits are retained and maintained ac nted Information Procedure.	cording to the
Responsible Authorities also which includes:	J'	
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5.4 Responsible Quality Personnel

The individuals designated as Quality (QM) and/or Safety Manager (SM) understand

Each Responsible
Authority (RA) is

The RA's have primary responsibility for

The RA ensures

5.5 Resource Management

The Company has the resources
The responsibility, authority and the interrelation of personnel

that includes

5.5.1 Personnel

Personnel assigned to key positions and those performing field operations provide according to the *QMS-06***Training Procedure**. Unless otherwise noted personnel can

Specifically, individuals responsible for Quality and Safety management do not Qualified personnel are assigned to manage the following functions revise as required)

- •

5.5.2 Buildings, Workspace, Equipment and Associated Utilities

The facility consists of

The areas and buildings are

Adequate space is provided for are defined in the QMS-22

Application of Complex Protective Coatings Procedure.

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Ambient conditions are provides

QMC#:

The facility also

5.5.3 Erection Tools and Equipment

The Company has under its control

5.6 Internal Communication

To ensure proper communication between and throughout all levels of employees within the Company, internal communication is

which is documented in the QMS-04 Management Procedure.

Management periodically communicates with employees to

Employees are encouraged to

This system requires management to

5.7 Quality Manual

The primary purpose of the Quality Manual and QMS Procedures is

The Quality Manual has been developed by

The quality manual is approved

The quality manual is

Additional procedures and work

instructions

For instance:

- •

5.7.1 Organization

Review meetings are

Reviews are

reported and records are retained and maintained. The controls for management review are defined in the **QMS-04 Management Procedure**.

The organizational chart

which are further defined in the QMS-05 Responsibilities and Authorities

Procedure.

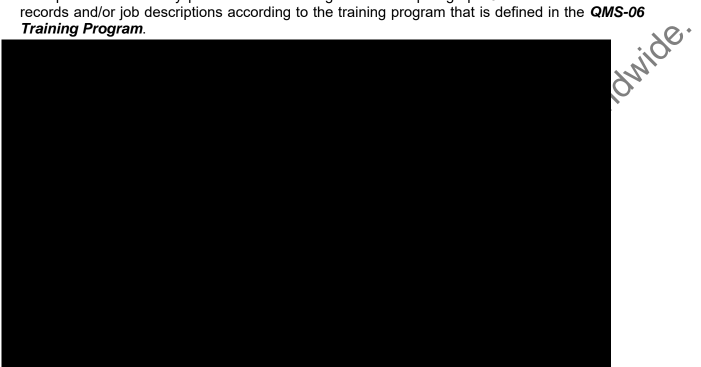
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The qualifications of key personnel and managers listed in paragraph 5.4.1 are maintained in records and/or job descriptions according to the training program that is defined in the QMS-06 Training Program.



5.7.2 Approval

This manual is issued under the authority of

Management ensures the QMS is

5.7.3 Order of Precedence Value-Added

The order of precedence of order-specific documentation is

- 1.

Safety Manual

The Company ensures Employees according to the QMS-03 Construction Safety Program, QMS-04 Management Process and QMS-06 Training Program.

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a material at the fall accions informations.	vhich
	dail
5.9 Policy for Safety Executive management is responsible for	
5.9 Policy for Safety	
Executive management is responsible for The policy for safety includes:	
Executive management according to the QMS-04 Management Process Procedure . Safety goals	are
5.10 Responsible Safety Personnel	
Executive management designates The designated management representative for sadoes	afety
The designated management representative(s) has the ability, responsi	ibility
8,	

6.0 Construction Document Review and Communication

The Company performs contract and project specification review according to the **QMS-07 Proposal Development and Contract Review Procedure**. The review as well as

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The review considers The procedure provides
to assure
The Company communicates according to <i>QMS-21 Approval of Approval Documents Procedure.</i>
Communications include:
according to QMS-21 Approval of Approval Documents Procedure. Communications include: Decisions made in the process of these communications are
Decisions made in the process of these communications are
Contract review records may
Contract review records may
Project requirements are distributed to production and performance records are retained and maintained according to contract requirements and <i>QMS-01 Control of Documented Information Procedure</i> . The controls for contract review are defined in the <i>QMS-07 Proposal Development and Contract Review Procedure</i> .
Communications with Authorities Having Jurisdiction (AHJ) are documented by using the
Request for Information (RFI) form (or your form). The RFI is
A number is assigned to the <i>RFI</i> and then recorded in the <i>RFI Log</i> . The <i>Log</i> documents the following:
6.1 Customer Requirements

The Company captures

Contract Review process.

as part of the **Proposal Development &**

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Documents are controlled to Documents are

The Company

determines

QMC#:

which is defined in the QMS-07 Proposal Development & Contract Review Procedure.

7.0 Reserved - N/A

8.0 Control of Management System Documents and Project Documents

8.1.1 Quality Management System Documents

The Quality System ensures

This assures

Necessary records of activities are retained and maintained. Quality improvements

structured

The System is

8.1.2 Review and Approval of Quality Management System Documents

Internal documents that affect quality are Revisions to the Quality Manual and other quality management system documents are

Revision controls are defined in the QMS-02 Configuration Management Procedure.

8.1.3 Revision Control of Quality Management System Documents

Controlled management system documents that are

The Quality Manual has a cover page showing the current revision date and the name and location of the Company. The revision is

The Company has established a method to ensure

Documents are controlled

The controls for document control and configuration management are defined in the QMS-01 Control of Documented Information Procedure and QMS-02 Configuration Management Procedure.

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8.1.4	Access	to	Quality	Manag	gement	Sys	stem	Docum	ents
-------	---------------	----	---------	-------	--------	-----	------	--------------	------

Relevant and current Company policies, procedures, safety requirements and project

8.1.5 Communication of Changes and Revisions to Quality Management System Documents Changes and revisions

are

QMC#:

according to the QMS-02

Configuration Management Procedure and applicable Change Order(s)

8.2 Project Documents

A method has been established and maintained

The controls are defined in the QMS-01 Control of Documented Information Procedure and QMS-02 Configuration Management Procedure.

8.2.1 Tracking Project Documents

have been established to

which indicate

8.2.2 Revision Control of Project Documents

Controlled project documents that are

The Company has established a

method to ensure

Documented procedures control

A process has been established to

The controls for document control and configuration management are defined in the QMS-01 Control of Documented Information Procedure and the QMS-02 Configuration Management Procedure.

8.2.3 Access to Project Documents

Relevant and current

The controls are defined in the QMS-10

Steel Work Procedure.

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9.0 Maintenance of Quality Records

All quality records and final inspections are

All quality records are

Records that document quality typically include:

9.1 Retention of Quality Records

The control of records is defined in the *QMS-01 Control of Documented Information Procedure*.

9.2 Storage of Quality Records

Records are controlled according to the QMS-01 Control of Documented Information Procedure.

9.3 Retrieval of Quality Records

Proprietary records are

10.0 Purchasing

Purchasing is **08 Purchasing Procedure**. The Company

The Company does not

10.1 Purchasing Data

Purchase documents

Purchasing documents for includes

The purchasing process is fully defined in the QMS-08 Purchasing Procedure.

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10.2 Selection of Subcontractors and Suppliers

The purchasing process

QMC#:

The supplier evaluation process is fully defined in the Subcontractors

QMS-08 Purchasing Procedure.

10.2.1 Fabrication/Erection Subcontractors

When required by contract, the Company

10.3 Verification of Purchased Product, Materials and Services

The responsibility for quality of subcontracted products

Documented

procedures are

Purchased products are

The methods used for verification of purchased items are defined in the **QMS-09 Receiving Procedure**.

10.3.1 Material Receipt Inspection

Materials received are

The Receiver

Deliveries are

Defective supplies are

Nonconforming supplies are

Documented procedures are established and maintained for Deliverable items

When certification test reports are

The methods for performing receiving

inspections are defined in the QMS-09 Receiving Procedure.

10.4 Control of Customer-Furnished Material

A negotiated agreement

The QMS-10 Steel Work Procedure has been established and

maintained for

Verification includes

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The methods for the control of Customer-furnished materials are defined in the QMS-10 Steel Work Procedure.

10.5 Purchasing Records

Purchasing documents,

QMC#:

are retained and maintained according to

the QMS-01 Control of Documented Information Procedure.

10.6 Customer Verification of Product

the Customer or nominated representative is

The methods used for the control of Customer verification are defined in the QMS-08 Purchasing Procedure.

11.0 Material Identification

A documented procedure has been established and maintained for identifying

The procedure provides for

Purchasing documents

includes

The filing and retention

Records are retained according to the QMS-01 Control of Documented Information Procedure. The methods for the control of supplied materials and identification of deliverable items are defined in the QMS-10 Steel Work Procedure.

12.0 Erection Process Control

Processes may include Corrective action is Procedures and records are including The procedures include

Effective implementation of the following documented procedures is required as a minimum: evise as required)

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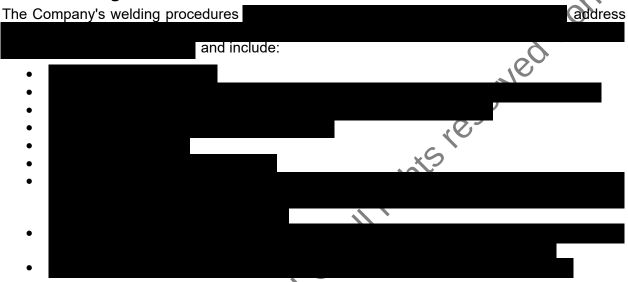
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The methods for the control of the erection process are defined in the **QMS-10 Steel Work Procedure**.

12.1 Welding

QMC#:



The methods used to develop and manage welding operations are defined in the *QMS-10 Steel Work Procedure*.

12.2 Bolt Installation

The Company's bolting procedure QMS-25 Bolting Procedure is and includes:

The methods used to develop and manage bolt installation are defined in the **QMS-10 Steel Work Procedure**.

12.3 Material Preparation for Application of Coatings

The Company's prepares material for coating application according to the coating manufacturer's recommendations, product data sheets and project specifications.

12.4 Coating Application

The Company applies	according to	
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12.5 Equipment Maintenance

12	.5 ⊑quipi	Herit Maii	Renance				
Α	documented	preventive	maintenance	program	QMS-24	Maintenance	Procedure is otherwise, the
Cor	mpany						
				Preve	entive main	itenance activiti	es are
				1 1010	mare man		
						It is accept	able to
							,O
12	6 Loydo	wn/A 222n	nhly			-, Je	9
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The	Companys	documented		the follow	ing:	40	
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13	0 Insne	ction and	d Testing	<i>'</i> Ο.			
	-		_	of the apr	olicable en	ection project,	
	Onouro com	omanoo to	requirements	an the app		ecks occur	
		nspection co	ongista of				
		rispection co		methods for	or the cont	rol of the inspec	tion and testing
							s are controlled
acc	ording to the	QIVIS-14 Co	ntrol of Nonco	ontormand	es Proced	iure.	
13	.1 Assigr	nment of (QC Inspect	ions and	d Monito	oring	
	inspectors a					á	according to the
	S-06 Trainir nstructi on p er	ng Procedur	e,		under t	ne following con	ditions:
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13.2 Receipt Inspection

Materials received are
The Responsible Authority

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The Responsible Authority The methods for performing receiving inspections are defined in the **QMS**wide. 09 Receiving Procedure.

13.3 In-Process Inspection

The Company retains and maintains in-process inspection plans

All materials used in the

work product are

QMC#:

Applicable inspection instructions indicate In-process inspections are

In-process inspections

are

In-process inspections

may

Records of in-process inspections are maintained and

retained according to the QMS-01 Control of Documented Information Procedure.

In-process inspections are

The following inspections are described in the QMS 10 Steel Work Procedure: (revise as Aties, L.C. required, here and in QMS-10)

13.4 Final Inspection

Qualified inspectors

Inspection

records identify

Records of fibal inspections are maintained and retained according to the QMS-01 Control of Documented Information Procedure.

13.5 Inspection Records

Inspection records provide

Inspection records are retained and maintained according to the

QMS-01 Control of Documented Information Procedure.

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14.0 Calibration of Inspection, Measuring and Test Equipment

Company owned, rented or borrowed measuring and test equipment instruments and devices

The controls for such equipment and calibration activities are defined in the **QMS-15 Calibration Procedure**.

15.0 Control of Nonconformances

Nonconformances are

QMC#:

Documented procedures are established and maintained for

The methods used to control nonconformances are defined in the *QMS-14*Control of Nonconformances Procedure.

15.1 Nonconforming Quality Management System

The Company conducts

according to the QMS-12 Internal Auditing Procedure. The Company assigns

according to the QMS-04 Management Procedure.

15.2 Nonconforming Work

When a nonconformance occurs,

the Company

according to the QMS-13 Corrective Action Procedure and QMS-14 Control of Nonconformances Procedure. The Company evaluates

The Company implements

The Company ensures

The Company retains and maintains records

according to the QMS-01 Control of

Documented Information Procedure.

16.0 Corrective Action

√the Company has implemented and maintains

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Corrective actions are applied

Adaptive action according to QMS-04

Management Process Procedure,

Corrective action is applied when:

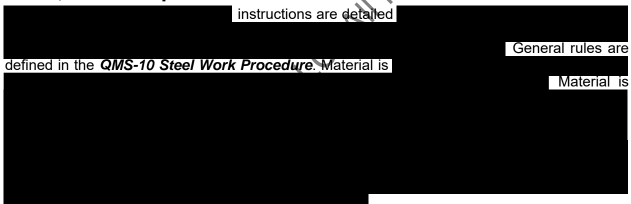
The Company determines

In addition to the preventive measures

The corrective action process is defined in the QMS-13 Corrective Action

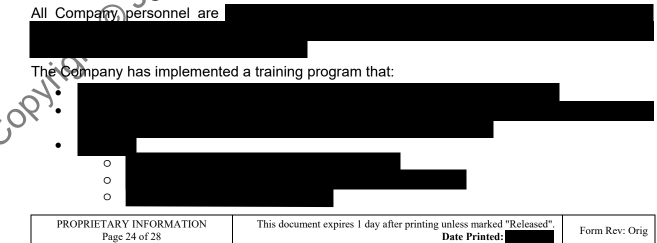
Procedure.

17.0 Handling, Storage and Delivery of Materials, Fabricated Work, and Components



The handling and shipping process is defined in the *QMS-11 Shipping Procedure*.

18.0 Training



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Management conducts . The training program is defined in the QMS-06 Training Program.

19.0 Internal Audit

Internal quality audits are which is accomplished by Audit requirements include The internal audit process is defined in the QMS-12 Internal Auditing Procedure.

20.0 Erection Plan

The Company prepares

The erection plan includes the following information as appropriate for the project:

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The erection plan is All io All revisions are 21.0 Safety Plan The Company prepares a safety plan The safety plan may A safety plan considers The safety plan includes The safety plan includes the following information as appropriate for the project: The safety plan is and is

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All revisions are

QMC#:

Prior to the start of the erection project, the Company retains and maintains documentation other evidence)

Important:

23.0 Safety Management System

23.1 Documentation Requirements

The QMS-03 Safety Program contains the following information:

23.2 Safety Training

Safety training includes

Safety training includes

The safety plan described in 21.0 is

√he Company provides training according to

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Date:	Latest Revision Date
Project: 5	Customer, Unique ID, Part Number
Document Status:	Draft, Redline, Released, Obsolete
Document Link:	Location on Server (if used)

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1.0 PURPOSE OF DOCUMENT AND RECORD CONTROL

This procedure defines the requirements for the control of documents within the quality management system (QMS). 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. A record is

3.0 DOCUMENT TYPES

- 3.1. Quality Manual: this document provides
- 3.2. QMS Procedures: these documents provide
- 3.3. General Work Instructions: these documents provide
- 3.4. Inspection Instructions: these documents are
- 3.5. Forms: these documents are

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Control of Documented Information

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3.6. Records that are created for temporary retention of miscellaneous information are

4.0 QUALITY MANUAL

4.1. Creating the Quality Manual

The Quality Manual has been developed by top management of the Company, which includes the Company's Vision and Governing Policies.

4.2. Review and Approval

The Quality Manual is reviewed and approved by top management before release. Approval is indicated by

4.3. Distribution

The Quality Manual is distributed electronically through the Company's internet server.

The Document Control Center may retain older hardcopies or softcopies for historical purposes, but these are

In some cases, a hardcopy of the Quality Manual may be given to an employee, department or Customer. If the document is needed for more than thirty (30) days it is marked "Released" and dated with the month and year of release by the Responsible Authority (RA).

Each employee must

4.4. Change Control

Any employee may request a change to the Quality Manual. Requests for changes may be made by

5.0 QUALITY MANAGEMENT SYSTEM PROCEDURES

5.1. Creating New QMS Procedures

QMS procedures should be created as soft files (MS Word, etc.). It is recommended that files of a similar type

5.2. Review and Approval

QMS Procedures are to be reviewed and approved by top management. At least one member of top management that is responsible for reviewing the document should be responsible for the area affected by the document. Approval is indicated by

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5.3. Distribution

QMS procedures 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 historical purposes, but these are

In some cases, a hardcopy of the procedure may be given to an employee, department or Customer. If the document is needed for more than thirty (30) days it is marked "Released" and dated with the month and year of release by the Responsible Authority (RA). Each employee must

5.4. Change Control

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

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 quality related 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 soft files (i.e., MS Word, etc) and then submitted to the Configuration Control Board (CCB) for review and approval. Work instructions should include, as applicable:

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 and subject to the controls of the Configuration Management Procedure - not this procedure. 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 historical purposes, but these are

In some cases, a hardcopy of the work instruction may be given to an employee, department or Customer. If the document is needed for more than thirty (30) days it is marked "Released" and dated with the month and year of release by the Responsible Authority (RA). Each employee must

6.4. Change Control

Changes to general work instructions are performed in the same manner as the Quality Manual. When general work instructions are changed, the revision history table is updated and the revision indicator advanced.

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7.0 INSPECTION INSTRUCTIONS

7.1. Creating New Inspection Instructions

New inspection instructions are developed by or under the supervision of the Quality Manager using requirements from

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.

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 historical purposes, but these are

In some cases, a hardcopy of the inspection instruction may be given to an employee, department or Customer. If the document is needed for more than thirty (30) days it is marked "Released" and dated with the month and year of release by the Responsible Authority (RA). Each employee

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 Quality Manager. 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

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 notify the Responsible Authority of the approval by providing one software copy of the form for upload onto the Company's internet server and/or intranet in the current forms directory. It is the appropriate manager's

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responsibility to

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,

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 approval as originals but must have their revision indicator advanced.

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

9.2. Third party specifications and engineering drawings, including those of the Customer, are controlled according to the **QMS-02 Configuration Management Procedure**. 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 filed as needed to request changes or updates.

11.0 CONTROL OF RECORDS

11.1 The controls for each type of record are defined in *Appendix A* of this procedure.

The listed "controller" must ensure their assigned records

11.3 Records for active contracts are maintained in the quality department handling the operations. Records are removed from the active files at

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11.4	The Document Control Center maintains archive files for records. Records shall be maintained a minimum of
11.5	Records that are discarded after retention shall be
11.6	Hardcopy records are to be stored in suitable cabinets that prevent damage or deterioration When archived records are stored elsewhere,
11.7	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.
11.8	Records are verified for
11.9	The Company does not require vendors to maintain records for the Company; instead,
11.10	To ensure protection of records, electronic records are subject to
11.11	Local computer data that is stored on company computers must
11.12	When making corrections to written record entries, the error is
	Correction fluid or correction tape is not to be used on any quality records.

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APPENDIX A: RECORD RETENTION MATRIX

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Required Record or Document Type	Company Record	Controller	Type	Location	Minimum Retention
Calibration records	Calibration		Form		70.
Contract review records	Contract review		Form		
Control of Nonconformances	RFS		Form	6	
Corrective actions	RFS		Form	70	
Design change records	Engineering order		Form	Sel	
Design input records	Engineering order		Form G	0	
Design review records	Engineering order		Form		
Design validation records	Construction inspection		Form		
Design verification records	Production inspection	C).	Form		
First Article Inspection	First article		Form		
Internal audit records	Internal audit	•	Form		
Lost, damaged or unsuitable Customer property	Customer property		Form		
Management review meeting minutes	Management review report		Form		
Record of realization process	Engineering order		Form		
Record of release of product	Construction inspection		Form		
Supplier evaluation	Supplier review		Form		
Traceability records	Construction inspection		Form		
Training records	Training record		Form		

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CONFIGURATION

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		on Date: XXXX
	Document Identifier:	Configuration Management
	Date:	Latest Revision Date
	Project: 5	Customer, Unique ID, Part Number
	Document Status:	Draft, Redline, Released, Obsolete
SPE	Document Link:	Location on Server (if used)
ment describes config		

This document describes configuration management procedures.

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DOCUMENT CHANGE RECORD

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1.0 PURPOSE

This procedure defines the requirements for the management of the configuration of products produced by the Company's configuration management activities include the following:

The following are not governed by this control procedure:

2.0 THEORY

Part configuration includes a variety of aspects of a given part, including its shape, function, internal components, chemical analysis, raw materials, suppliers used and more. Because a given product may change over its life, typically due to design improvement activities or customer requirements, it is important to maintain control and records over changes.

This dramatically improves future design and production efforts.

This procedure has been developed based on practices defined in ISO 10007 and MIL-STD-973.

3.0 CONFIGURATION DOCUMENTATION

3.1. The current configuration of a given part is identified through applicable technical documents. These may include, but are not limited to:



3.2. All such technical documents are developed by Engineering and approved by the CCB. (See section 4.0) They are then controlled according to this procedure.

3.3. The baseline documentation is entered into a database that maintains current data for every configuration item. As new configuration items are generated, approved and placed in the release system, they are

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3.4. Configuration documents and Customer intellectual property received by Contracts is

4.0 CONFIGURATION CONTROL BOARD (CCB)

4.1. The Responsible Engineering Authority (REA) and Quality Manager serve as the Configuration Control Board, which has full authority and responsibility for

4.2. The Chairperson of the CCB is any specified member dependent upon the circumstance. The Customer may be invited to attend CCB meetings.

4.3. The CCB serves as the point of authority to resolve all program configuration management questions at all levels of activity, e.g.,

4.4. CCB responsibilities include:

OB responsibilities include:

5.0 BASELINE MANAGEMENT

5.1. The Company may establish a configuration baseline to identify and create the initial configuration identification of deliverable supplies at specific times during the contract cycle. The baselines provide

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5.2. contai	All descriptions of the baselines used to state product performance and design requirements are ned in configuration documents.
5.3. 5.3.1.	For configuration management purposes, four major baselines may be required as discussed below.
3.3.1.	Pre-Nelease Dasellile.
5.3.2.	Functional Baseline:
ć:	At the Functional Baseline, the
config	uration management system is operating and the released documents have described the following:
•	
•	
5.3.3.	Allocated Baseline:
include	These
•	
•	
5.3.4.	Product Baseline.
This b	
I nis b	aseline prescribes:
•	
This !	accline and approved changes conve as the configuration reference point for all subsequent resistance
	aseline and approved changes serve as the configuration reference point for all subsequent reviews. ed technical documents may be used if

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5.4. **Baseline Maintenance**

Once established, the baselines serve as the approved departure points for updating by incorporation of changes that have been approved by the CCB. The baselines plus the approved changes represent the product configuration at any point in time. Configuration documents are

The release system is shown in Figure 1, which...,

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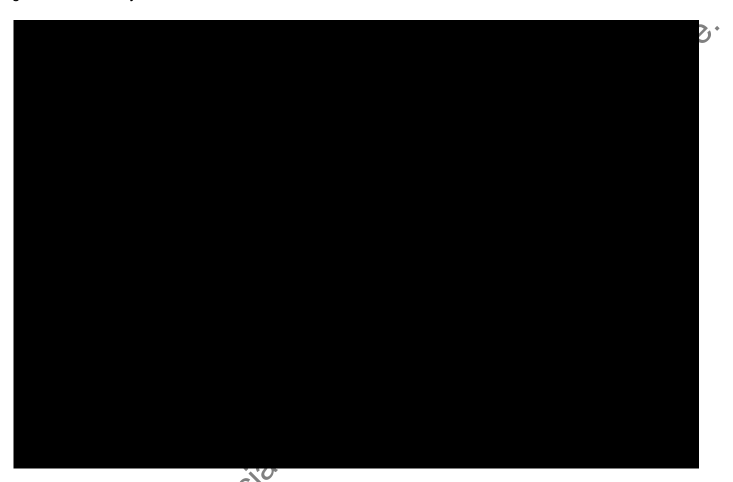
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Figure 1: Release System Flowchart



5.5. Document approval is indicated by any of the following methods:

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5.6. The Document Control Center prepares the release package after

6.0 CONFIGURATION CHANGE CONTROL

6.1. Configuration change control is the process of maintaining the baseline identification and regulating all changes to that baseline. The 'as-designed' technical documentation must equal

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6.2. confia	Change control is vested in the Configuration Control Board. Any employee may request a change to a puration. All proposed changes to the baseline documents are
6.3. anoth	Joint change control authority is established where any program shares a commonly identified item with er program.
6.4.	Evaluations of changes include the consideration of
6.5.	The evaluation will take into consideration
6.6.	All associated changes and affected hardware items or computer programs are
6.7.	Types of Configuration Change ges to the configuration are implemented after approval of engineering changes, deviations or waivers.
	efinition for each is as follows:
6.7.1.	Engineering Change:
672	Deviation:
0.7.2.	Deviation.
673	Waiver:
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6.8.	Change Classification
	ges in configuration are classified by the CCB as either Class I or Class II. The change classification ned by the CCB is entered on the Engineering Order, which

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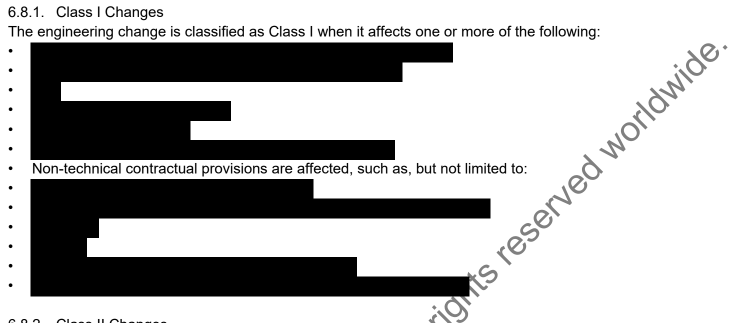
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6.8.1. Class I Changes

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The engineering change is classified as Class I when it affects one or more of the following:



6.8.2. Class II Changes

Any change that does not fall within the Class I definition is a Class II change. Class II changes are

- 6.9. Change Implementation
- 6.9.1. All approved changes are implemented under the guidance of the configuration management function.
- 6.9.2. Configuration Management maintains approval records for all configuration changes.

These records identify

6.9.3. The Quality Group verifies that changes have been incorporated into affected units and that the associated configuration status records have been revised.

6.9.4. Superseded revision levels of electronic documents are

6.9.5. During the evaluation of the ECP, EO or RFS, the CCB determines what implementation actions are required to accomplish the approved change and affected areas are identified.

6.9.6. The CCB provides a complete description of the effort required to accomplish the approved change. The definition of the actual tasks required is

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6.9.7.	Deviation:						
6.9.8.	Waiver:						
6.9.9.	Supplemen	nt Releases:					
6.9.10					(6		
6.9.11	. Proposed	Class I engine	ering changes are ange Proposal (E	approved by	the CCB and a pineering Orde	re submitted to th	e Customer in
A Clas	ss I Enginee	ering Change is	s not		j	(20) 30 10 9 311	
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Figure 2: Change Control Flow



6.9.12. Re-identification Practices Q

Part numbers are changed whenever complete item interchangeability is not possible for all products shipped and for all current and future products. When complete item interchangeability is not possible,

6.9.13. All deliverable items are fabricated and assembled according to the configuration defined by the appropriate engineering drawing and its authorized changes.

6.9.14. No oral instruction or other random or unwritten authority is accepted in place of formal change control (see the Baseline Management section herein). Redlined technical documents may be used if

7.0 SUBCONTRACTOR AND VENDOR CHANGES

7.1. Only those subcontractors having a funded design effort are permitted to implement Class I or II changes with submittal to the Company for review and concurrence or non-concurrence in classification.

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- 7.2. For all vendors used by suppliers, proposed changes to baseline documents are
- 7.3. Suppliers and vendors are controlled according to the

8.0 MANAGEMENT DIRECTIVES

- 8.1. Management members of the CCB/MRB issue their binding policies, procedures and directives to personnel within their exclusive organization in the form of a Bulletin.
- 8.2. The Bulletin is completed as required by individual format. The Bulletin is the only accepted form of correspondence for intra-company and inter-company requests for work to be performed or when providing instruction for performing work. The signed and completed Bulletin is

9.0 CONFIGURATION RECORDS AND REPORTS

The following lists are revised as required to include the latest configuration status of listed documents. Dependent upon contract requirements, records and reports may include:

9.1.	1. Numerical lists:	
9.2.	2. Indentured Lists:	
	68	
9.3.	3. As-Built Parts List	
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9.4.	4. EO Status	
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9.5.	5. Data Lists:	

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9.6.	Configuration Account Record for Integrated Systems:		
			0
9.6.1.	Configuration Item Identification Report:		
0.00	As Britters As Basiness I Confirmation	NI	
9.6.2.	As-Built vs. As-Designed Configuration:		

10.0 PRODUCT AND TEST SOFTWARE CONTROL

Production of software for integration into deliverable products is controlled according to

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CONSTRUCTIONS erved worldwide. SAFETY PROGRA. SAFETY PROGRAM

Origination Date: (month year)

Revision Level (Orig, A, B, C, etc) Revision Date: (month and year) Released By: (your issuing authority or EO#) Jak SPE

Abstract:

This document describes the Company's safety program.

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1.0 RESPONSIBILITIES

1.1 SAFETY DIRECTOR

Education/Orientation:

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Enforcement:

Execution of Work:

•

Inspection/Correction:

• Insure that any reported unsafe condition, hazard or potential hazard will be:

Safety Meetings/Training:

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Execution of Work:

•

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Hazard Communication:		46/2
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Injuries/Accidents:		SO

Inspection/Correction:

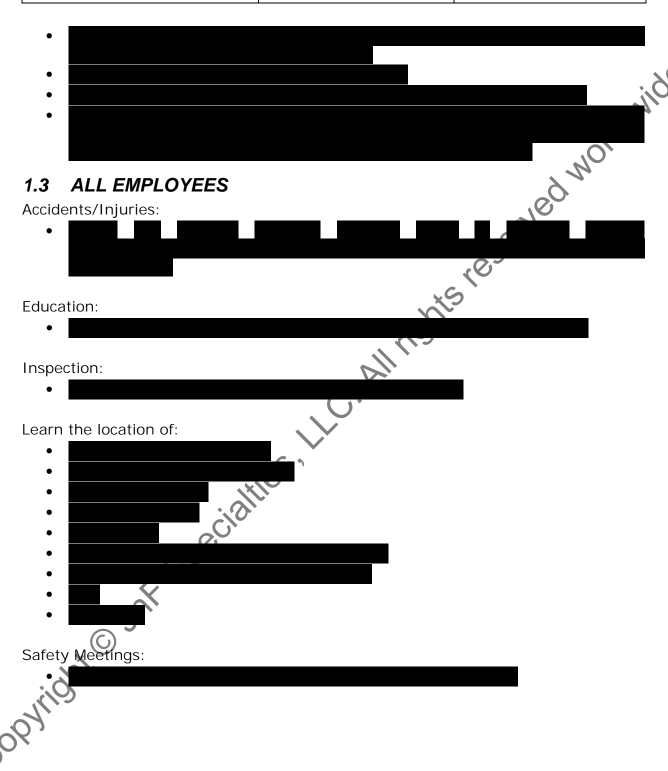
0 Reporting:
Following procedures and/or Contractor procedures, investigate and report all:

Safety Meetings/Training:

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2.0 SAFETY RULES

2.1 General

Alcohol/Illegal Drugs:



Emergency Procedures and Facilities:

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Hazard Reporting:

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Inspection of Equipment:

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Know the location of:

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operating Equipment:

• Equipment:

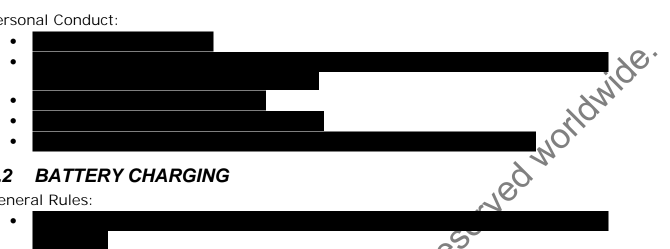
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Personal Conduct:

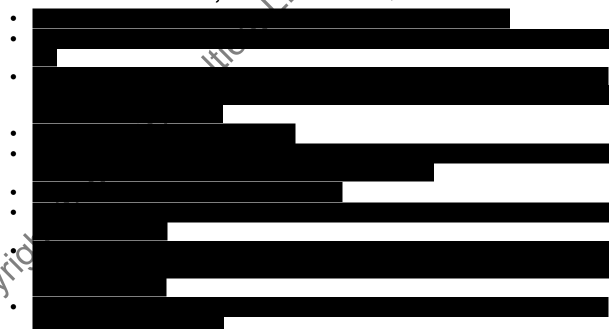


BATTERY CHARGING 2.2

General Rules:



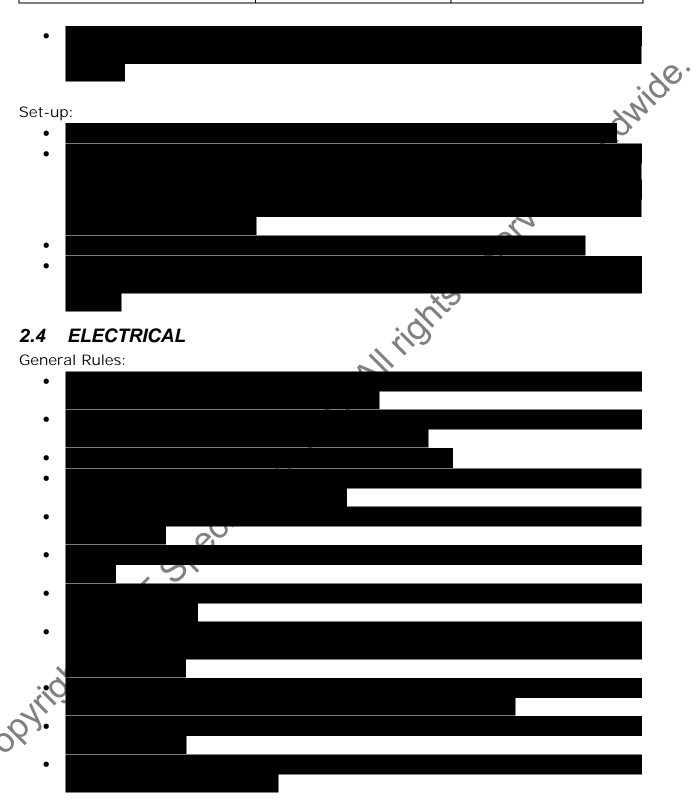
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2.5 Floor	FALL HAZARDS and Wall Openings:		25
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Other	Fall Prevention Rules:	().	
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Other	Safety Devices to Prevent Falls:	•	
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2.6

FIRE PREVENTION AND PROTECTION Learn what the right type of extinguisher is for different types of fires: Fire Extinguishers: 0 Housekeeping: Other Safety Precautions: Storage Facilities:

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Storage Locations:

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GOOD HOUSEKEEPING

General Housekeeping:

HAND AND POWER TOOLS Tools: 2.8

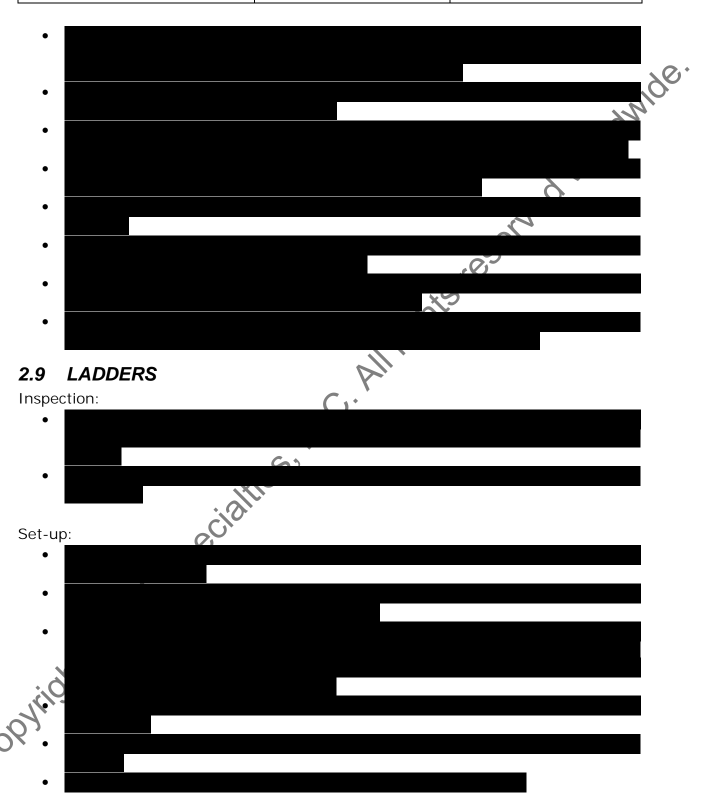
Hand Tools:

Power Tools:

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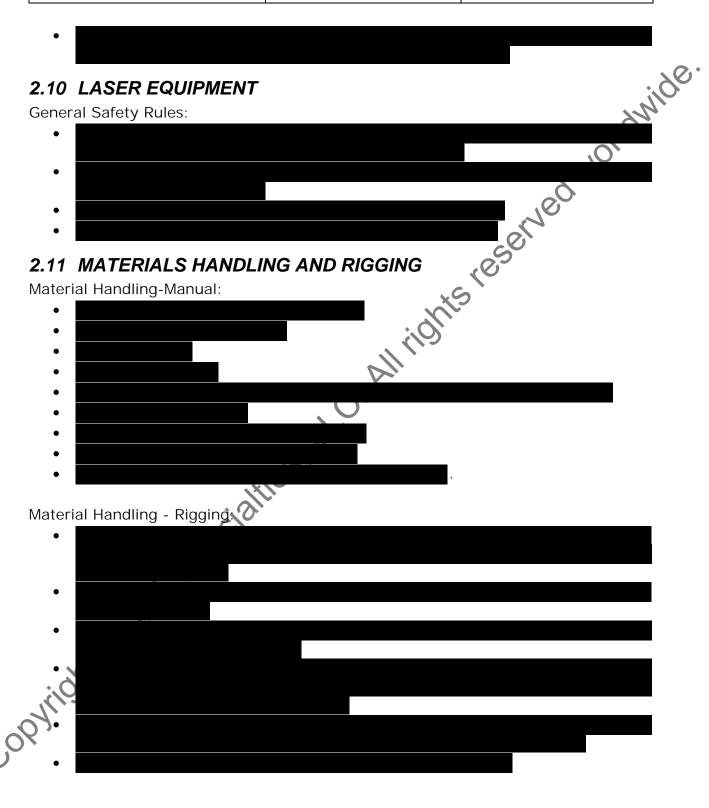
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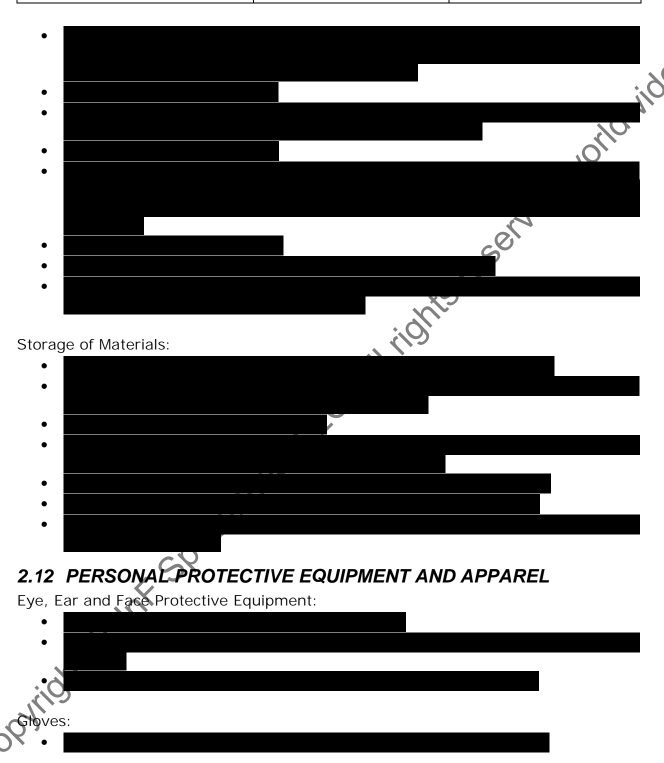
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Shoes:

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2.13 SCAFFOLDS

General Safety Information:

2.14 SIGNS, SIGNALS AND BARRICADES

Signs, Signals and Barricades:



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Bracket Scaffold: Orlichi Mobile Scaffolds: Tubular Welded Frame Scaffold (Safeway Type)	•					
Mobile Scaffolds: •	•					
Mobile Scaffolds: •	Bracket Scaffold:					digh
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Tubular Welded Frame Scaffold (Safeway Type)	Mobile Scaffolds:					
Tubular Welded Frame Scaffold (Safeway Type)	•			J.		
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	Tubular Welded Frame Sca	affold (Safe	າ way T	ype)		
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Tyro-point Suspension (Swinging) and Single-point Suspension (Spider-type) Scaffold:	Tyro-point Suspension (Scaffold	(Swinging)	and	Single-point	Suspension	(Spider-type)
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2.15 WELDING AND CUTTING

Fire Prevention:

Portable fire extinguishers shall be provided at all locations where welding of the control o

Inspection/Use of Equipment:

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Personal Protection:

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Storage/Placement of Equipment:



3.0 FALL PROTECTION

The Company has implemented a fall protection plan to protect personnel from falls. The Company is firmly committed to the health and safety of all individuals on our job sites as well as complying with all applicable safety standards. This program allows us to

	. 00	
	DESCRIPTION	APPLICATION
PART 1	OSHA SUBPART R	ALL PROJECTS
PART 2	FALL PROTECTION STANDARDS AND REQUIREMENTS	ALL PROJECTS
PART 3	SPECIFIC FALL CRITERIA	ALL PROJECTS
PART 4	FALL PROTECTION	PROJECT SPECIFIC
PART 5	SENRAC SUBPART R – STEEL ERECTION (DRAFT ONLY)	PROJECT SPECIFIC
PART 6	OWNER REQUIREMENTS –	PROJECT SPECIFIC

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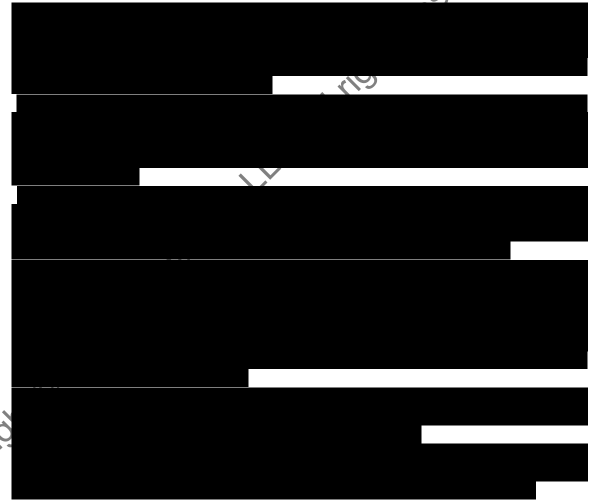
PART 1 - OSHA Subpart R

PART 1.1 1926.70 FLOORING REQUIREMENTS

(a) Permanent flooring - skeleton steel construction in tiered buildings.

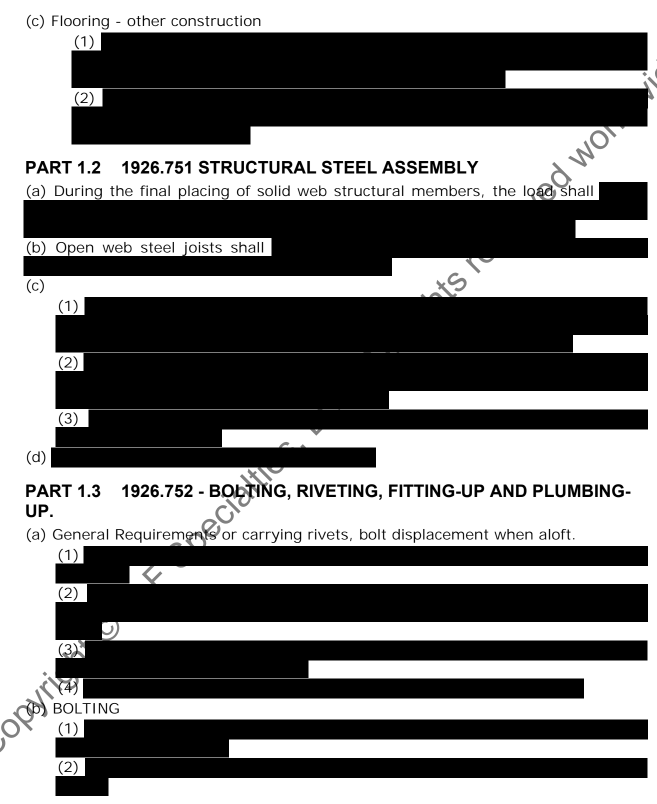


(b) Temporary flooring - skeleton steel construction in tiered buildings.



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	ROTECTION STANDARDS AND REQUIREMENTS
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Employee Qualifica	LIUII3

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General Site Conditions	
Ladders	
Lifts	, 110,
Material Staging	
Minimize Employees	
Narrow or Small Surfaces	
Personal Fall Protection Equipment	
Precast and miscellaneous steel	
Prefabricate	
Recognition	

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Safety Program

Safety Continuing Education		
Secured Members		1974
Site Specific Pre-Construction N	Meeting	9 W
Steel/Joist	(8)	
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Tools and Equipment		
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PART 3 - SPECIFIC FALL CR	ITERIA	

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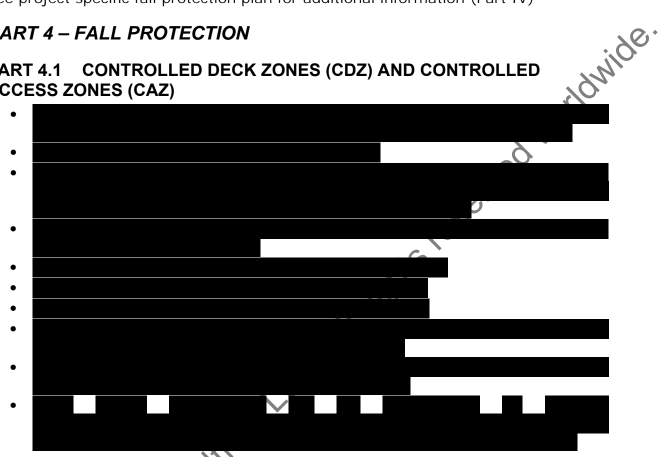
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See project specific fall protection plan for additional information (Part IV)

PART 4 – FALL PROTECTION

PART 4.1 CONTROLLED DECK ZONES (CDZ) AND CONTROLLED **ACCESS ZONES (CAZ)**



FALL PROTECTION SYSTEMS **PART 4.2**



Fall protection systems such as warning lines, controlled access zones and safety monitors, may be utilized in controlled work environments provided the following is established:



The above items to be addressed in site specific safety plan.

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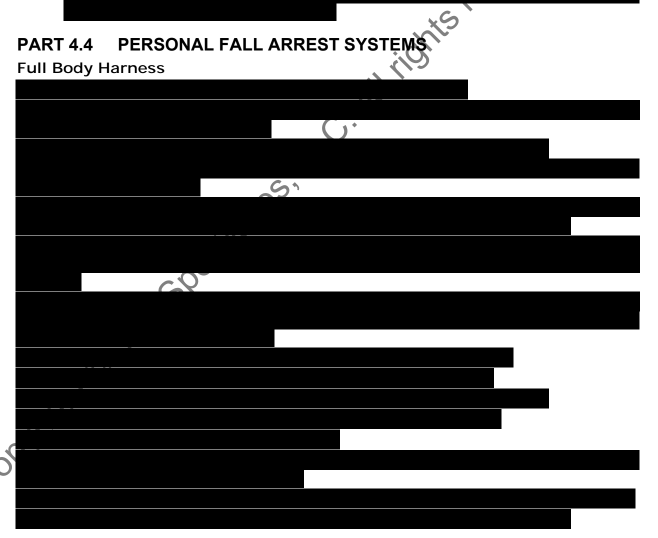
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PART 4.3 GUARDRAIL SYSTEMS [PER OSHA.1926.502 (9) & (6)]



PART 4.4



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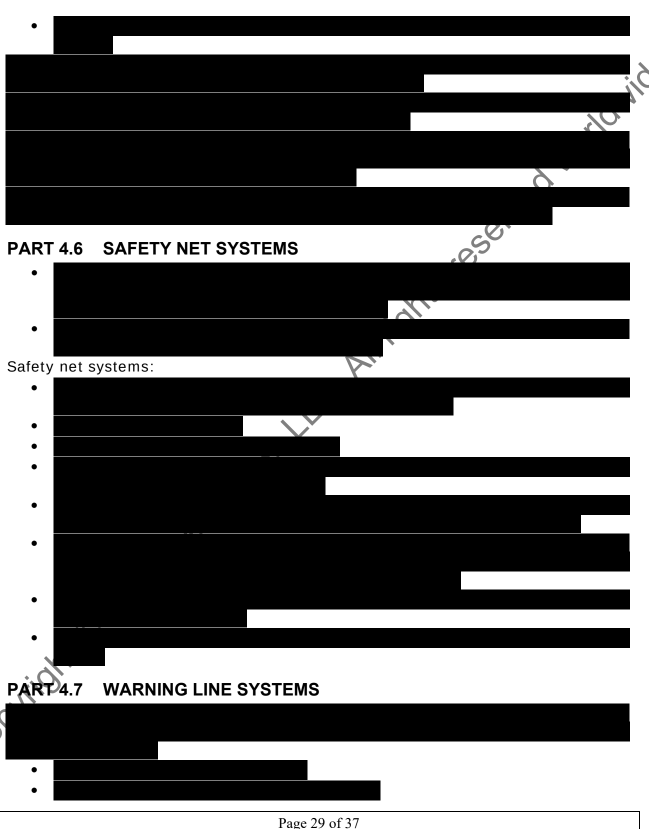
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Retractable Lifelines	
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Horizontal and Vertical Lifelines	Ne
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Positioning Devices	
PART 4.50 SAFETY MONITORING SYSTEMS	
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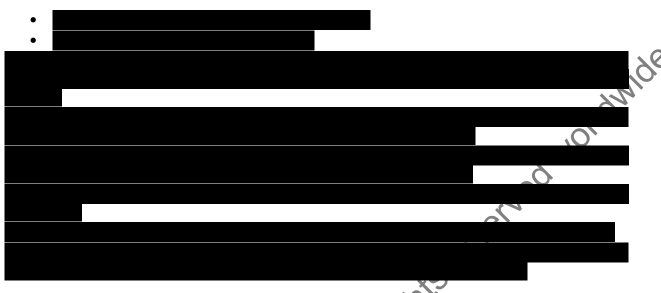
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PART 5 - SENRAC (DRAFT ONLY)



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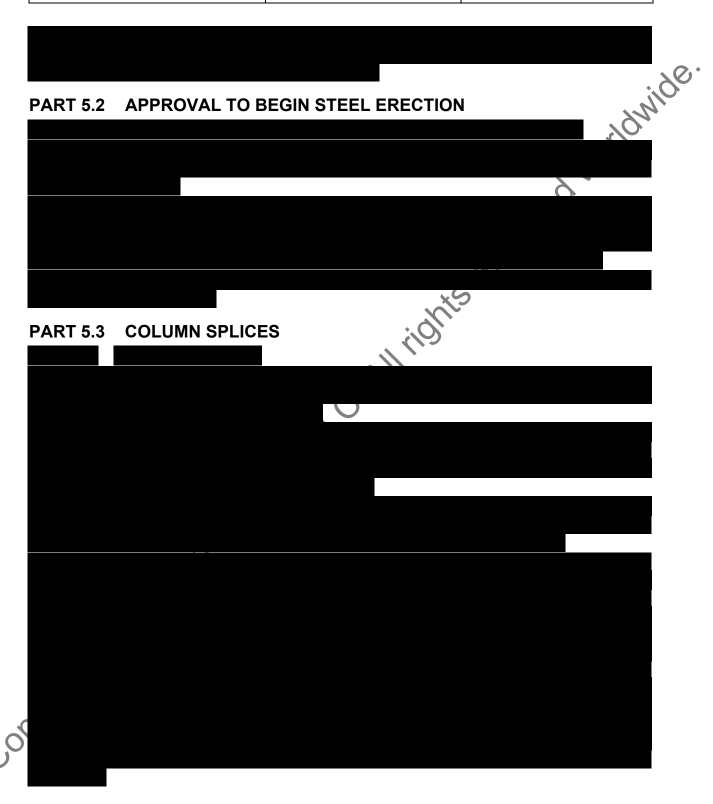


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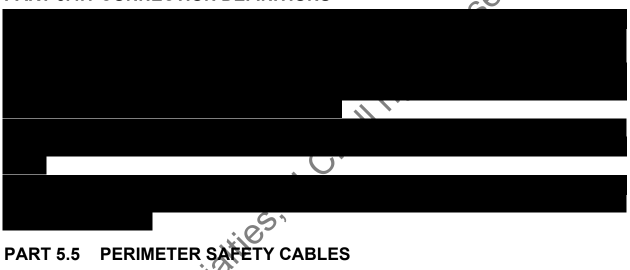
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PART 5.4 DOUBLE CONNECTIONS

1926.756 Beams and columns



PART 5.4.1 CONNECTION DEFINITIONS



1926.756 Beams and columns



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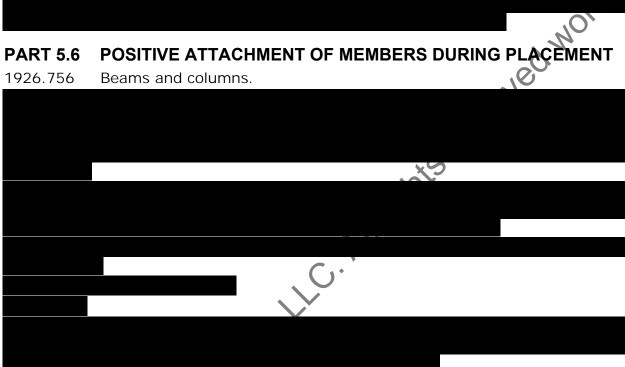
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1926.760 Fall protection

PART 5.6

1926.756

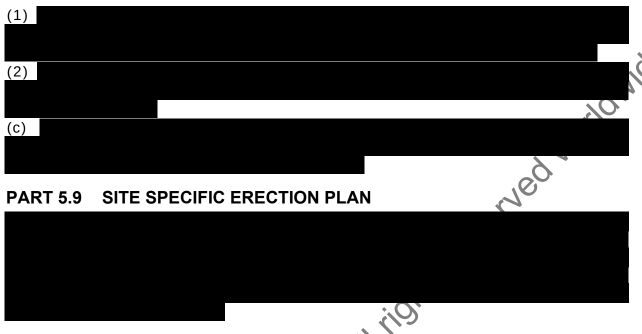


ROOF AND FLOOR OPENINGS **PART 5.7**

PART 5.8 SITE LAYOUT AND ACCESS

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PART 5.10 SLIPPERY SURFACES

1926.754 Structural steel assembly (c) Walking/working surfaces



PART 5.11 STRUTJOIST BOTTOM CHORD STABILIZER PLATE

1926.757 Omen web steel joist.

PART 5.12 TRIPPING HAZARDS

1926.754 Structural steel assembly. (c) Walking/working surfaces.

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4.0 HAZARDOUS COMMUNICATIONS

4.1 LABELING



4.2 OSHA INSPECTIONS



4.3 WHAT IS HAZ-COM?

"Right to Know"

Hazard Communication, Haz-Com or "Right to Know" all refer to

Here is a partial list of materials, considered hazardous, common to construction sites:



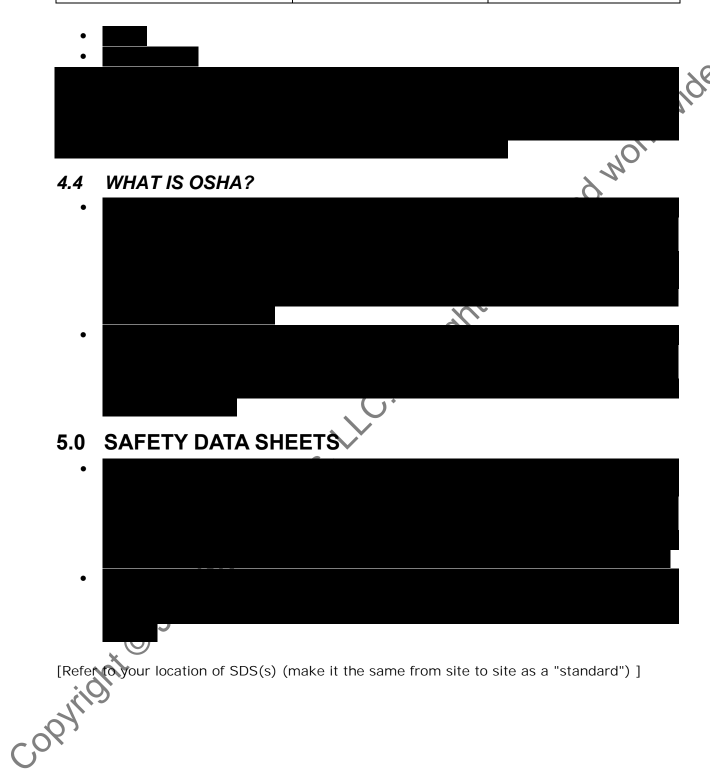
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This document describes the management review process.

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Management	Process
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1.0 PURPOSE

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

2.0 THEORY

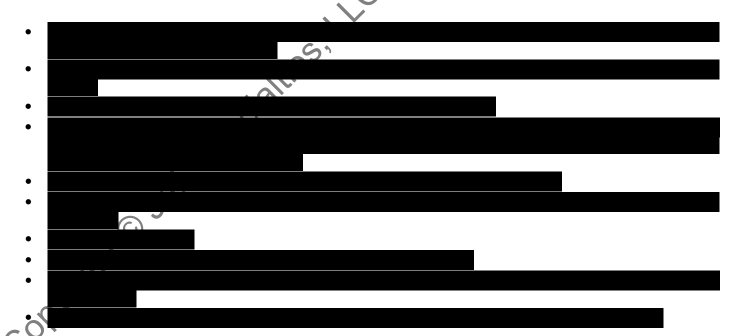
The Company believes in "intelligent management," which enables the Company to make decisions based on facts, data and verifiable evidence. Intelligent management reduces the need to make decisions based on personal opinion, whims or mood and ensures results of decisions are measurable.

3.0 MANAGING AS A PROCESS

The Company recognizes that it has to manage processes identified in the Quality Management Policies and Procedures handbook; however, management itself must also be treated as a process. This means that management activities must have inputs, outputs, controls and reaction plans (when things do not work out as expected.) The Company must consider the results of analyses and evaluations and the outputs from management reviews to determine if there are needs or opportunities to be addressed as part of continual improvement.

The process map in the Appendix of this document identifies how Management is treated as a process and provides an overview of how management is performed.

Management is responsible for implementation and application of the following QMS requirements:



PROCEDURE: MANAGEMENT REVIEW

4.1 The management of the Company performs formal management review of the Quality Management System a minimum of The minimum attendance for Management Review

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Your Company Name

Management Process

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4.2	This review shall include
4.3	Minutes of the meetings are
4.4	The Management Review meeting should include analysis of the following inputs:
•	
•	
•	
•	
•	
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4.5 of	Management shall use action items or the corrective action system to take recorded actions as a result. This includes
	See the QMS-13 Corrective Action Procedure.
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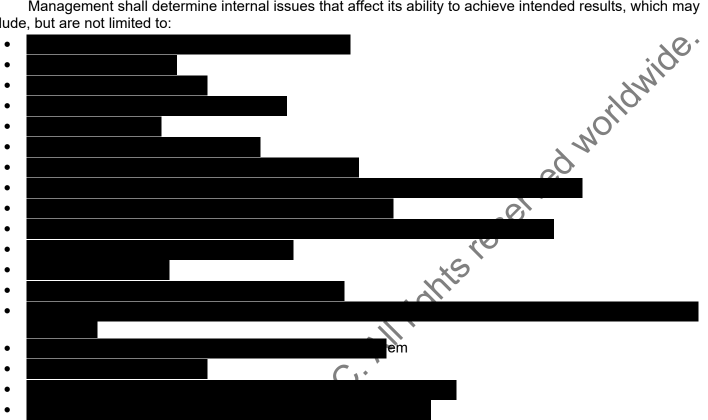
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4.6 Management shall determine internal issues that affect its ability to achieve intended results, which may include, but are not limited to:



Management shall determine external issues that affect its ability to achieve intended results, which 4.7 may include, but are not limited to:



PROCEDURE: MEASURING AND MONITORING PROCESS **OBJECTIVES**

Each process identified in the Quality Management System has at least one objective. The objective is

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5.2	Each process objective must
5.3	Top management will Throughout the year, assigned managers and staff will
5.4	Throughout the year, assigned managers and staff will
5.5	During Management Review the data will
5.6	When a process does not meet a goal, corrective action shall
	(O)
5.7	The current metrics, standings, previous goal and revised goals shall
5.8 deteri	Over time, management shall assess performance of each process against the goals as a means of mining if continual improvement is being made. If not
6.0 6.1	PROCEDURE: INTERNAL and EXTERNAL COMMUNICATION Internal communication is an important facet of the way the Company does business. By this we mean
0.1	internal communication is an important facet of the way the company does business. By this we mean
The fo	ollowing methods are used for internal communications:
•	
•	
•	
•	
6.2	External communications that are relevant to the quality management system must be limited to
6.2.1	Confidential Company Information
	pany Employees must not reveal Confidential Company Information to External Parties except
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Your Company Name

Management Process

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6.2.1.1 Basic Company Information

Company Employees must not communicate Basic Company Information to External Parties excep

Only Authorized Responsible Authorities may communicate about the Company or its business, or C. All rights re communicate as a representative of the Company, with any of the following External Parties:

Only Authorized Responsible Authorities may communicate about the Company or its business or communicate as a representative of the Company on

6.2.1.2 Written Company Information

All Written Company Information must conform to

All Written Company Information must be approved by the appropriate Responsible Authority before it is communicated to any External Party.

With respect to any Written Company Information regarding new business, clients, or other contract counterparties, or other Third Parties with a business relationship with the Company, care must be exercised to

Written Company Information regarding material transactions, contracts, or other significant corporate events or circumstances, or prepared in response to requests from governmental or regulatory bodies, must



Your Company Name

Management Process

Rev: Orig

7.0 PROCEDURE: RESOURCE MANAGEMENT

7.1 The management of resources is a critical component to the management activities of the Company. Resources requiring such management includes:



7.2 Like other management activities, resource management must be based on data.

7.3 To manage resources, top management must

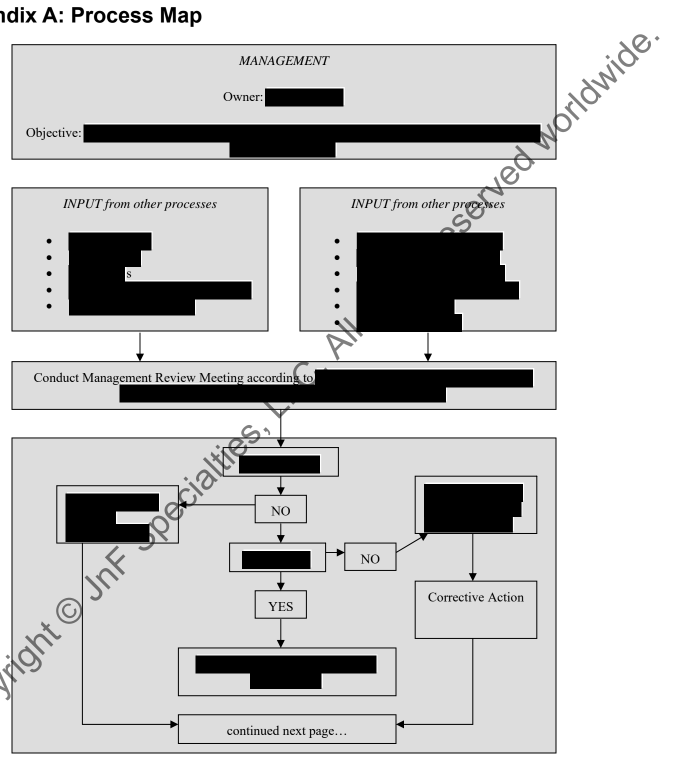
7.4 During Management Review, managers shall

7.5 From that data, top management can

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Appendix A: Process Map



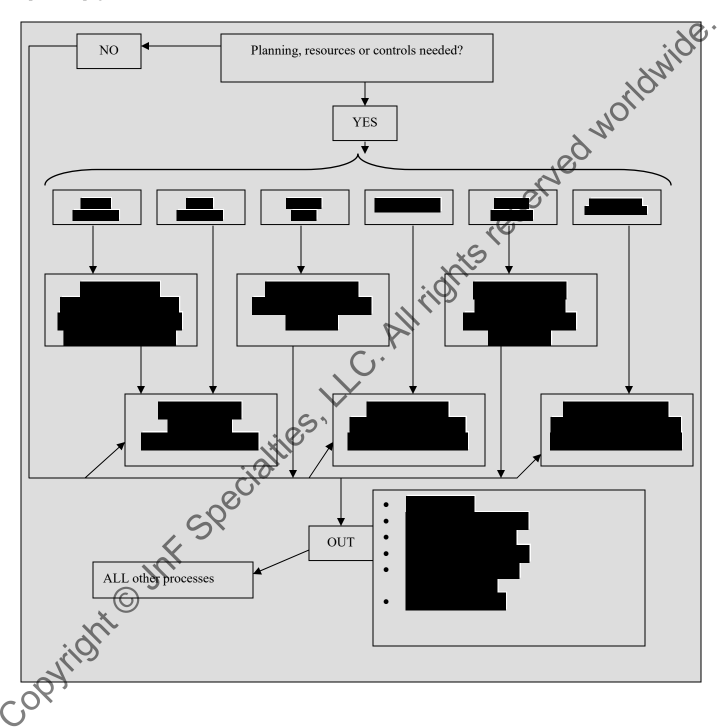
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Abstract:

This document describes responsibilities and authorities of Company personnel.

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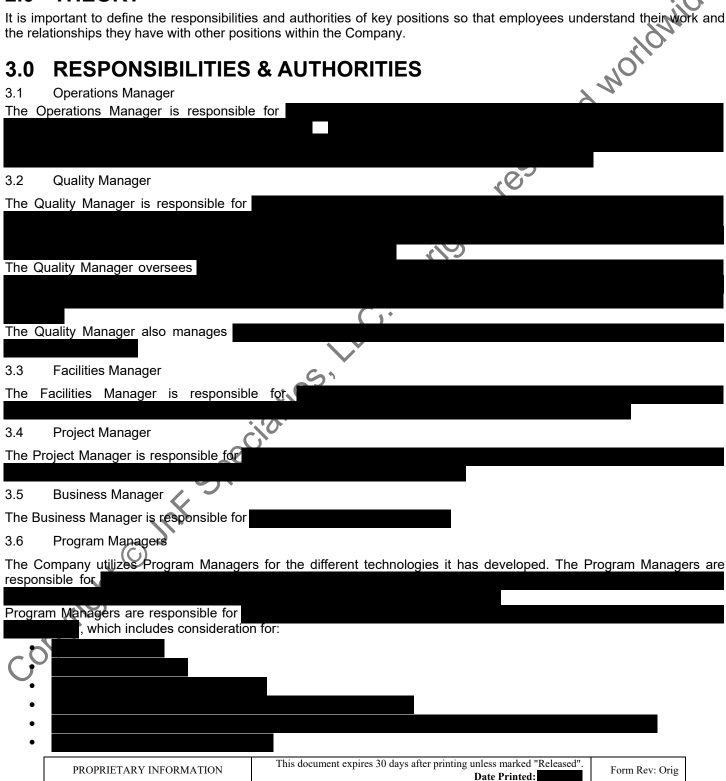
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Vour Logo	(Vous Company)	Responsibilities and Authorities
Your Logo	(Your Company)	Rev: Orig

PURPOSE 1.0

This document provides an overview of the responsibilities and authorities for key positions within the Company.

2.0 THEORY



Your Logo (Your Company)

Responsibilities and Authorities

Rev: Orig

The Administrative Assistant is responsible for

3.8 Accounting Manager
The Accounting Manager is responsible for
3.9 Environmental Health & Safety Manager
The EHS Manager is responsible for

3.10 Quality Group Staff & Inspectors (including Receiving)
The Quality Group includes

3.11 Construction Operators
Construction operators include

3.13 Shipping Personnel

3.7

Shipping personnel are responsible for

Internal Auditors are responsible for

Administrative Assistant

3.14 Human Resources Staff

Human Resource staff is responsible for

3.15 Purchasing Staff

Purchasing staff is responsible for

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Abstract:
This document describes training program and requirements.

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3	3.4 Documented Training Program	
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PURPOSE 1.0

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 served. of abilities and on-the-job training to enhance those abilities.

TRAINING PROCEDURE 3.0

3.1 Hiring

Employees are hired on their basis to best meet the requirements for the position

To accomplish this, potential candidates are compared against the requirements of the QMS-05 Responsibilities and Authorities Procedure as well as job descriptions for the open position. These job descriptions typically

Initial Indoctrination and Orientation 3.2

Once hired, new employees are assigned to their position and undergo initial indoctrination and orientation. This introduces the employee to

On the Job Training 3.3

Once an employee has completed initial indoctrination they undergo on-the-job training relative to their position. This training is specific to the area and equipment on which they work and is typically

Documented Training Program 3.4

3.4.1 Personne responsible for functions that affect quality receive initial and periodic documented training. including, but not limited to, project managers, detailers, inspectors, welding personnel, fitters, and painters.

3.4.1.1 Training is specific to the function or activities related to the job description, such as

3.4.2 Personnel providing training shall have appropriate training or experience in the subject they are teaching. Training course outlines include

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Additional Training 3.5

At the discretion of management, training may be conducted at any time, which may be necessitated by

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Date:	Latest Revision Date
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Abstract:

This document describes the procedures used to review contracts and develop proposals.

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Proposal Development and Contract Review

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1.0 PURPOSE

This document defines the Proposal Development and Contract Review process including or making reference to procedures for the various activities within the process.

2.0 **THEORY**

The Company can only meet Customer requirements by ensuring that all such requirements are obtained from the Customer, then reviewed and understood. This process ensures the suitable capture of requirements and ensures that the Company's understanding of those requirements is communicated to the Customer prior to and through contract acceptance.

3.0 **PROCEDURE**

When addressing Customer needs and industry trends, the Company consider

Documentation is not required for contract review and proposal development for Customers that purchase items on a recurring basis or when the dollar-value of the purchase order is

The Company determines its capability to meet Customer requirements by:

a)				
b) establishing the c	riteria for:	(),		
1)				
2)				
c)				
d)				
e) determining, retai	ning and maintaining	required records that de	emonstrate:	
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Proposal Development and Contract Review

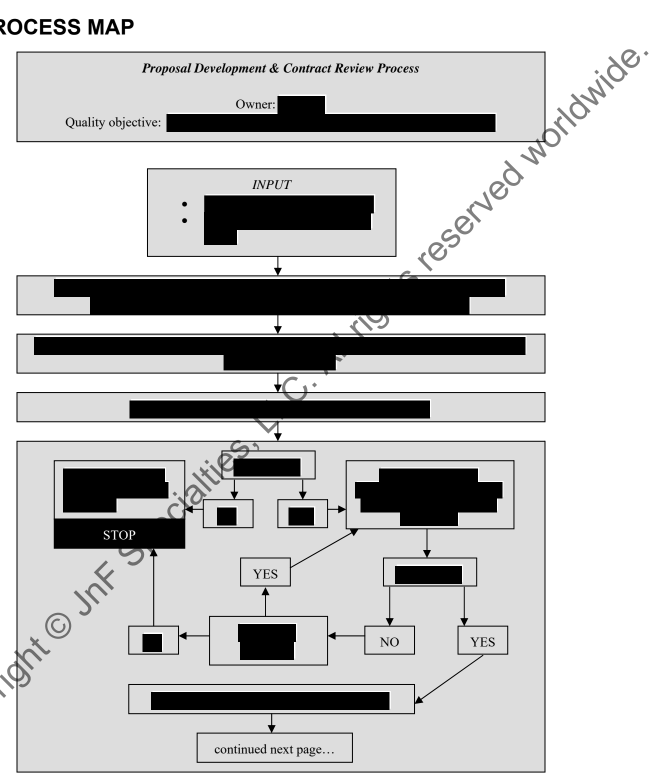
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4.0 PROCESS MAP

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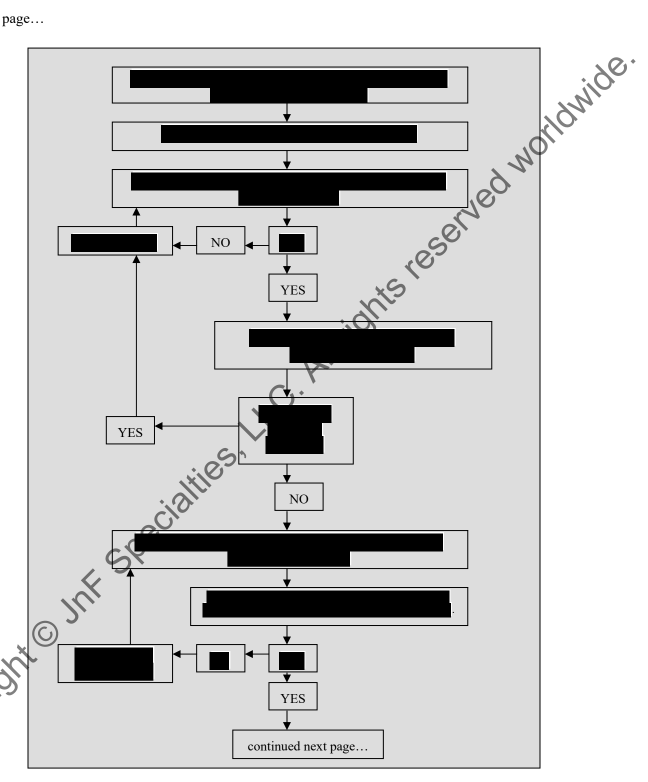


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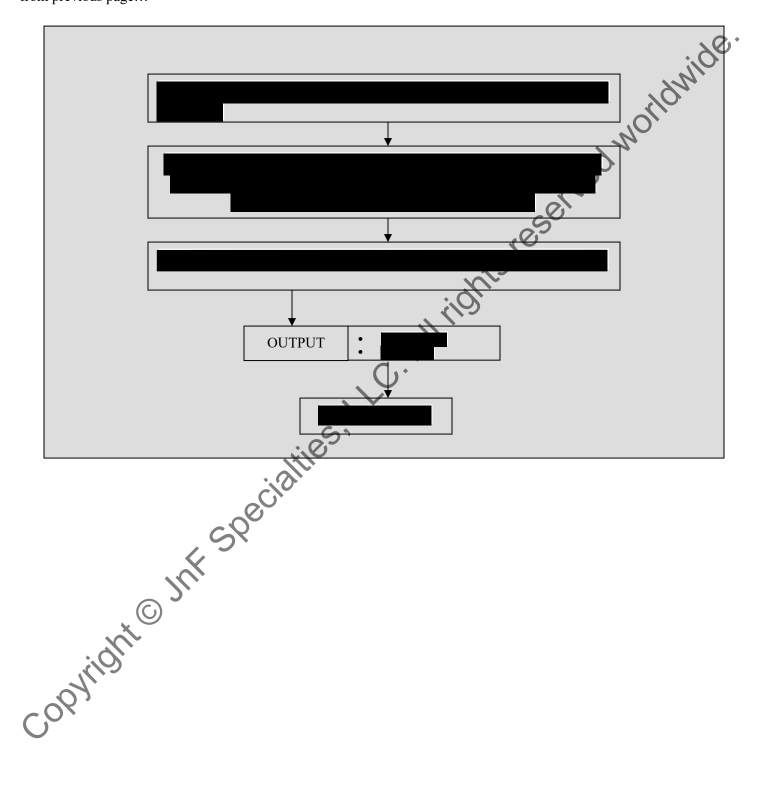


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Abstract:
This document describes the work instruction for reviewing purchase order content.

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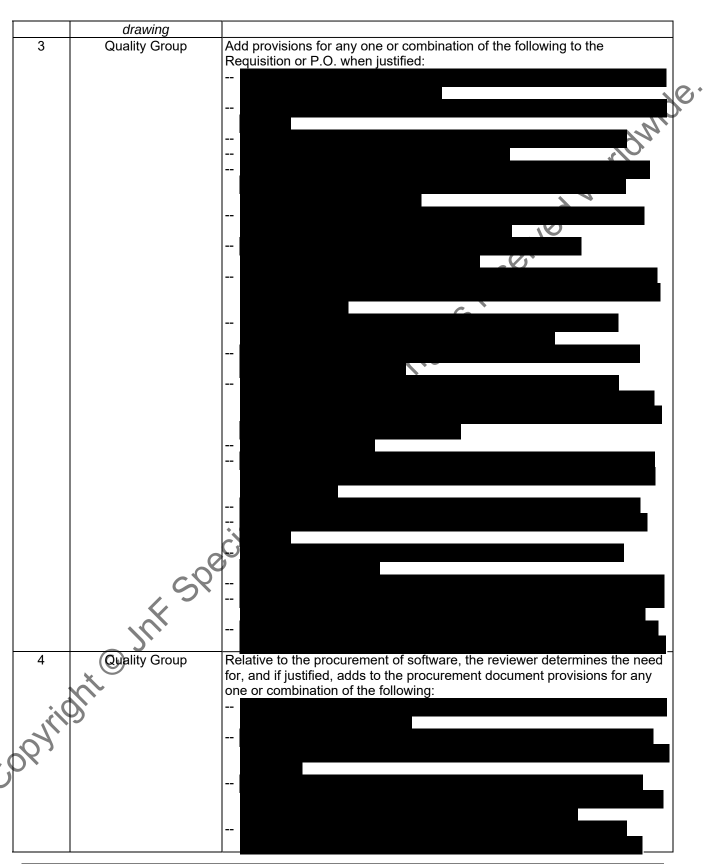
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1	Quality Group	The reviewer determines the need for, and if justified, imposes the requirements of Supplier Quality Requirements to the Requisition or
		P.O. Complete the Used-On and Contract# sections on the cover page of the
		PO
		Used-On = J/N or Program Acronym; Contract# = P.O.#
		Check-off applicable requirement boxes on Requisition
2	Quality Group	Forward Requisition to
		Check mark the appropriate field in the "Type of Certs" section; multiple
		types of Certs may be required Verify Raw Material Requirements are recorded on Requisitions, <i>except</i>
		verily Naw Material Nequilients are recorded on Nequisitions, except
		Supplier should be evaluated a south of the Complian Fusion to
		 Suppliers should be evaluated according to the Supplier Evaluation Determine if a Supplier has been designated by the Customer - notify
		Purchasing when a sole-source Supplier is designated by the Customer
		Add known OA requirements to the requisition for entry on the BO:
		Add known QA requirements to the requisition for entry on the PO; such as
	IF	THEN
2.1	Older Revision	Contact the applicable Project Engineer and process the Requisition
	Supply Required	
2.2	Requisition is marked	
	"Under Revision"	
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	X	
0.0	(A) M () 1	
2.3	A Raw Material Requirement is not	
	Specified	
2.4	Deviation to drawing is	Validate each exception by examination of the applicable drawing or
3/1	noted on Requisition	specification to determine the appropriateness of the deviation, e.g.,
76,	such as "Less Note" Deviation to drawing is	
Y T	noted on Requisition	
	such as "Less Note"	
2.5	Order in for construction	
2.5	Order is for construction activity without	
	reference to engineering	
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5 Discrepancy in Requisition or P.O. 5.1 Supplier Quality Requirements applies 5.2 P.O. requires additional conditions related to inhouse processing 5.2.1 P.O. requires additional conditions related to inhouse processing 5.2.2 Requisition or P.O. Ok 6 Quality Group Forward Supplier Evaluation to the Supplier; perform required follow-up routines.	Requisition or P.O. 5.1 Supplier Quality Requirements applies			
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Abstract:
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S	PROCESSING REQUISITIONS AND PURCHASE ORDERS	

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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 construction 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 construction or services affects everything we make. As a result, it is important to monitor and control the quality of both construction and services that we receive as well as the suppliers of such construction and services.

3.0 PROCEDURE: SUPPLIER EVALUATION AND SELECTION

- 3.1 All suppliers of construction related materials or services must be evaluated unless these suppliers are:
- 3.2 Supplier evaluation is conducted by following the formation the Supplier Evaluation Form.
- 3.3 The Supplier Evaluation Form ensures
- 3.4 Once approved through the Supplier Evaluation Form, the Quality Manager 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 Quality Manager will

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3.8 Using the results from combination of the following functions for product suppliers, the Quality Manager will determine if the Supplier should be increased in rating to		

3.9 For suppliers providing product, incoming inspection results are recorded on the Subcontractor Performance Rating Spreadsheet, which calculates the Supplier's current quality rating based on parts received and parts accepted. A new Supplier that rates 100% on their first delivery may be upgraded to 3.10 If a new Supplier rates

3.11 If any Supplier rates less than

3.12 If items are returned to any Supplier

3.13 Any Supplier may be de-rated to

3.14 Management may override

3.15 During management review, the entire Approved Supplier List is subject to

4.0 PROCESSING REQUISITIONS AND PURCHASE ORDERS

4.1 During review of each requisition, the Quality Group will

4.2 Responsible Authorities take into consideration

4.3 Responsible Authorities ensure the adequacy of requirements prior to their communication to a Supplier, which includes:

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•	
•	When appropriate the purchase order defines acceptance criteria for
•	ajo.
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1.4	When appropriate, the purchase order defines acceptance criteria for
4.5	As applicable, purchase order information includes:
a)	
0)	
c)	
1) roa	uirements relative to:
. TCq	difficitis relative to.
e)	
·)	
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g)	
	C.C.
4.6	The requirements for delegation are defined when
1.7	When the Company or its Customer needs to perform verification activities at a Supplier facility
	(C)
4.8	See the process map herein.
4.0	Fundamental Community of the Community will prothesize the shift forester and/on the
4.9 nainte	Emergency Purchasing Authority: The Company will authorize the shift foreman and/or the enance foreman emergency purchase authority for
5.0	OTHER PURCHASING RULES
5.1	In all instances, the Purchasing Department will

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5.2	Any employee of the Purchasing Department that has any financial or other interest in a supplied
comp	pany, either directly or through any member of his/her immediate family, shall
- 0	The acceptance by myselectical measured of wife an evertible from a small one is
5.3	The acceptance by purchasing personnel of gifts or gratuities from suppliers is
5.4	The acceptance of items intended for the purpose of advertisement and bearing the name of the
Supp	lier is
5.5	The Purchasing department will cooperate with Customer-related activities and will
5.6	The Purchasing department will not,
5.0	The Turchasing department will not,
5.7	The Company will abide by all Government clauses or other statutory or regulatory requirements as
	enced by the order, contract or other requirements document.
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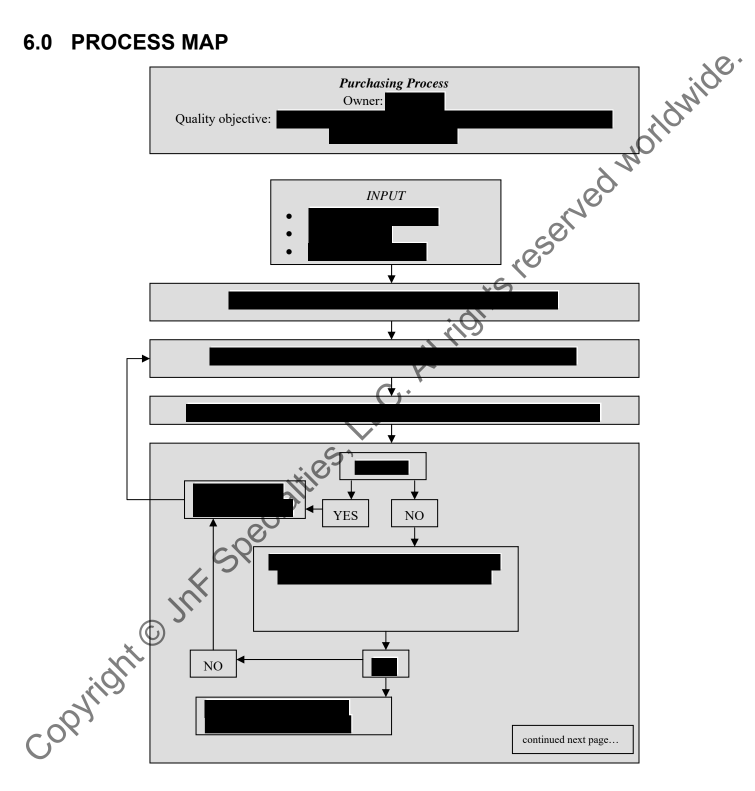
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6.0 PROCESS MAP

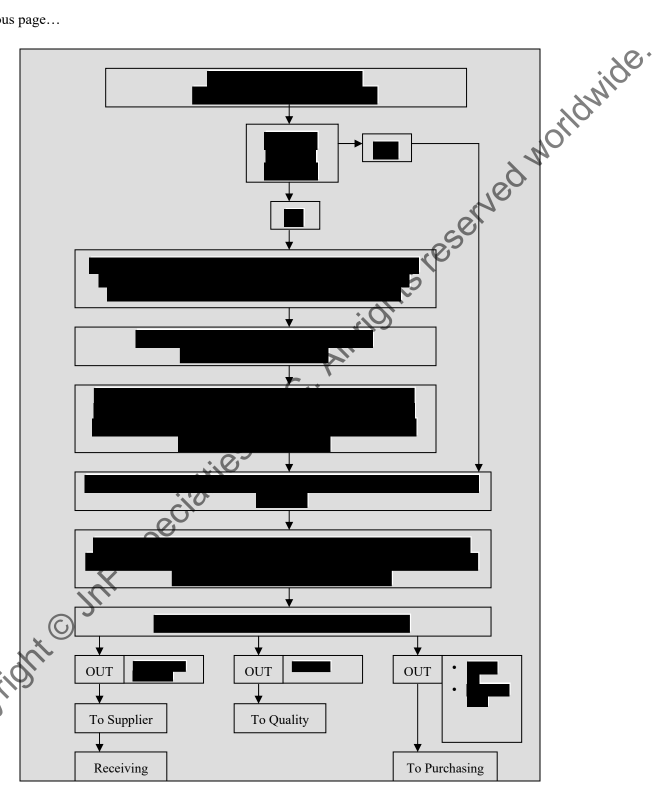


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Abstract:
This document describes the receiving and inspection process.

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1.0 PURPOSE

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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 Company process or item 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 item 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 item 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

All deliveries other than mail or express carrier are routed to

- The Responsible Authority (RA) shall
 - If the RA notices any obvious damage to the item's packaging, they
 - The Responsible Authority (RA) shall
 - The RA shall

4.0 PROCEDURE: BECEIVING INSPECTION

- 4.1 The inspector will receive the items and original paperwork from the RA.
- 4.2 Inspections are performed according to **Appendix A** or as required by

5.0 MATERIAL IDENTIFICATION

5.1 Received materials for production/fabrication are identified by one or a combination of the following methods:



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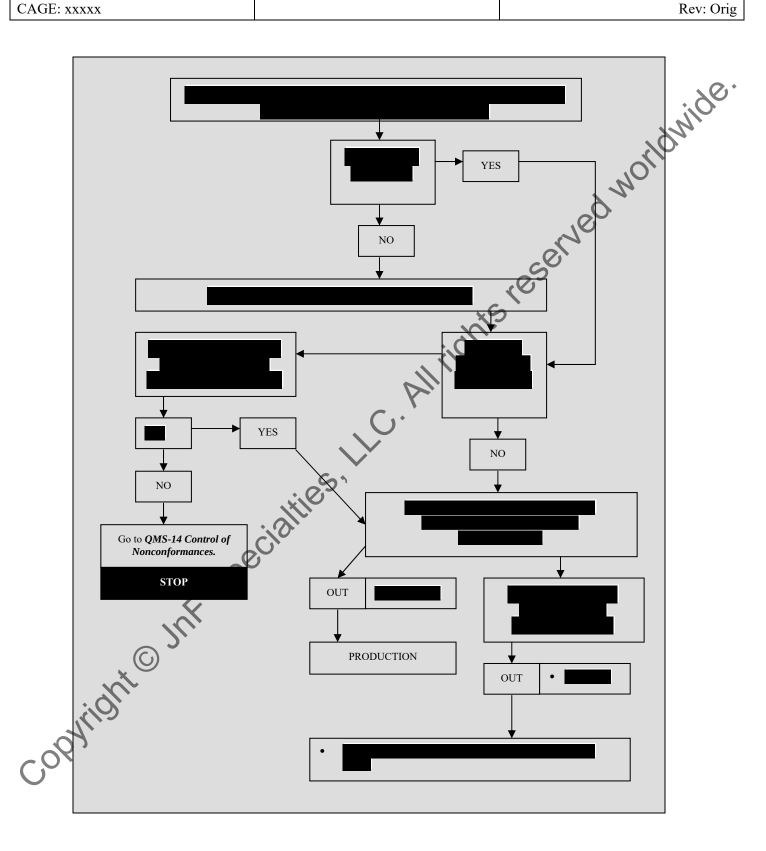
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PROCESS MAP



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APPENDIX A - Receiving Inspection Work Instructions

APPENDIX A - Receiving inspection work instructions
Op 1: Acquire copy of applicable purchase order. Perform
Op 2: Count the quantity of items received. Items exempt from counting include
Count the quantity of home received: home exempt from establing molade
Op 3: If the supply is a Catalog/Commercial item,
Op 4: Perform First Piece Mechanical/Visual inspection on a new production part number to determine
Op 5: SAMPLING PLAN: Randomly select items for
Op 6: Verify dimensional conformance of selected items according to
Op 6. Verify differsional comormance of selected items according to
Op 7: Verify conformance to the required chemical composition according to
Op 8: Verify lot/heat number traceability is
Op 6. Verify formeat number traceability is
Op 9: If the Supplier is a distributor of the supplies, verify traceability is
Op 10: If supplies are nonconforming or their conformance cannot be determined within
Op 10. Il supplies are noncomorning of their comornance cannot be determined within
Op 11: If the supply is obviously unfit for use
Op 12: Complete inspection report and
Op 13: Complete shelf life expiration log for Op 14: Record the quantity and date received on the <i>PO</i> then initial each item to indicate acceptance. Process
the <i>Purchase Order</i> according to Appendix B .
Op 15: If the Supplier's packaging is
On 16: Irraport Customer Supplied materials upon receipt to
Op 16: Inspect Customer Supplied materials upon receipt to
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APPENDIX B - PURCHASE ORDER PROCESSING

Step	IF	THEN	11
1	Items on PO not		
	received (back order	r)	•
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2	Items on the PO were	re	
	received in full		
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1.0 Purpose

This document defines the overall construction process and includes or makes reference to the procedures necessary for the process.

2.0 Theory

С	Construction operations or tasks must be conducted under controlled conditions to achieve	ve th	e highest	quality
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Problem Resolution

All employees are instructed to immediately notify a Responsible Authority (RA) whenever a process or construction related problem occurs that cannot be corrected according to

It is understood that the Responsible Authority (RA) occasionally may not be available for support; in that event, No disciplinary action may be attached to an

employee's attempt to resolve a problem.

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Construction Documentation

All revision controlled construction documents are available at the point of use and

In addition to this process procedure, additional construction documentation may be required for a construction operation. When required,

Documentation includes

Records that are created for temporary retention of miscellaneous information are not

Material Identification

Construction/fabricated materials are to be identified by one or a combination of the following methods:

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When re	equired, lot traceability or individual serialization of materials (major mark numbers) is maintained or
	Con the OMC 44 Control of Nonconformance Dropolisms
Any ma	See the <i>QMS-14 Control of Nonconformances Procedure</i> terials not marked with a tag are
	Match-Marking S
	Match-Marking
	arked.
lse nai	nted marks, attached metal tags, other durable methods which do not degrade the finish of the piece.
	tress type steel die stamps to identify and match mark pieces. If steel die stamps are used, they must
As an al	ternate location for tub girder bottom flange splice plates, place the mark
Mark gir	ders and beams on
nsure	that during fabrication, the heat number is

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7.0 Material Handl	ing
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Work instructions and/or training instructs operators on the proper and safe handling of material In all cases, operators are	_
In all cases, operators are	S
	2,
8.0 Preservation	
Operators employ proper handling and packaging (protection) and cleaning of materials and c	onstituent parts
9.0 FOD – Foreign Object Damage and Detection	
Work instructions and training methods ensure that handling and preservation practices reduce of foreign objects (FOD) into the construction when applicable. In these cases, hold points are	the introduction
or loreight objects (FOD) into the construction when applicable. In these cases, hold points are	
10.0 Customer and Government Property Control	
Customer and Government property (C&G.Property) means all hardware or property owned by	or leased to the
Customer or Government or acquired by the Customer and Government under the terms of a includes:	contract, which
•	
•	
•	
All Customer and Government furnished property and/or equipment is inspected upon receipt a	according to the
QMS-09 Receiving Procedure. Any nonconformities or shortages are	according to the

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11.0 Validation of Processes

Unless otherwise specified by engineering requirements, a certificate of conformance (CofC) is used to declare results of validation and verification of activities.

Provisions for validation and verification includes:



12.0 Inspections and Tests

12.1 Scope of Examinations

At suitable intervals, the Inspector observes

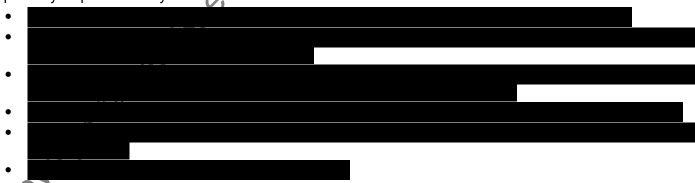
12.2 Extent of Examination

The Inspector examines the work to ensure

12.3 Preparatory Inspections

When required, preparatory inspections are conducted prior to beginning all definable segments of work as well as at the beginning of all phases of the contract. The Customer inspector and other involved personnel are notified

Preparatory inspections may include:



12.4 Initial Inspections

Initial inspections are held when

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12.5 Follow-Up Inspections

Follow up inspections are

12.6 Inspection of Work and Records

Except for final visual inspection, which is required for every weld, the Inspector

Size, Length and Location of Welds

The Inspector

12.7 In-Process Testing

In-process tests are conducted during construction to ensure ongoing quality of work. These are done randomly at the discretion of management or via planned quality control inspections according to the contract and inspection & test plan.

Testing plan procedure:

- •
- •
- •
- .

12.8 Completion Inspection

Once all operations are complete, the shop manager and quality manager

12.9 Final Inspection

When required, the quality manager, project manager or their designee and Customer representative are in attendance at this inspection. The final inspection is

(2)10Inspection and Test Status

The status of construction, inspection and testing is

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12.11 Documentation and Control

Records of inspection that provide evidence of conformance to requirements are retained and maintained according to **QMS-03 Records Control Procedure**.

13.0 Bolting

This section covers two grades of high-strength bolts, **ASTM A325** and **ASTM A490**, along with their installation and inspection in structural steel bolted joints. This section is used in conjunction with **AISC** and **RCSC**.

References:

- ANSI/AISC 360-10 Chapter "M" Fabrication and Erection
- RCSC Specification for Structural Joints Using High-Strength Bolts
- ASTM A325 Standard Specification for Structural Bolts, Steel
- ASTM A490 Standard Specification for Structural Bolts, Alloy Steel
- ASTM F436-09 Standard Specification for Hardened Steel Washers
- **ASTM F959-09** Standard Specification for Compressible-Washer-Type Direct Tension Indicators for Use with Structural Fasteners
- Research Council on Structural Connections

Drawing Information

The Engineer of Record specifies the following information in contract documents:

- The ASTM designation and type (Section 2 of RCSC) of bolt to be used;
- The joint type (Section 4 of RCSC) and method of installation
- The required class of slip resistance if slip-critical joints are specified (Section 4 of the RCSC); and,
- Whether slip is to be at the factored-load level or the service-load level, if slip-critical joints are specified (Section 5 of the RCSC).

The type of bolted connection(s) referenced on the contract documents determines the level and frequency of required testing by the **RCSC**. The types are Slip Critical, Snug Tight and Pre-Tensioned.

For all pre-tensioned types,

14.0 Protective Coatings

General

The type of coating system(s), coatings manufacturer, surface preperation and **DFT** requirements is

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Surface Preparat	ion			\Q ₁ .
The painting supe	ervisor or inspector v	verifies		107
Before any coating	g operations begin, th	e coatings supervisor ar	nd/or inspector verifies	70.
Surfaces shall				
Blast Cleaning				
Blasting abrasives	s shall		30	
5				
Final Surface Co	ndition / Profile			
The surface to be				
		\ <u>\</u>		
Application of coa	atings may be done l	by		
Areas not to be co	nated are			
A TOUS HOT TO BE O	sated are			
		. 65		
At the end of each	n coat, the applicator	inspects the work and lo	ooks for	
Curing of Protec				
The curing proces	ss and times for prot	ective caotings are		
Dry film thickness	(DFT) readings are			
. 41.				
15.0 Weldin	g			
The inspector sho	ould be an AWS certi	fied welding inspector o	r have the experience, kno	wledge and ability to
Welding procedu	re specifications (W	PS), procedure qualific	ation records (PQR) and	welder perfomance
qualifications (WP	Q) are			·
Welding filler me	tals are			
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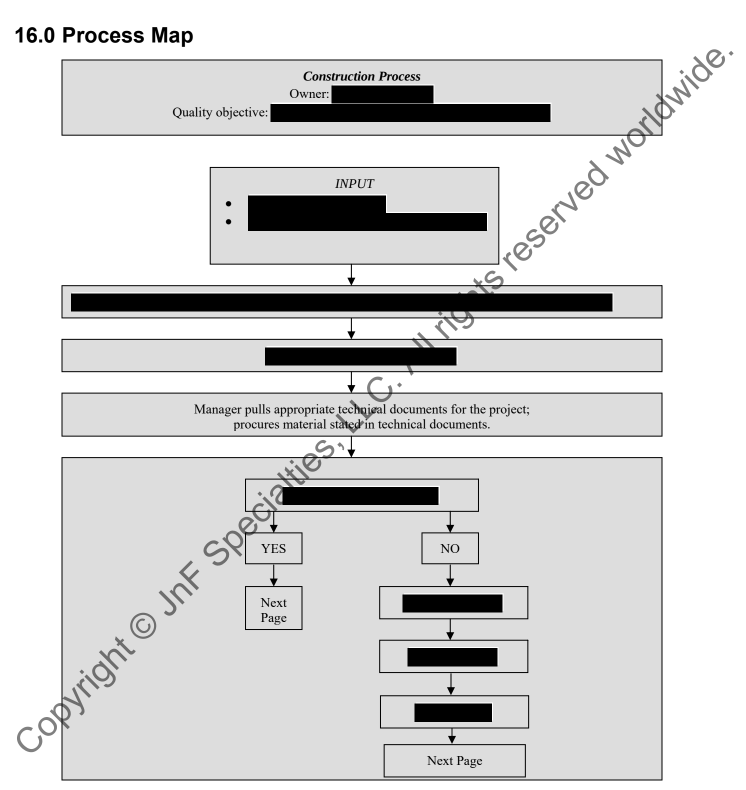
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16.0 Process Map

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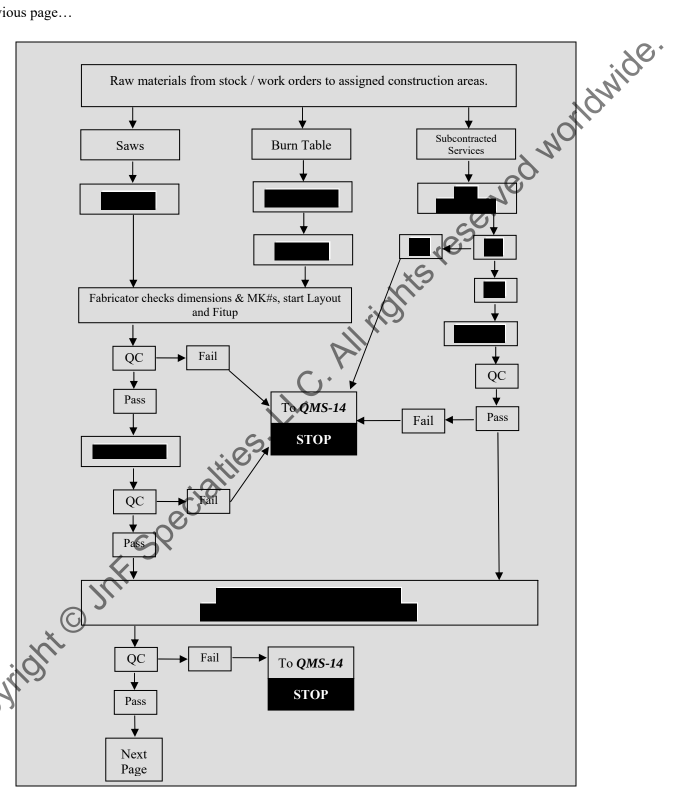




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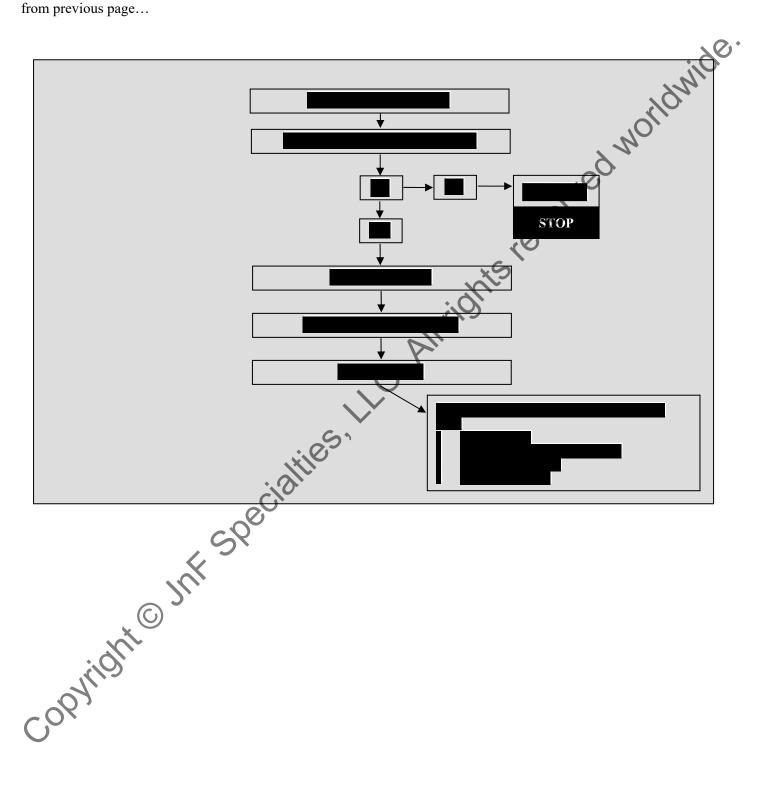
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Abstract:
This document describes the shipping process.

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1.0 **PURPOSE**

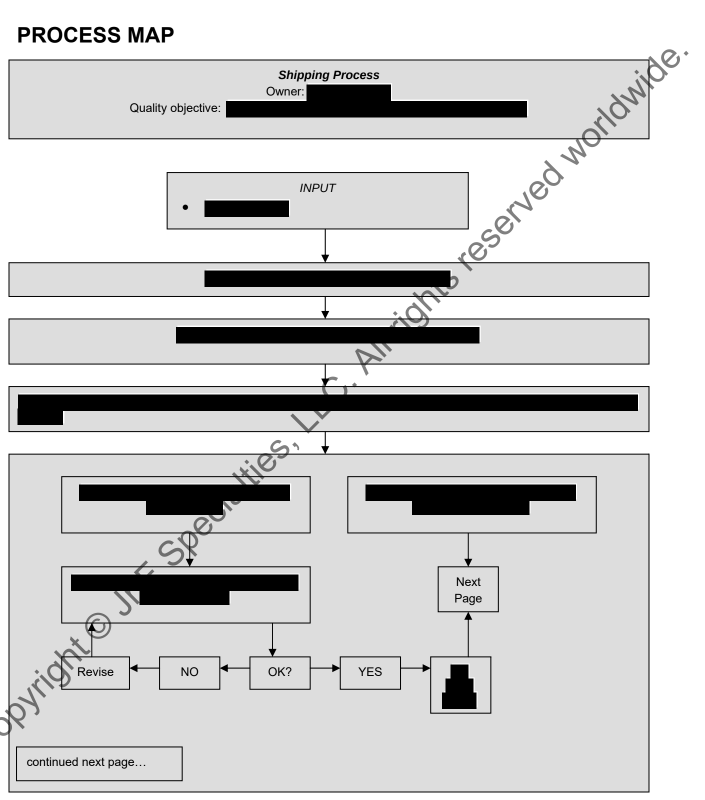
2.0 THEORY
The final packaging and arrangement of shipping is critical to the quality of fabrications as received by the Customer; as a result, Copyright. Infr Specialties, I.C. Amrights reserved in

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4.0 PROCESS MAP



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This document describes the procedure used to audit the quality management system.

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Internal Auditing

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1.0 PURPOSE

This document provides details and procedures for the internal auditing process.

NOTE: At this time, only quality system audits are conducted. When environmental system or other audits are implemented, this procedure will be amended to include rules for additional audits.

2.0 THEORY

Internal auditing of a Company's quality system is critical for maintaining good processes and documentation and for identifying areas for improvement opportunity.

3.0 INTERNAL AUDITING PROCEDURE

The Resonsible Authority takes into consideration 3.1 Internal quality audits are conducted by Audit requirements include 3.2 3.3 Auditors may Minimum auditor training requirements are as follows: 3.4 The Quality Manager plans audits according to 3.5 The Quality Manager maintains the Internal Audit Schedule that records this information. 3.6 Using the Internal Audit Report, the Lead Auditor 3.7

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3.9	The internal audit	
3.10	During the corrective action effectiveness review,	'9 ₄
3.11	The completed Internal Audit Report is	
3.12	Copies of the completed audit report are	
3.13	The results of internal audits are	
3.14	In all cases, auditees are	
	In all cases, auditees are	
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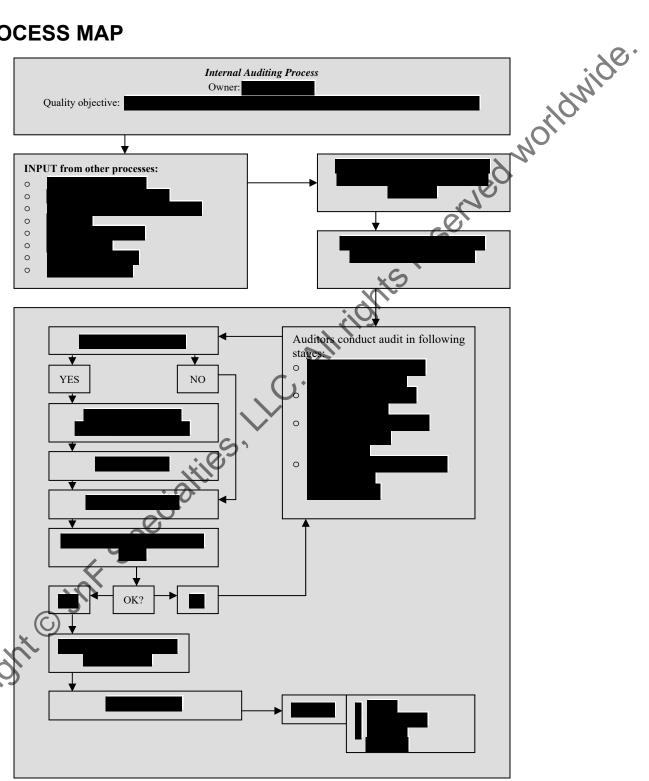


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4.0 PROCESS MAP



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This document describes the procedures used to correct nonconformities.

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Corrective Action Procedure

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Corrective Action Procedure

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PURPOSE 1.0

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

2.0 THEORY

Corrective action is taken to correct nonconformities, which could be defects found during construction, 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.

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 construction, processes and work environment.

PROCEDURE: INTERNAL REPORTS 3.0

- The Company utilizes a Request for Support (RFS) form to record nonconformances related to its construction, processes and quality system as well as compliments or positive feedback. The form and system are used for
- ALL employees are empowered with the ability to 3.2
- 3.3 No disciplinary action may be attached to the submission of RFS's.
- The Quality Manager has been assigned the role of RFS Administrator. 3.4
- 3.5 For the processing and routing of RF S's see Process Map.
- If the responsible manager determines they are not responsible for the issue involved, they 3.6
- 3.7 Actions taken shall
- The Quality Manager shall 3.8
- 3.9 In addition to corrective action efforts, management shall

hich

shall be used to address potential nonconformances. These shall be reported to management for review.

The management review process shall



Corrective Action Procedure

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3.11 Where construction is suspected of a nonconformance, the Company shall

4.0 PROCEDURE: INVESTIGATION & CORRECTIVE ACTION REQUESTS (ICAR's)

- 4.1 Any purchasing agent
- 4.2 ICAR's are processed through the same steps as the RFS but are routed to
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 Copyright on the Specialties, I.C. All rights Failure of a Supplier to respond to an ICAR or to respond with an insufficient action plan may mean

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5.0 PROCESS MAP



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This document describes procedures for control of nonconformances.

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Control of Nonconformances

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1.0 PURPOSE

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

2.0 THEORY

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

3.0 GENERAL PROCEDURE

- 3.1 "Nonconformance" is any item made by the Company or raw material used by the Company or returned from the Customer that does not meet:
- 3.2 Nonconforming items must
- 3.3 All employees are empowered to
- 3.4 Upon discovery of a nonconforming item, an employee may
- 3.5 When an employee cannot bring the item into conformance through immediate rework, the employee shall
- 3.6
- The employee shall complete the top portion of the RFS form, filling in all pertinent spaces. The employee shall then

Your Company Name

Control of Nonconformances

Rev: xx

3.8	The employee shall
3.9	Upon receipt of the RFS, the Quality representative will
0.40	
3.10	Quality will then
3.11	If the nonconforming item is ascertained or estimated to be the fault of a Supplier, Quality may
3.12	Quality will also
3.13 MRB a	The RFS shall then be submitted to the Material Review Board (MRB) for review and disposition actions that affect configuration may be immediately implemented when
3.14	The MRB consists of the following managers, at a minimum: er
	MRB Qualification erial Review Board member must:
3.15	In the event of a non-unanimous
3.16 or safe	The Company shall provide timely reporting of delivered nonconforming items that may affect reliability ety. Notification shall include

4.0 DISPOSITIONS

4.1 Dispositions are classified as Major, Minor or None.

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4.1.1	Major:		
4.1.2	Minor:		N'
4.1.3	None:		
4.2 4.2.1	MRB dispositions may include Clarification	, but are not limited to:	3 MO
4.2.2	Conditional Acceptance		
4.2.3	Non-Deliverable	, C·	
4.2.4	Notification		
4.2.5	Precautionary		
4.2.6	Repair (Non-Standard and Sta	ndard)	
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4.2.7	Request for Waiver/Deviation		.9e.
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4.2.8	Return to Supplier (Receiving I	nspection)	
420	Dowerk (Non Standard and St	andord)	
4.2.9	Rework (Non-Standard and Sta	andard)	
4.2.10	Scrap		
F 0	CHETOMED DISPOS		
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5.0 5.1	Major: A Waiver/Deviation disp	position is	
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This document describes calibration procedures.

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Calibration Procedure

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1.0 **PURPOSE**

This document defines the procedures necessary for calibration of measuring equipment.

2.0 **THEORY**

Measurement results are only valid when M&TE of known accuracy is used. This calibration procedure ensures M&TE is properly verified for accuracy against known standards. Measurement devices that are used to indicate process feedback are not subject to calibration, such as short-circuit or open-circuit, hot or cold, off or on, etc; however, when a measurement device is used to determine conformance to a Customer requirement, then the device should be properly verified for accuracy.

3.0	DEFINITIONS	eserve
•	Accuracy Ratio –	
•	Adequacy -	
•	Calibration:	
•	Gages –	
•	Inspection Aid -	
	M&TE - Measurement and Test Equipmen	
•	Procurement of M&TE -	*
•	Procurement of Mare -	
•	Recall -	
•	Significantly out-of-tolerance -	
•	Special Equipment	
•	Standards	

GENERAL CALIBRATION PROCEDURE

4.1 Calibration is performed by			
4.2 Measuring instruments are to be calibrated at a temperature of	and		relative
humidity. Sufficient temperature stabilization time is allowed before calibration. For	cases	where	calibration
must be conducted in the construction area,			

Your Company Name

Calibration Procedure

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A number is issued when a gage does not provide its own serial number. 4.3 4.4 All M&TE are kept clean and when not in use are 4.5 A recall log is maintained on all M&TE and standards. The log provides The number of items scheduled for monthly recertification is 4.6 4.7 In addition to the recall log, a Calibration Report is kept on each Company-owned gage/standard. The purpose of this report is to 4.8 Calibration intervals may be established based on one or more of the following criteria: Adjustable M&TE is periodically recalibrated based upon 4.9 TABLE I, Calibration Intervals alibration Cycles to

Interval Adjustment: M&TE whose calibration error is recorded as being greater than the last recorded 4.10 calibration error but not significantly out of tolerance

New Calibration Cycle

4_11 M&TE calibration intervals may be extended or adjusted

Qualify for New

Calibration Cycle

Calibration Cycle

Annual **∢** Bi-Annual 3 - 4 Years 5 Years

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4.12 Overdue items are

4.13 A calibration sticker is used to identify individual or groups of items of M&TE. The sticker displays

4.14 Calibration Standards/Special Equipment

The following is the position of the National Conference of Standards Laboratories (NCSL):

Calibration of standards/special equipment is conducted by checking against laboratory standards available at outside laboratories. Approved calibration laboratories are listed in the *Approved Supplier's List*.

When calibrations are made for standards/special equipment, the calibration lab is required to submit a report that contains, as appropriate:

- 4.15 A calibration record and recall log is maintained on all Transfer Standards, indicating
- 4.16 The calibration department places all Customer furnished inspection gages in the calibration system unless
- 4.17 Traceability: Inspection work instructions specify measurement and test equipment utilized for construction conformance inspection.

When specified,

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	Non-Calibrated M&TE: Upon alibrated measurement devices ng conditions:		non-calibra	ited M&TE	may be	submitte	ed for	calibration. under the
1) 2)								ajide
A non-	-calibrated measurement device	that is ver	rified accura	te		λ.	1-0[
4.19	Calibration Not Required M&TI	≣				180		
4.19.1 calibra	tion; however,						are e	xempt from
4.19.2 howev					a a	ire exemp	ot from	calibration;
4.20 and ar	Per e placed on a calibration schedu		ng or gages	owned by	employee	s are cali	brated	prior to use
4.21	Storage and Handling of Mo	&TE:						
4.22	M&TE requiring transportation	to a calibra	ation laborat	ory is				
4.23	M&TE storage areas are) 1					
4.24 storag	Archive / Long-Term Storage e if it was not:	M&TE doe	es not requir	e accuracy v	verificatio	n prior to	archive	/ long-term
•								
M&TE	that has been calibrated a	and stored						
5.0	OUT-OF-TOLERANG	CE EQU	JIPMENT	AND T	OOLIN	IG		
5.1 exhibit	Calibrated M&TE that is foun ing some other form of anomald			ut of tolerar	nce, dam	aged, ino	perative	e, erratic or
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5.2 M&TE found significantly out of tolerance at recalibration for 2 interval cycles is

5.3 An instrument whose calibration error is significantly out-of-tolerance over a short portion of a specified range may

6.0 LOST EQUIPMENT

6.1 Measurement and test equipment that cannot be located is classified as "Lost".

APPENDIX 1

Setting and/or selecting a reference standard to calibrate a measurement device.

Requirement:

The measurement range of a device being checked for accuracy must

VOLTMETER:

A voltmeter shall be verified for accuracy within an equivalent range on the reference standard:

A voltmeter reference standard may have scales that range from 2-20V, 20-200V, etc.

The voltmeter being checked for accuracy must be set to bracket within a range of the reference standard - or -

OTHER MEASUREMENT DEVICES:

Any reference standard whose maximum measurement range is the same as the device being checked for accuracy must

For instance,

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Calibration Procedure

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APPENDIX 2

Nonadjustable M&TE is inherently stable and includes
The Operator is only required to shock inherently stable MSTE for damage prior to each use because
The Operator is only required to check inherently stable M&TE for damage prior to each use becaus
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For instance,
To control the inventory of inherently stable M&TE, the Responsible Authority
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1.0 **PURPOSE**

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and"

anational Organization for Standardization

anaties Measurement and Test Equipment
MRB: Material Review Board
MTR: Mill Test Report as defined in Section 14 of ASTM A6
NCP: Nonconforming Product
NCR: Nonconformance Report
IDT (NDE): Nondestructive Testing (Nondestructive Testing (Nondest This document provides the accepted definitions and abbreviations for terms used by the Company.

2.0

- CCB: Configuration Control Board

- R&D: Research and Development
- RA: Responsible Authority
- REA: Responsible Engineering Authority
- RFCA: Request for Corrective Action
- RFI *: A written request for information or clarification generated during the construction phase of the project
- RFP: Request for Price/Proposal
- RFS: Request for Support
- RQA: Responsible Quality Authority
- RCSC: Research Council on Structural Connections
- RTV: Return to Vendor
- SAE: Society of Automotive Engineers
 - SB (also S/B): "should be"
- S.E.: Structural Engineer
- SSPC *: The Society for Protective Coatings, which was formerly known as the Steel Structures Painting Council
- WPS: Welding Procedure Specification as defined by ANSI / AWS A3.0

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3.0 DEFINITIONS (GLOSSARY)

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Checking (of Shop Drawings, digital F	Production Model and Erection Drawings)	No
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Specifications	Office
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Abstract:

Abstract:
This document describes the procedures used to design and develop construction and services.

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	DEVELOPMENT PROCEDURE velopment planning

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1.0 PURPOSE

This document provides details on the Design and Development process.

2.0 THEORY

The Company performs research and development (R&D). Controlling the design and development activity ensures that construction designs meet all requirements and that parts produced are adequate as a result of the design.

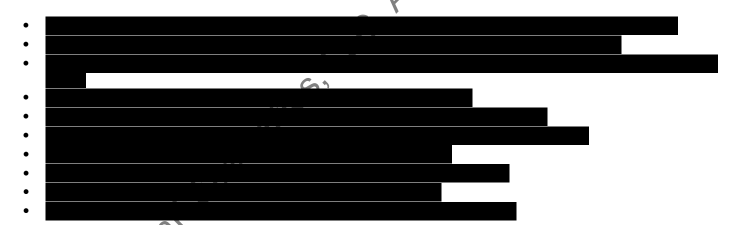
3.0 DESIGN & DEVELOPMENT PROCEDURE

3.1 General

The responsible engineering authority (REA) for design and development is assigned by the Operations Manager. Design and development personnel from various business groups may include

3.2 Design and development planning

The Company considers the following conditions when determining the stages and controls for design and development:



3.3 Design and development inputs

The Company considers the following conditions when it determines requirements essential for the specific types of construction and services to be designed and developed:



The Company determines that design and development inputs are

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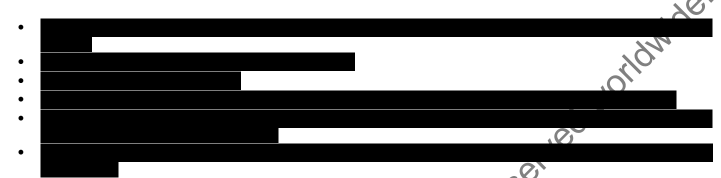
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3.4 Design and development controls

The Company applies controls to the design and development process to ensure that:



3.5 Design and development outputs

The Company ensures that design and development outputs:

The Company retains records for design and development outputs.

3.6 Design and development changes

The Company identifies, reviews and controls changes made during or subsequent to the design and development of construction and services to the extent necessary to ensure

The Company retains records for

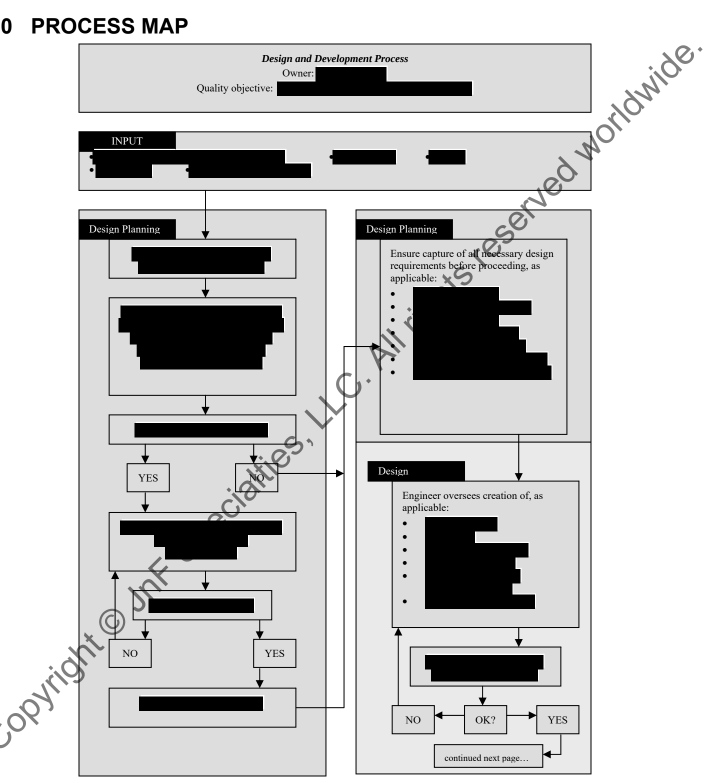


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4.0 PROCESS MAP



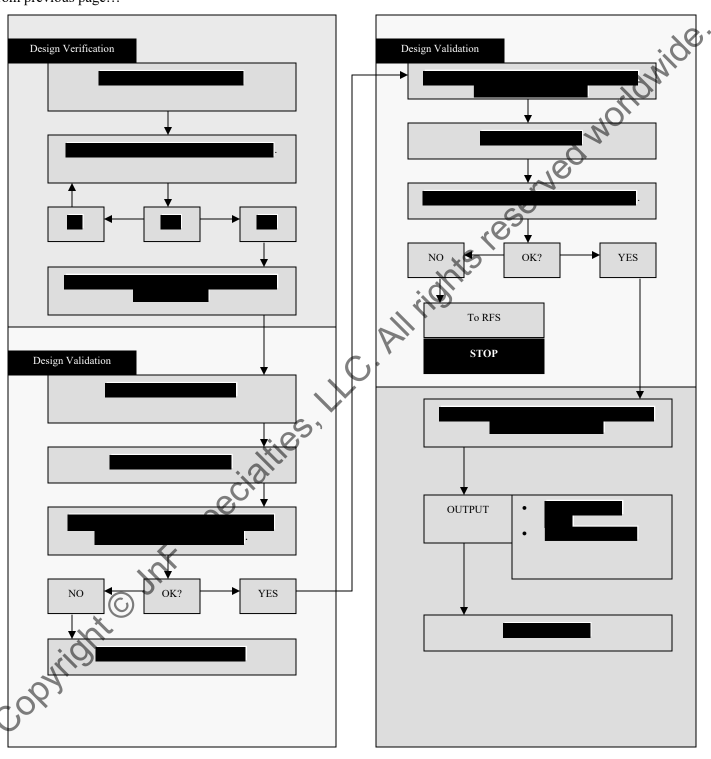


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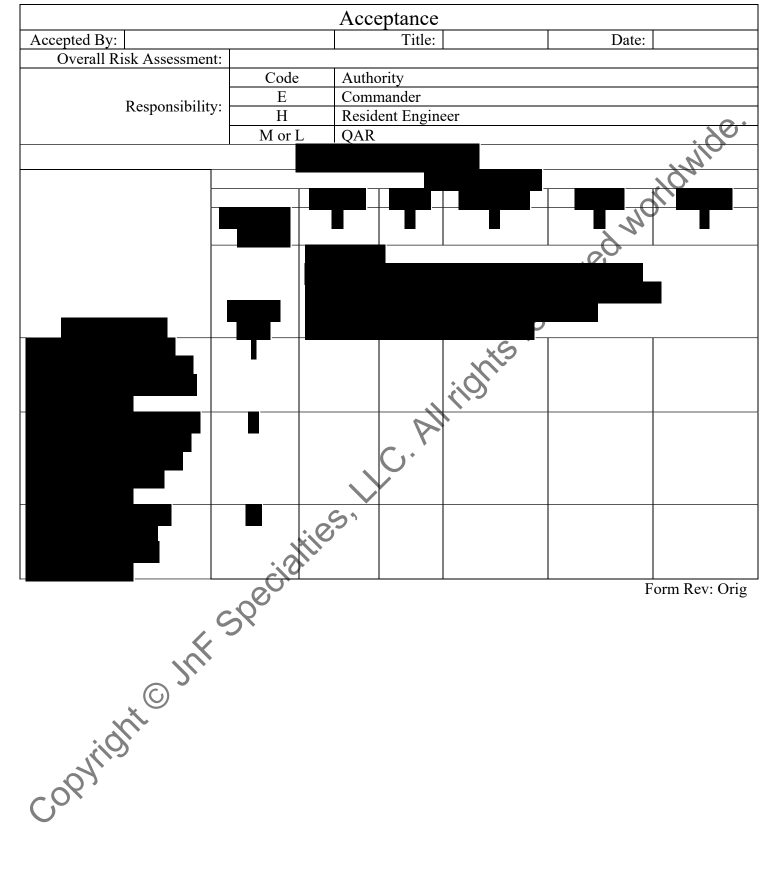
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such permission. Procedure:
Supplier evaluation:
The Quality or Purchasing Group forwards Supplier Survey for completion by Supplier.
Supplier evaluation is required for deliverable materials.
Supplier evaluation is <i>not</i> required for *.
A new Supplier is submitted to management for review. Management has discretionary
authority to
Supplier capability/approval is determined by:
Supplier capability/approval is determined by:
Acceptable Practice:
Suppliers are
Non-deliverable material Suppliers are
Suppliers that provide process materials that affect production of deliverable items are
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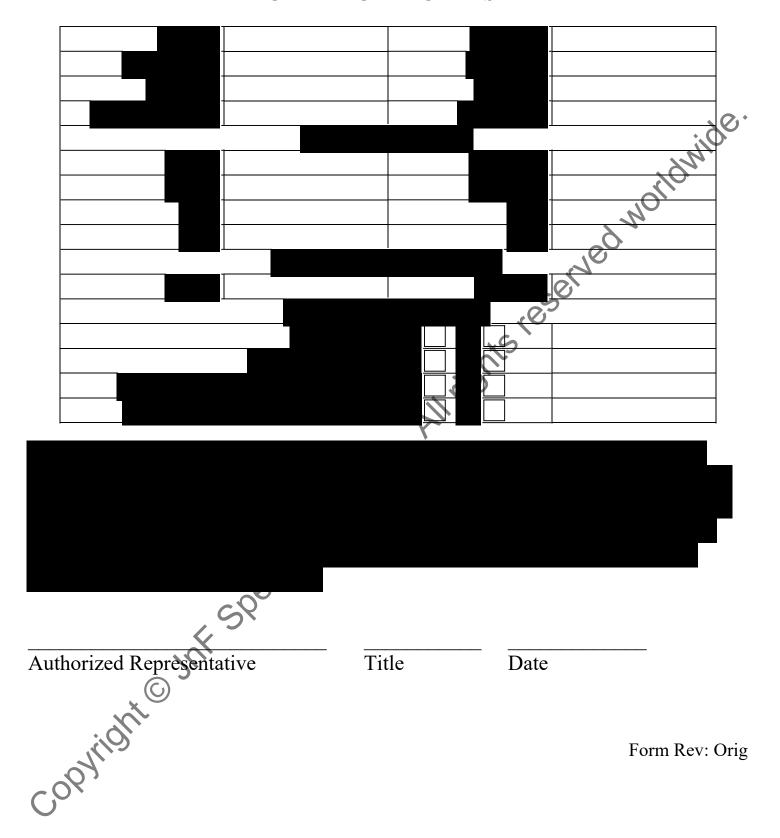
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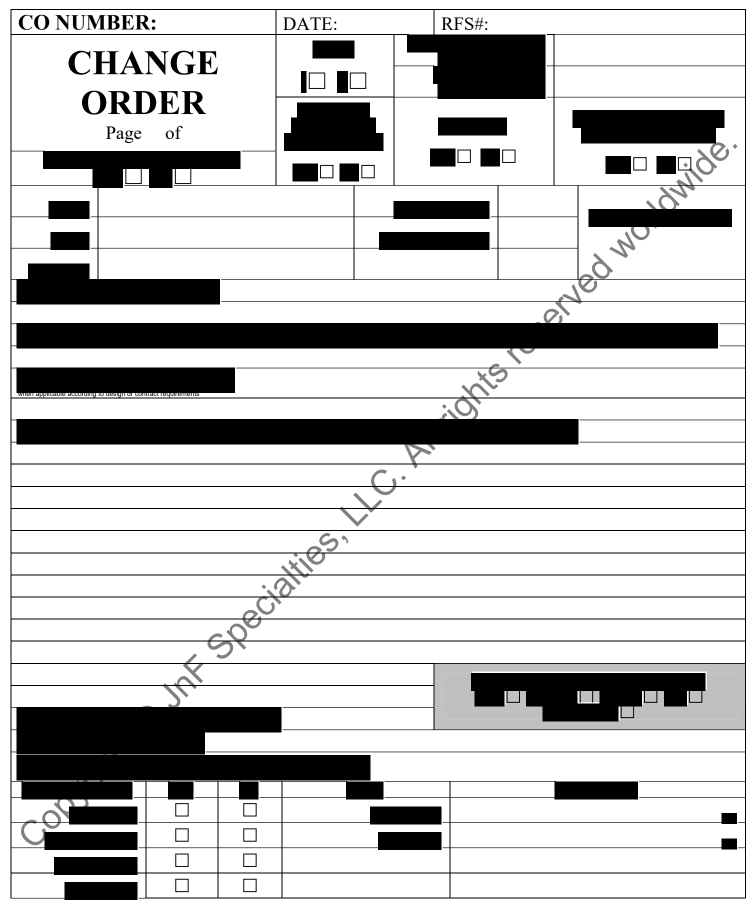
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Jnk specialis Abstract:
This specification establishes a standard of quality for the construction of Customer facilities.

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PART 1.0 GENERAL

1.1 RELATED DOCUMENTS

Procurement documents and general provisions of the Contract

See Environmental Protection Standard ined wo

See Temporary Facilities and Control Standards

1.3 LEADERSHIP IN ENERGY AND ENVIRONMENTAL DESIGNALEED

See Sustainable Design Standards.

1.4 CODES AND ORDINANCES

The Company shall comply with all currently adopted codes, ordinances, laws and regulations applicable to the work. The Company shall be fully responsible for

1.5 REFERENCES

General:

The Company shall comply with the applicable provisions of the referenced standards except as modified by governing codes and the Contract Documents.

SAFETY AND HEALTH STANDARDS 1.6

The Company shall comply with the Federal safety orders as set forth in OSHA and comply with

1.7 FIRE SAFETY

Existing building sites:

The Company shall do all things reasonably necessary to preserve the capability of protecting interior building areas from perils of fire.

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New building sites:

The Company shall do all things reasonably necessary to ensure delivery of an adequate and reliable water supply to the construction site for fire protection. Prior to the time exterior walls and roofs are erected,

Notify the Project Manager 48 hours in advance of any connections.

The Company shall do the following and all other things reasonably necessary to protect the work from the hazards of fire and wind.



Notification

The Company shall provide the Customer prior notification of

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1.8 **FURNISHED PRODUCTS**

The Company will furnish products indicated - the Work includes providing support systems to receive Jordwide. equipment.

1.9 SYSTEM DESCRIPTION

See applicable contract design requirements and performance requirements

1.10 **SUBMITTALS**

Product Data: Submit for action.

The Company shall

Shop Drawings: Submit for action.

The Company shall

Samples: Submit for action.

The Company shall

Calculations: Submit for information.

Quality Assurance/Quality Control Submittals: Submit for information, as required:

Document Review:

The Company shall

Closeout Submittals: Submit for Owner's documentation, as required:

OUALITY ASSURANCE

Definition of Qualified Installer:

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The Company shall

Pre-Installation Meetings:

Before the start of Work, Company representatives shall

1.12 DELIVERY, STORAGE AND HANDLING

Packaging, Shipping, Handling and Unloading: As required

Storage and Protection: As required

1.13 WARRANTY

Submit in Customer's documentation.

1.14 MAINTENANCE

As required to include required Extra Materials.

1.15 EMERGENCY POINT OF CONTACT

The Company shall

PART 2.0 PRODUCTS

2.1 MANUFACTURERS

All articles, material, and equipment shall

2.2 MATERIALS

The Company shall

2.3 SURPLUS MATERIALS AND EQUIPMENT

Existing materials and equipment that have been removed and are in reusable condition shall

Salvageable materials include, but are not limited to,

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Company owned construction materials and/or equipment or construction declared surplus but not in reusable condition, as determined by the Project Manager, shall

The Company is responsible for loading and unloading all materials. Only Company personnel shall

Refer to Environmental Protection Standards for any construction materials or equipment suspected to contain hazardous materials or residue for disposition requirements.

2.4 FIXED ASSET EQUIPMENT

Fixed asset items are identified by Company tags or labels.

Fixed asset equipment include, but are not limited to,

Fixed asset equipment shall

Refer to Environmental Protection Standards for any construction materials or equipment suspected to contain hazardous materials or residue for disposition requirements.

2.5 FINISHES

Whether or not specifically required by other construction documents, paint all new construction and equipment as called out in Interior and Exterior Painting Standards.

2.6 SOURCE QUALITY CONTROL

Add requirements as necessary

2.7 MONITORING ACTIVITIES

The Company will

PART 3:x EXECUTION

3.1 GENERAL

Edit the following as required.

Manufacturer's Instructions:

Prepare substrates, apply primers and install (erect, apply) the work, including

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Coordination of Work:	
The Company shall	
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Refer to architectural drawings and industrial engineering layouts for inten-	ded layout and appearance.
Coordinate work with that of others to produce	
Maintain minimum 3 ft. 6 in. of clear space in front of	
Coordinate among trades to achieve required	
Remove and reinstall work that interferes with	
TIL C	
The Company shall	
Modification of New and Existing Construction:	
Do all cutting, fitting and patching required to adapt to site conditions a	nd as required to complete the
project, even if	
	

Lead-based paints have been used on buildings and metal structures because of their durability and corrosion resistance. Buildings and metal structures may be coated with lead-based paint. Any demolition work of painted surfaces must be performed in conformance to OSHA Lead in Construction Standard,

Page 9 of 12

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Title 8, CCR Section 1523.1

Equipment and Operations Noise:
Prior to start of any work in the affected area
Prior to start of any work in the affected area,
Gasoline or propane powered equipment shall
Dust Control: During construction, keep dust to a minimum and under control. Refer to Environmental Protection
Standards. Where walls or floors (non-hazardous materials) are removed, cut or installed, a large
commercial type vacuum cleaner shall
<u>Hile</u>
Existing Services:
Where existing utility, electrical or other services are temporarily disconnected because of demolition or other construction activity, they shall
SS
3.2 USE AND CARE OF PREMISES
General General
The Company shall
The Common and each enhantment identified with the contract that were been detailed to
The Company and each subcontractor identified with the contract that may have work to perform in any part of the premises in which mechanical apparatus and equipment of fixtures of any sort are installed, or
in the process of being installed, shall

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Welding and open flame operations:

Take special precaution at all times against fire. Advise the Project Manager 48 hours prior to start of any welding, torch cutting or open flame operations on the project.

The Company is responsible for
9
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Existing utilities:
Utility and/or service lines shall
Temporary loadings of floors and roofs:
Temporary floor or roof loadings for the storage of materials, equipment, lifting devices, and like items,
or transporting of material or equipment across a floor or roof shall
For transporting of heavy materials or equipment across the floor or roof, the Company shall
3.3 INSTALLATION Anchors and Fastenings:
3.3 INSTALLATION
Attended and Lastenings.
The Company shall
Attention is specifically directed to the following operations where violations to the above may occur.
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Drilling or other penetrations of flutes in metal decking from the underside is prohibited. Supports for
ceilings, lights, small piping, etc., from overhead metal deck must

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3.4 FIELD QUALITY CONTROL

Inspection:

The Company shall

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	Origination	n Date: XXXX	
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-	Document Identifier:	Construction Readiness Review	
]	Date:	Latest Revision Date	
]	Project:	Customer, Unique ID, Part Number	
	Document Status:	Draft, Redline, Released, Obsolete	
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This document describes the process for performing a construction readiness review.

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Your Company Name

Construction Readiness Review

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6.2	ACTION ITEMS	
	Specilalla	
	Spyright. Specialties, I.C.	

Your Logo	Your Company Name	Construction Readiness Review	
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1.0 **PURPOSE**

The purpose of the Construction Readiness Review (CRR) is to demonstrate overall construction readiness and assure compliance with the requirements of the contract. All necessary construction plans, travelers, tools, facilities and other resources shall be in place and available to ensure conformance to all quality and design requirements within the negotiated program budget and schedule.

2.0 **SCOPE**

- This procedure shall apply to all construction and outside subcontractors/suppliers. 2.1 Construction Readiness Reviews should be identified during the proposal phase of a program and shall be specified in the negotiated contract.
- 2.2 This document addresses issues related only to 'readiness to start construction'. In instances where a Supplier is responsible for design and analysis tasks, additional design reviews shall be required. Design/analysis reviews and how to conduct them are not in the scope of this document. However, any residual issues from design reviews that are related to construction shall be considered suitable for inclusion in the CRR agenda.

APPLICABLE DOCUMENTS 3.0

This document is subject to the requirements of the following subcontract documents in descending order of precedence.



GENERAL 4.0

A construction readiness review is required when any of the following conditions exist. 4.1



Your Logo
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Your Company Name

Construction Readiness Review

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4.1.5

CRR PROCESS, REQUIREMENTS AND RESPONSIBILITIES GENERAL **5.0**

5.1

5.1.1 A CRR is a formalized process of review and critique conducted jointly by the Customer and the Company to assess the overall construction readiness of structures or other equipment according to the subcontract document prior to starting the construction operations. The objective is

5.1.2 The review shall be conducted on-site by the Company Team and Customer Team assembled per paragraph 5.2. The Owner may

5.2 CRR TEAMS

The Company Team shall consist of

5.2.1 It will be the responsibility of the Subcontract representative

5.2.2 Similar to the Company Team, the Customer Team shall be comprised of

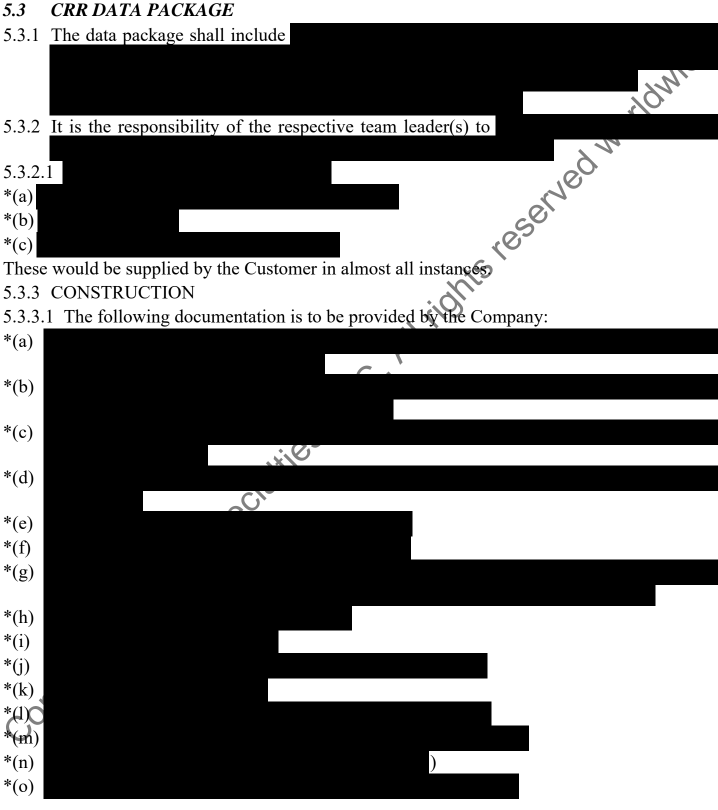
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Construction Readiness Review

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5.3.4	Materials Procurement and Subcontractors:
5.3.4.	1 The following documentation shall be provided by Subcontractors:
*(a)	:.8
*(b)	
*(c)	
	Program Management
	Program Management The following documentation shall be provided by Subcontractors: Program Management The following documentation shall be provided by the Company:
*(a) *(b)	
(c)	
(d)	
(e)	
	CRR SCHEDULE
5.3.6	
5.4	
5.4.1	The date of the CRR proceedings shall be set at the time of contract award, if possible,
	but no later than the published program schedule release (usually 30 days ARO) The CRR proceedings shall be scheduled to coincide with
	The CKK proceedings shall be senedured to comelec with
- 4 -	
5.4.2	A complete data package shall
5.5	CRR AGENDA AND PROCEDURES
5.5.1	The agenda for the CRR Proceedings shall,
5.5.2	The CRR proceedings shall be held on-site at the Company's facility and the Company's
4	Team leader, usually the Construction Manager shall act as the Proceedings Chairman.
ر,0	The agenda shall include

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Construction Readiness Review

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5.5.3	It is the responsibility of the respective team leaders to
6.0	POST-CRR EVALUATION AND ACTION ITEMS FOLLOW-UP CUSTOMER FEEDBACK AND READINESS RATING
6.1 6.1.1	CUSTOMER FEEDBACK AND READINESS RATING Following the CRR proceeding, the Customer Team shall An overall readiness rating shall be
> SA	assigned from the following three categories:
> CO	ACTION: NDITIONAL
<i>-</i> CO.	ACTION:
> UN	SATISFACTORY.
	ACTION:
6.2	ACTION ITEMS
	All action items generated through the CRR proceedings and the feedback briefings to Company management shall
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Construction Readiness Review

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Attachment 1 ACTION ITEM

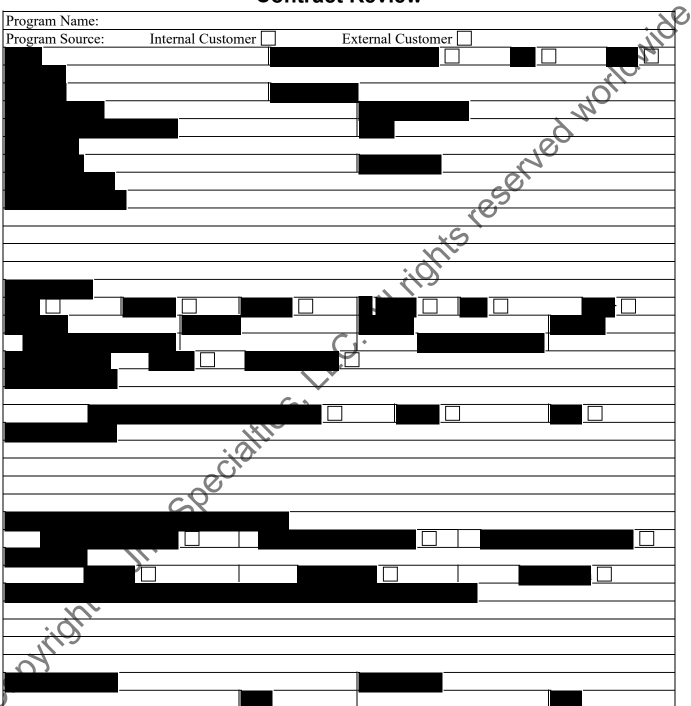
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Meeting:	Action Item Number: Due date:
	Due date: Due date: Due d



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Contract Review



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Compliance Matrix-1

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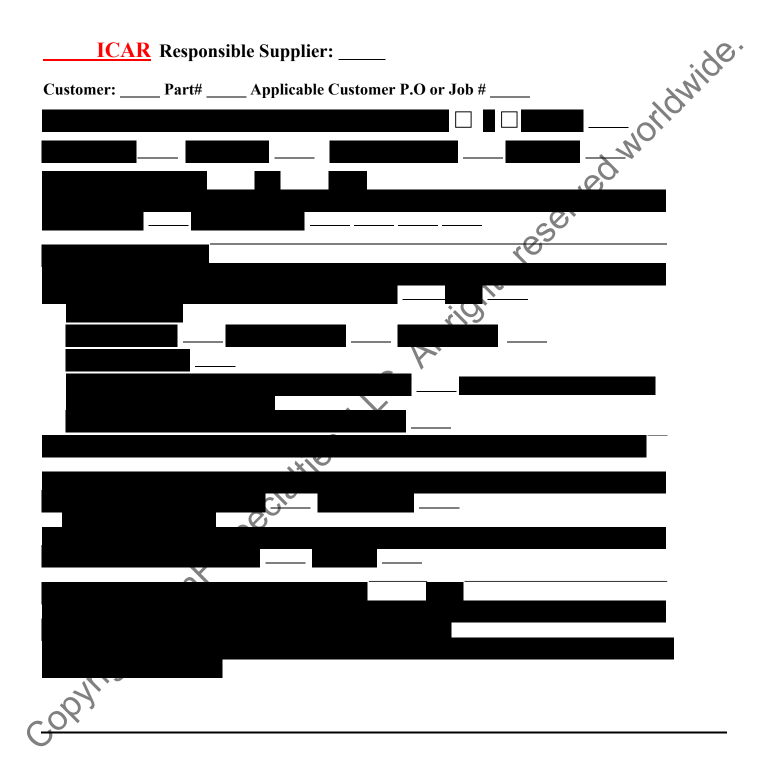
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REQUEST FOR CORRECTIVE ACTION



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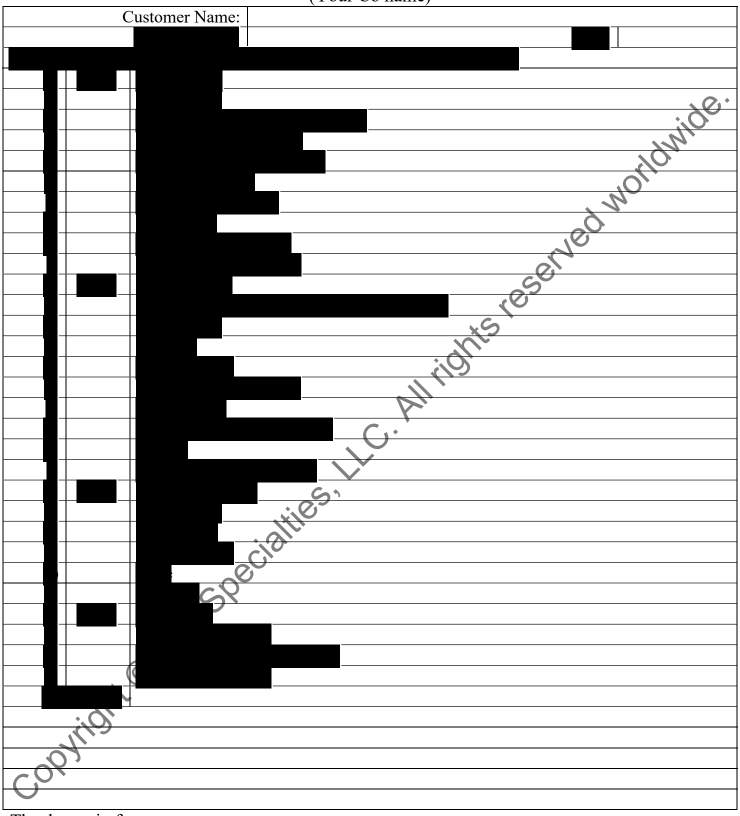
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CUSTOMER PERCEPTION SURVEY (Your Co name)



Thanks again for your support

Please Fax the completed survey to: (Your Name and Fax#)

CUSTOMER SATISFACTION SURVEY

Your Logo

Date: (input date)

To: **Customer Contact Name**

Directings,

We are asking you to spend a few minutes out of your busy day to please circle the number representing our particle.

Thank you for your participation in our survey - please fax your response to:

Your Name - Phone: Your# - Fax: Your#

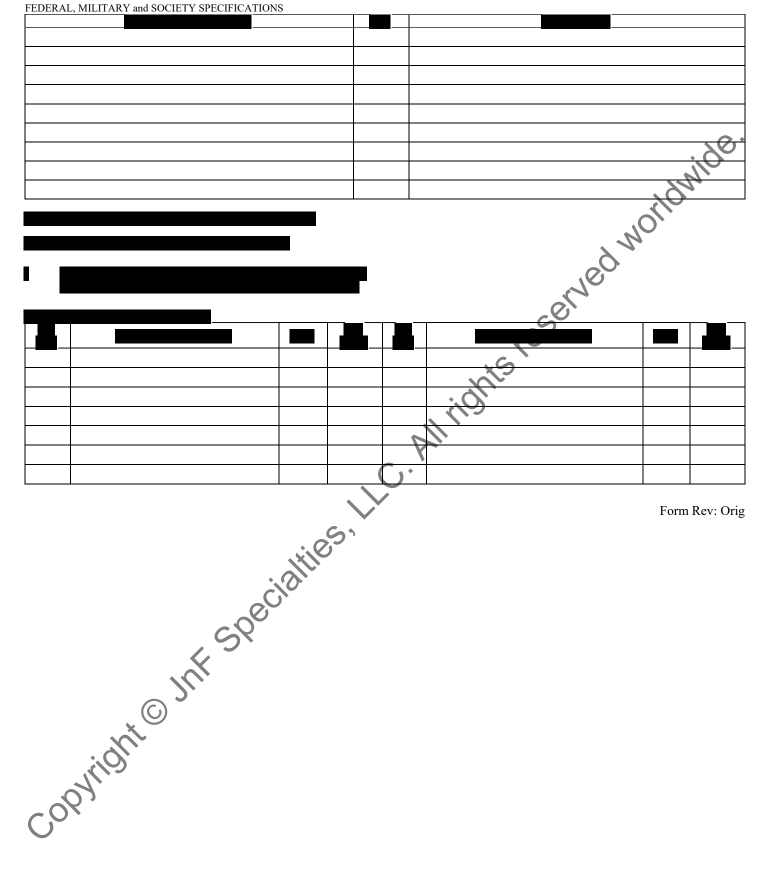
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DAILY CONSTRUCTION QUALITY CONTROL REPORT

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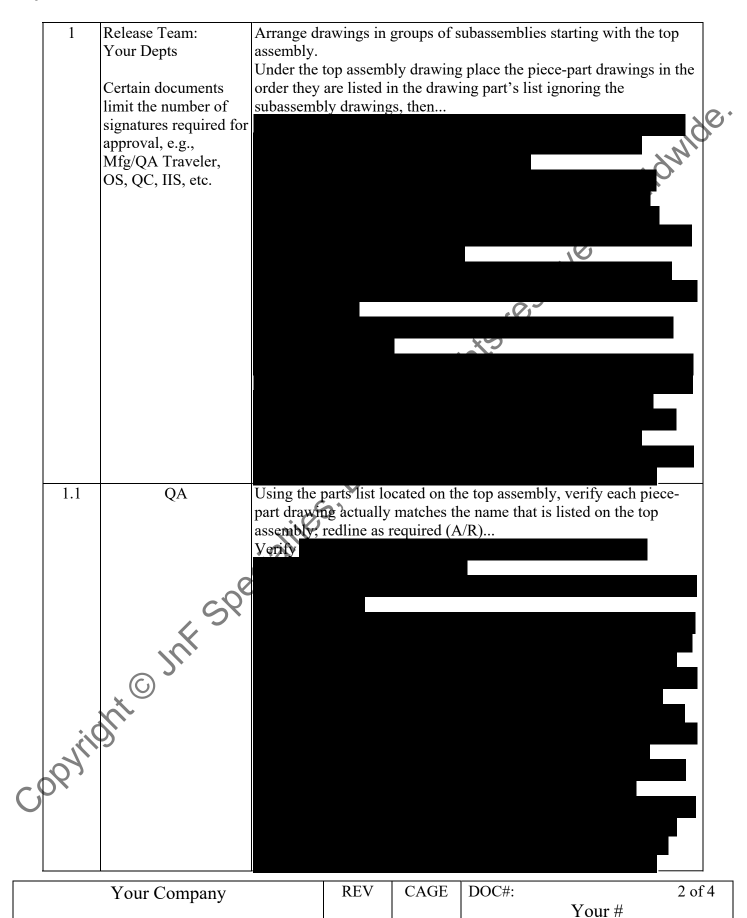
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Sample Equipment List

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FACILITY PLAN

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Project.	Customer, Unique ID, Part Number	
Document Status:	Draft, Redline, Released, Obsolete	

Abstract:

Specialité This document describes the facility plan for the (your project name).

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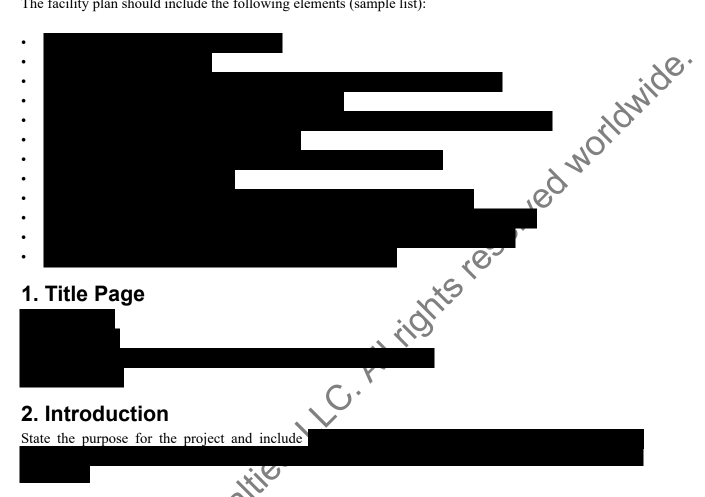
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2.	Introduction
3.	Existing Conditions and Projections
4.	Existing Facilities Evaluation
5.	Project Development
6.	Recommended Project
7.	Environmental Review
Fa	cility Plan / Map
COR	Title Page 4 Introduction 4 Existing Conditions and Projections 4 Existing Facilities Evaluation 4 Project Development 5 Environmental Review 5 cility Plan / Map 6
	Page 3 of 6

Page 3 of 6

The facility plan should include the following elements (sample list):



3. Existing Conditions and Projections

Indicate the planning area, the existing service area and potential future service areas on a map or sketch.

5. Project Development

Consideration should be given to key project conditions that must be met to complete the task. Include total project development cost analysis (sample list):



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6. Recommended Project

Provide the total project costs for the recommended project, which includes

For the recommended project, include all of the following (sample list):

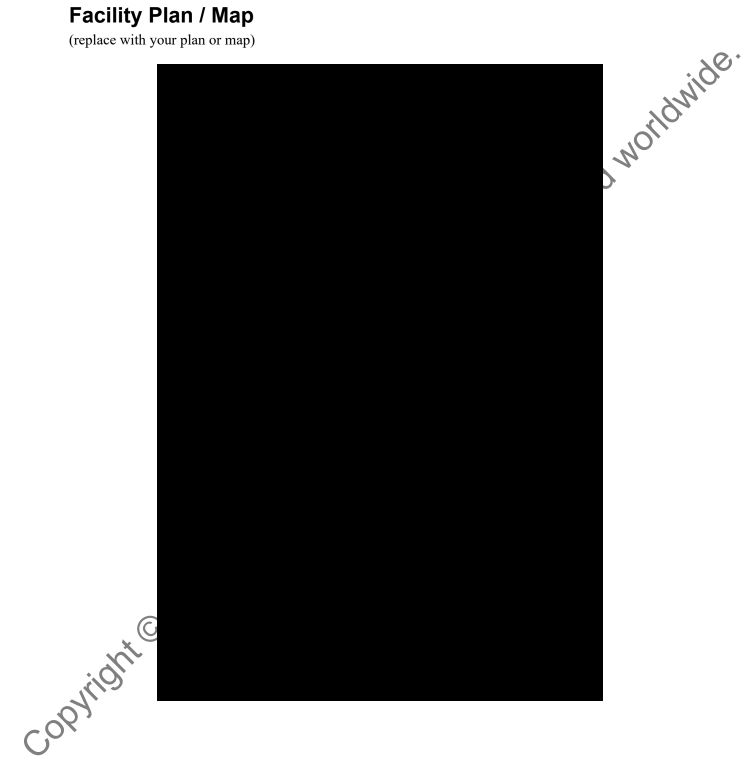
7. Environmental Review

Moildnige. Copyright July Specialties, I.C. All rights rest Provide an evaluation of the positive and negative impacts of the proposed project on the environment.

Page 5 of 6

Facility Plan / Map

(replace with your plan or map)



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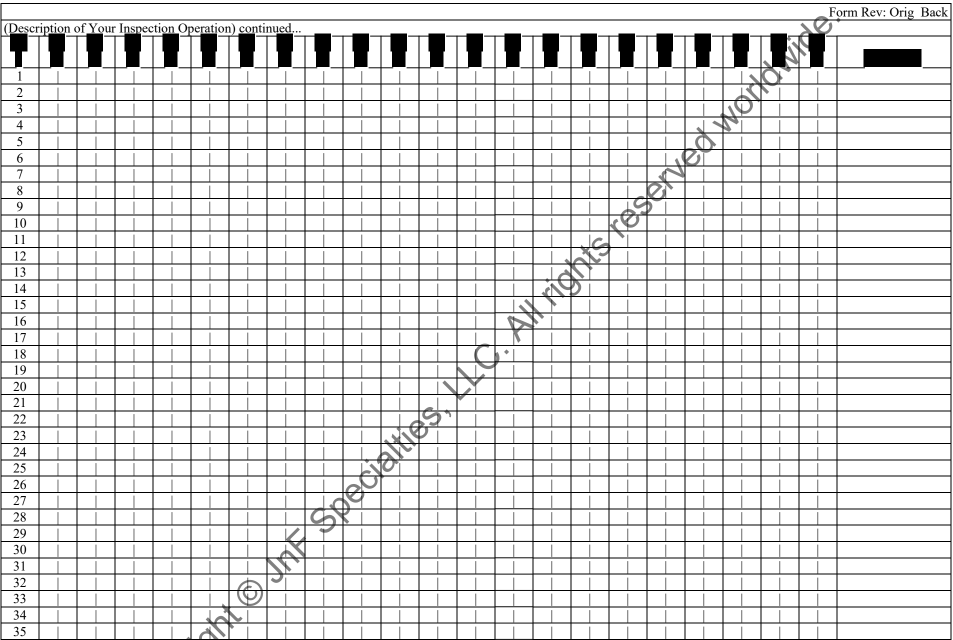


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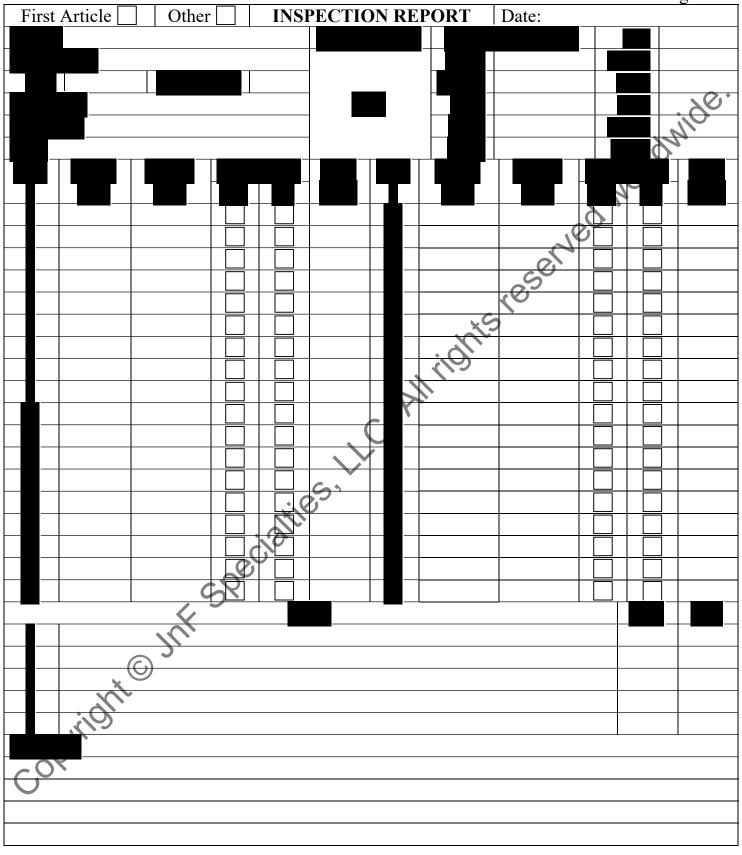
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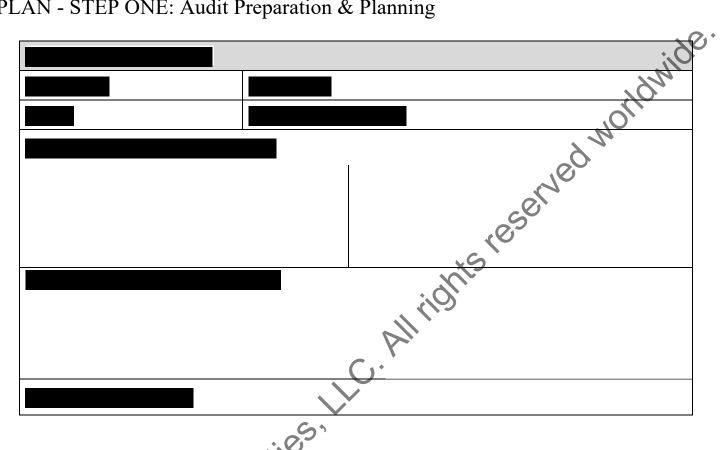
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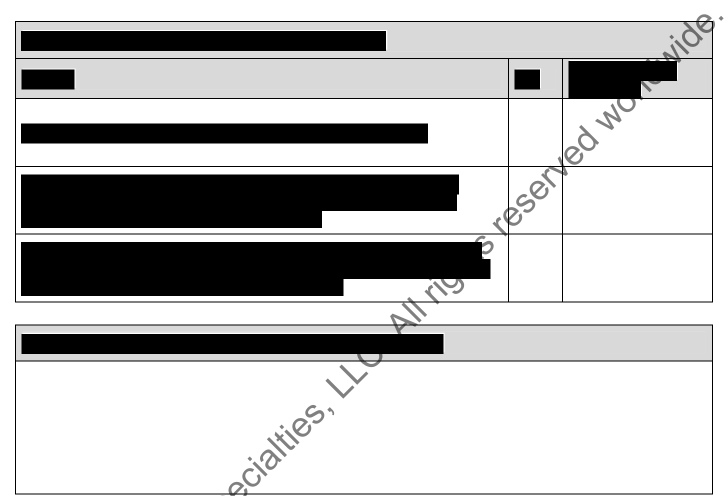
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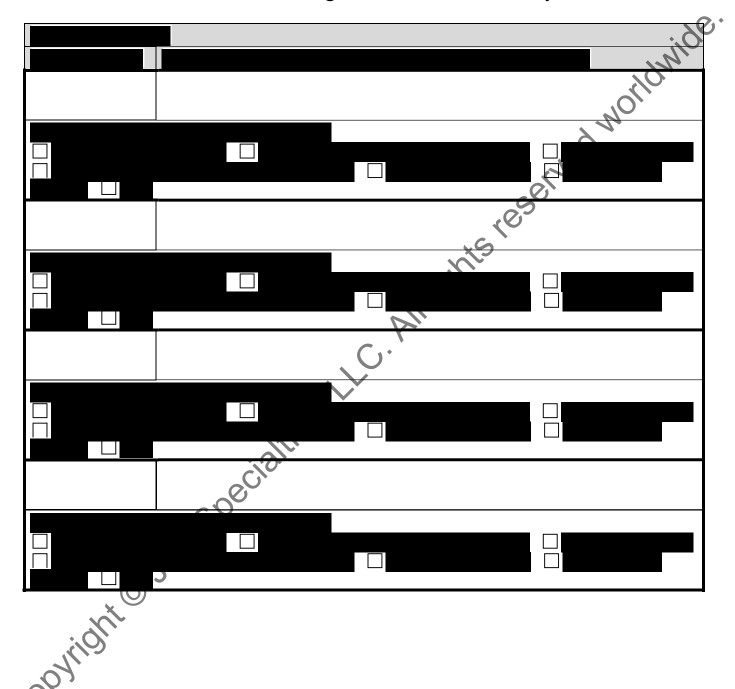
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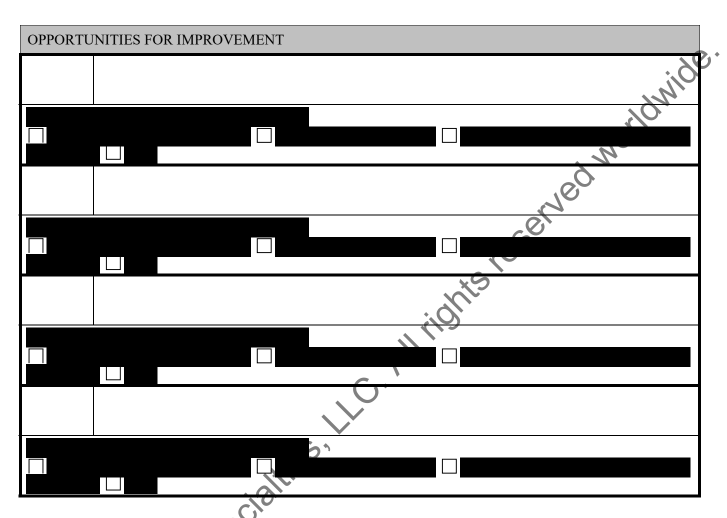
STEP FIVE: Summarize Your Findings for Nonconformance System



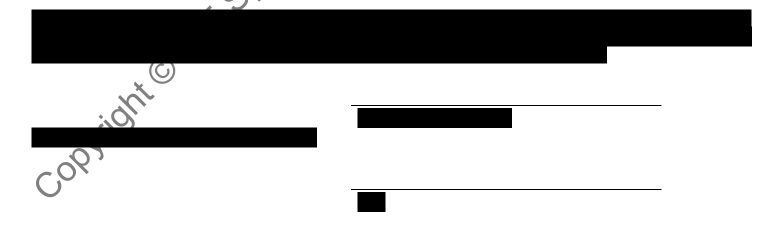
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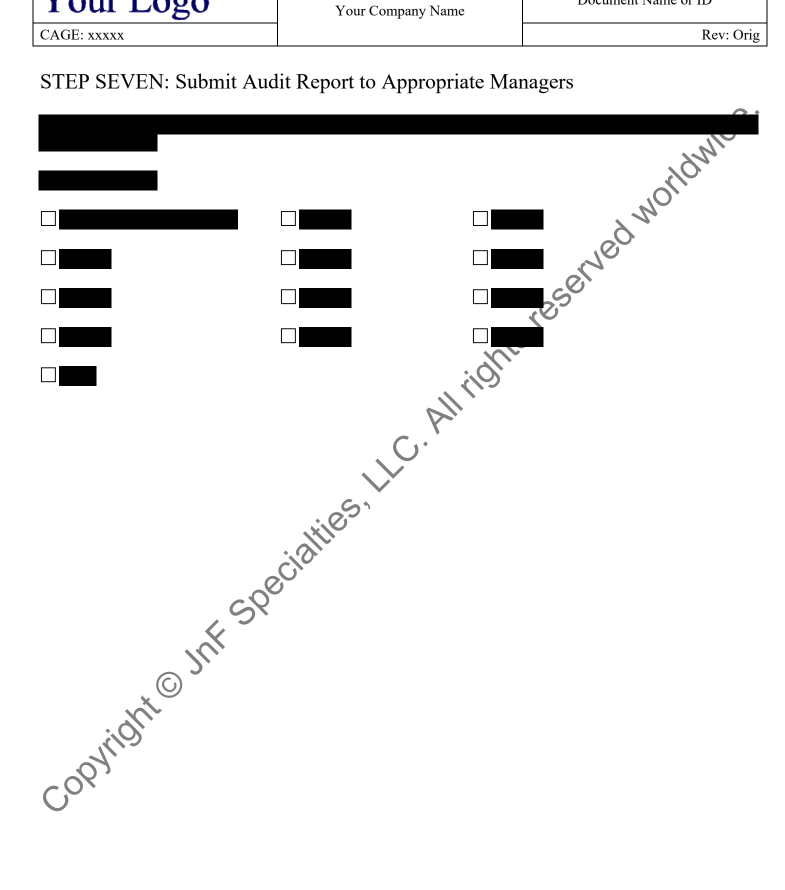
STEP SIX: Review Audit Report and Submit



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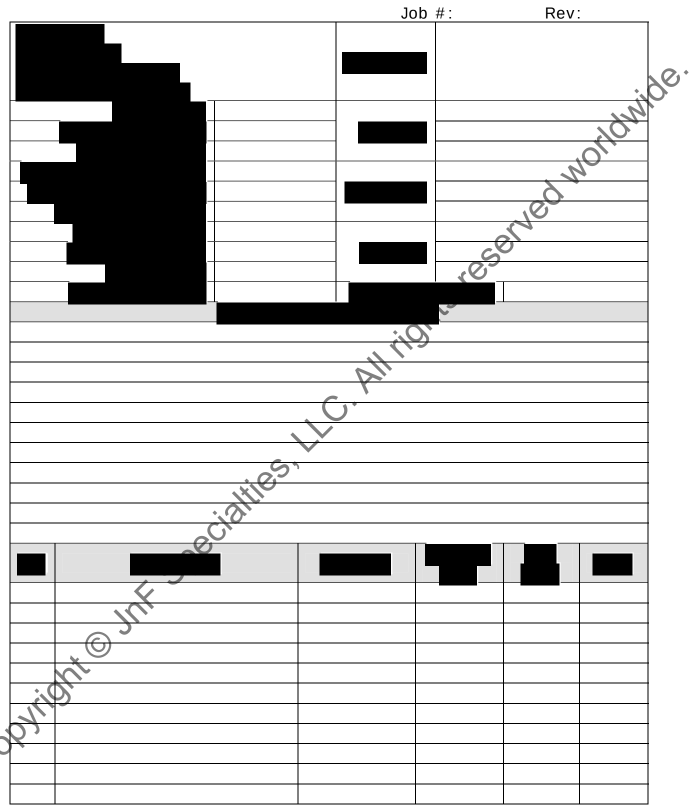
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ITEM 4: Review of resources needed to maintain and improve the effectiveness of the objects	quality management system.
ITEM 5: Review the effectiveness of current training programs and the effectiveness designated individuals. Includes	notidivide.
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ITEM 7: Review of quality objectives, data and goals. Review

Process	Quality Objective	Data Metric	Current Standing	Goal
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Internal Auditing			Mis les	
Proposal Development			- 20	
and Contract Review		Ali		
Purchasing		10.		
Receiving	•	65,		

ITEM 8: Discuss Customer feedback and complaints not already discussed as part of the NCR system review.

ITEM 9: Discuss the overall performance of the quality system, any changes to the Company that may affect the quality system or vice-versa. *Include*

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ITEM 13. NCR's FILED AT THIS MEETING:

ITEM 1	0: Note other	recommendations for management to					
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ITEM 11	. Note follow-	up activities from prior Management Review issues.					
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ITEM 12	ITEM 12. Set date for next Management Review:						
		recommendations for management to up activities from prior Management Review issues. next Management Review: ED AT THIS MEETING:					
ITEM 13	. NCR's FILI	ED AT THIS MEETING:					
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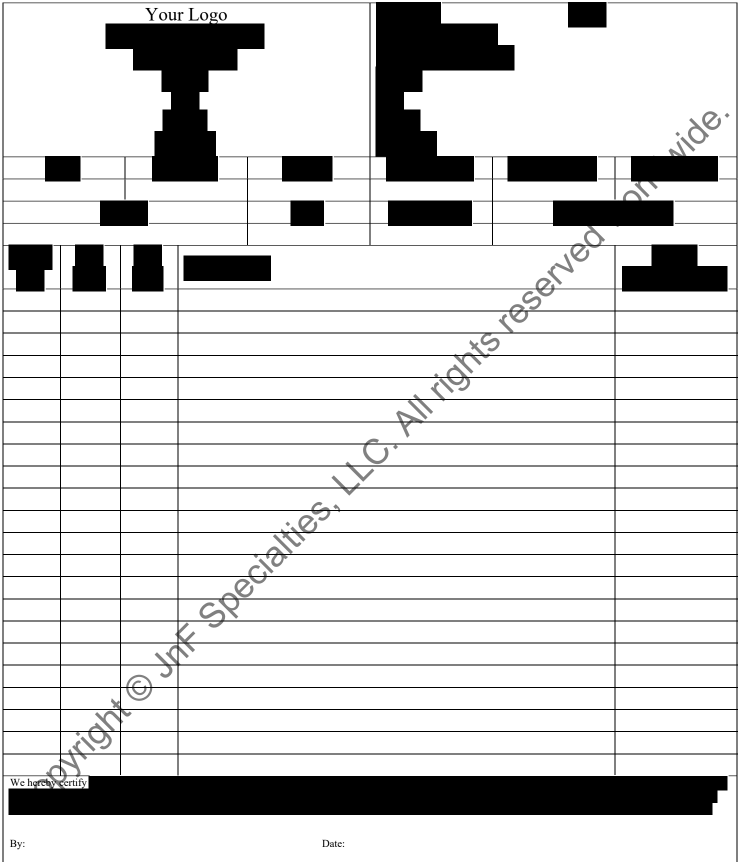
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Abbreviations:

CRR = Calculated Risk Release

CIO = Continuous Improvement Opportunity

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Dear (insert your appropriate name)
Our records show the Customer/Government property listed below is currently located at your facility. If you have knowledge of other property that should be included, please let us know by including the item(s) on your response.
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Construction Project Punch List





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Inspection Tags

Green = Good, Yellow = Withhold, Red = Bad

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Your Logo (Your Company)

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This document briefly describe	es (Your Comr	pany)'s quality management system.
Copyright © July Serio	es (Tour Comp	any) s quanty management system.
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The Company performs all project management functions according to Customer specifications. The following is a brief description of the quality management system that is used to achieve project goals.

The Company's quality management system (QMS) links numerous activities to transform inputs into outputs. The output from one process directly forms the input to the next process.

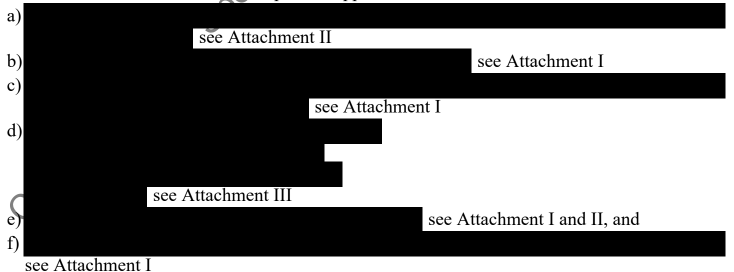
The application of a system of processes together with the identification and interaction of these processes and their management has become the Company's "process approach".

An advantage of this approach is the ongoing control that it provides over the links between and among the individual processes within the QMS as well as over their sequences and interactions.

The Company's process approach emphasizes the importance of:

- a)b)c)d)
- The Company's process approach was achieved by

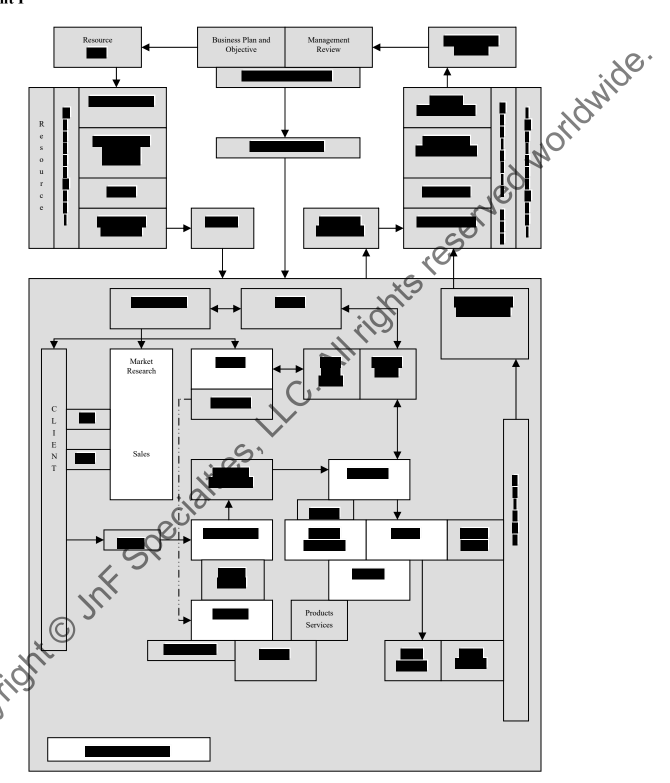
The Company's previous quality management system created an elemental structure of policies, procedures and work instructions but failed to show process interaction between inputs, outputs and their overall effectiveness. The process approach has enabled:



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Your Logo	(Your Company)	QMS Overview
The Company's quality manager	nent system (QMS) is complia	nt with
		ement that integrates Customer pany's primary tool for quality
Key functions of the QMS include	de: (value-added functions in b	pold font) Ned Ne
Another key function of the QI	MS is	
The QMS provides Users with a	access to	
The QMS is designed to		

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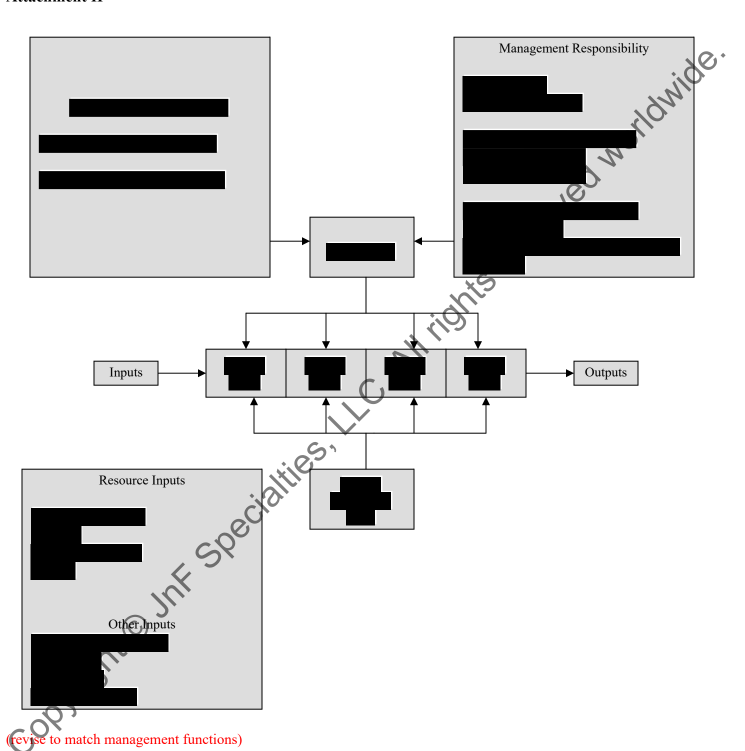
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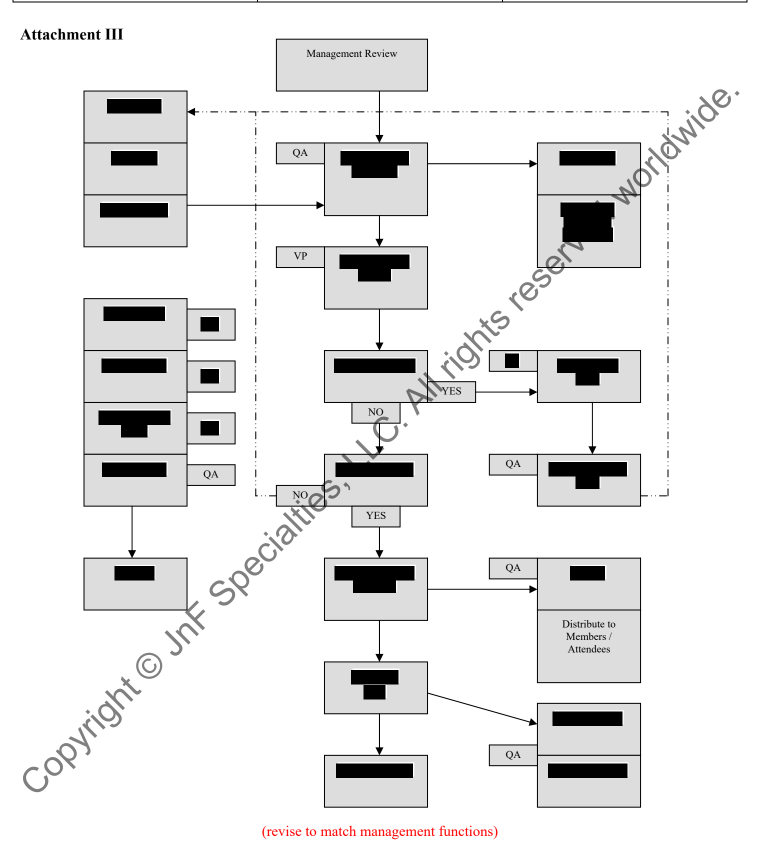
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Attachment II



Page 6 of 7



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Your Logo

PROJECT PLAN

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Document Identifier:	Name, Number, Unique ID
Date:	Latest Revision Date
Project:	Customer, Unique ID, Part Number
Document Status:	Draft, Redline, Released, Obsolete

(Your Project Description)

Under the Supervision of

(Your Custom CONTRACT NO. XXXXXXXXXXX

(Your Customer Name)

This document describes the quality control plan for xxxxxx.

Page 1 of 21

Your Logo	Your Company Name	Project Plan
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Project Plan

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3.0	SUBMITTALS	5
4.0	INSPECTION SYSTEM	6
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1.0 SCOPE

The Company has adopted a process-oriented method of management. This approach emphasizes the importance of:

a)

a)	
b)	29.
c)	.76
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The sequence and interaction of processes has been determined and are controlled by specific criteria and methods. Objectives are set for

2.0 RESPONSIBILITY AND AUTHORITY

All employees are empowered to

Project Superintendent

The Project Superintendent oversees all aspects of the job - responsibilities include:

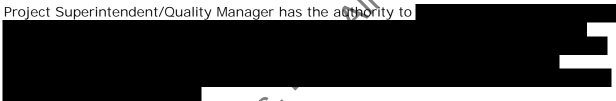
The Project Superintendent has the authority to

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Quality Manager

The Company's Superintendent/Contractor Quality Manager verifies conformance to all Plans and Specifications - responsibilities include but are not limited to:

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Alternative Contractor Quality Control Representative

In the event the Quality Manager is not present at the jobsite, the Alternative Quality Control Representative assumes

See Attachment 1 organization chart that shows lines of authority with the Quality Manager reporting to

3.0 SUBMITTALS All submittals are

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The Submittal Register is tailored to meet project schedules and is used as
General Submittal Procedure
Prior to submittal, all items shall
The Submittal Register may
Scheduling Procedure
The Company uses software program (your software name) to assure delivery of submittals
according to
4.0 INSPECTION SYSTEM
The engineering drawing, other technical documentation and identified critical items including key characteristics provides
the Quality
Manager oversees clarification of these criteria with the Project Superintendent.
Incoming materials are
Inspection consists of Preparatory, Initial and Follow-up Inspections and applicable records
for each Inspection.
Preparatory Inspections
This inspection will be conducted
This inspection will be conducted

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RECORD THE RESULTS OF THIS INSPECTION ON SEPARATE SHEETS AND ATTACH THEM TO THE DAILY REPORT.

Initial Inspections

This inspection will be held

Initial Inspections may include:



RECORD THE RESULTS OF THIS INSPECTION ON SEPARATE SHEETS AND ATTACH THEM TO THE DAILY REPORT.

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personnel may arrange with the Quality Mar	nager to be present for this inspection.
Follow-up Inspections may include:	
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Documentation and Control	, es
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Completion Inspection	
Completion Inspection Punch-Out Inspection:	
The Project Superintendent and Quality Man and develop a punch list of items that do no	ager shall conduct an inspection of the work t conform to the approved drawings and
specifications. The Responsible Authorities	//III
Pre-Final Inspection	
The Customer will perform this inspection to to be operated. A Customer Pre-Final Punch	verify the construction is complete and ready
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5.0 TESTING	2		, n
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6.0 DOCUMI	ENTS AND REC	ORDS	
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7.0 CONTROL OF NONCONFORMANCES

Construction design and construction deficiencies that are found to be nonconforming against specified requirements are

REWORK PROCEDURES

The Company has a long standing successful Noncompliance Management Program to ensure

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8.0 DOCUMENTATION		
Procedure		
All reportable records shall include	$i^{(0)}$	
All submittals of records will		
Test Reports will	Y	
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9.0 WORKMANSHIP

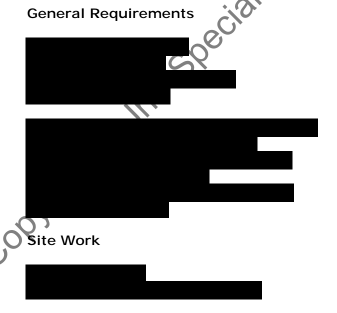
The Company plans and carries out construction activities that include workmanship requirements for:



10.0LIST OF DEFINABLE FEATURES OF WORK

(Tailor this section to address key elements of the project. A definable feature of work is a task that is separate and distinct from other tasks, has separate control requirements and may be identified by different trades or disciplines or it may be work by the same trade in a different environment. This list should be agreed upon during the coordination meeting.)

For instance – breakdown each work element from your contract Plans and Specifications:



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# Information Request

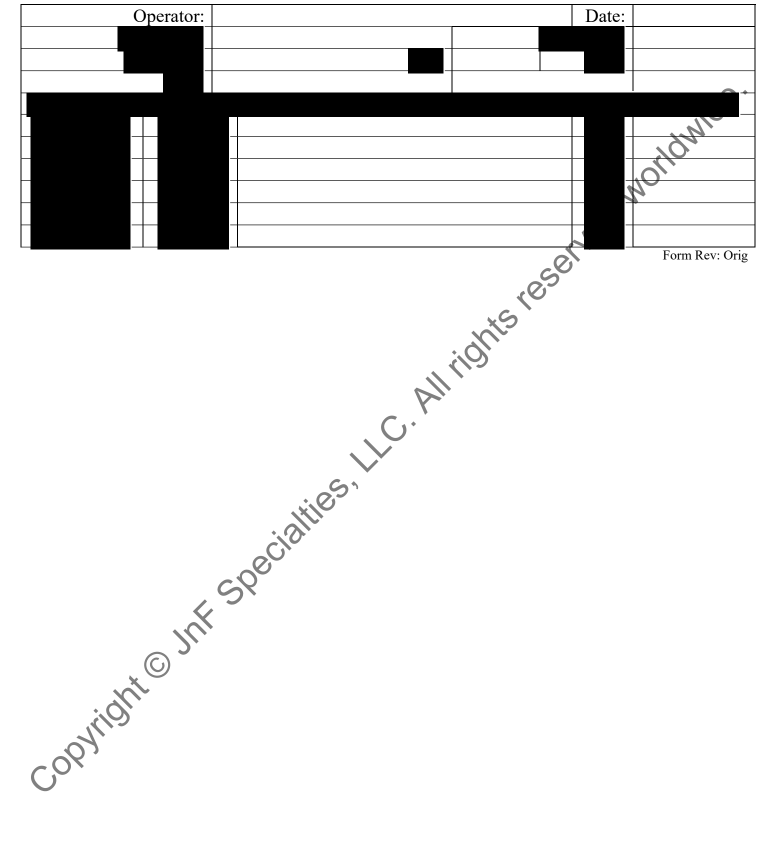
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### **ROUTING TICKET**

ACCOUNT#:



**Shelf Life Expiration Log** 

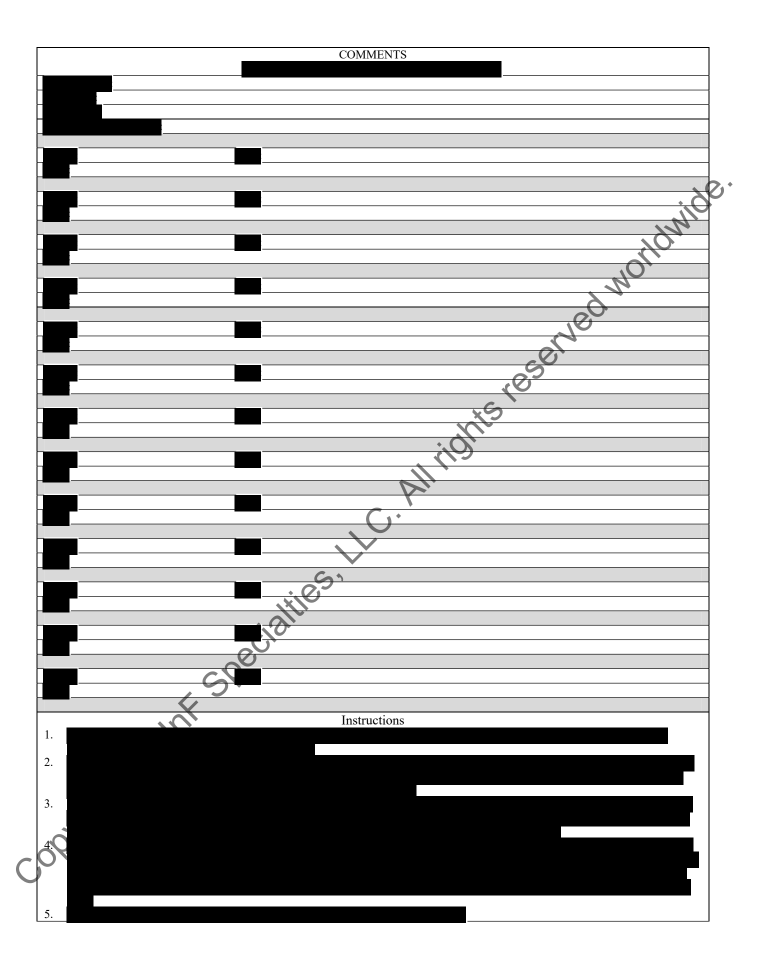
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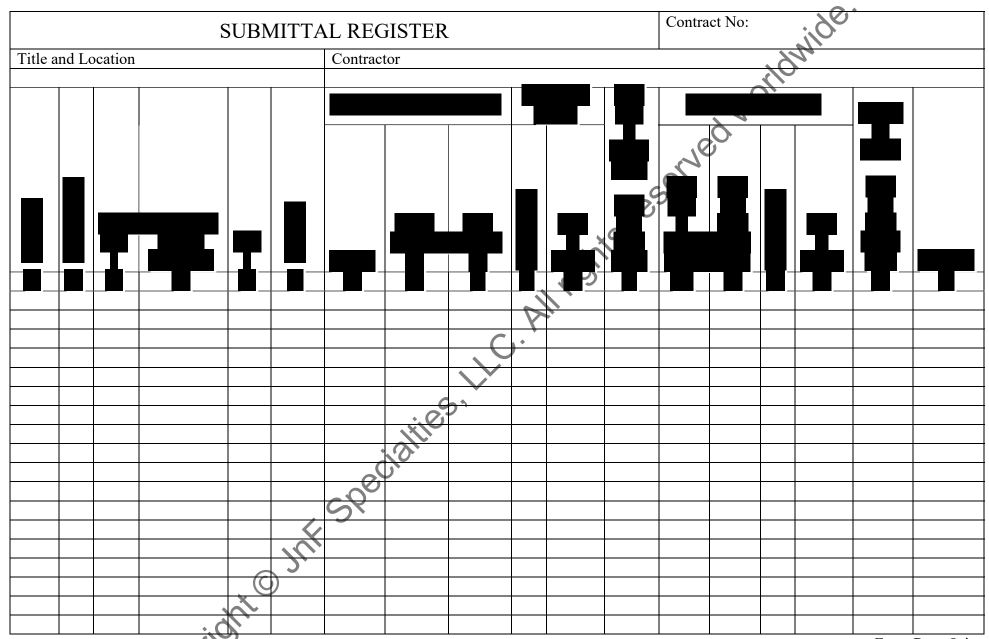
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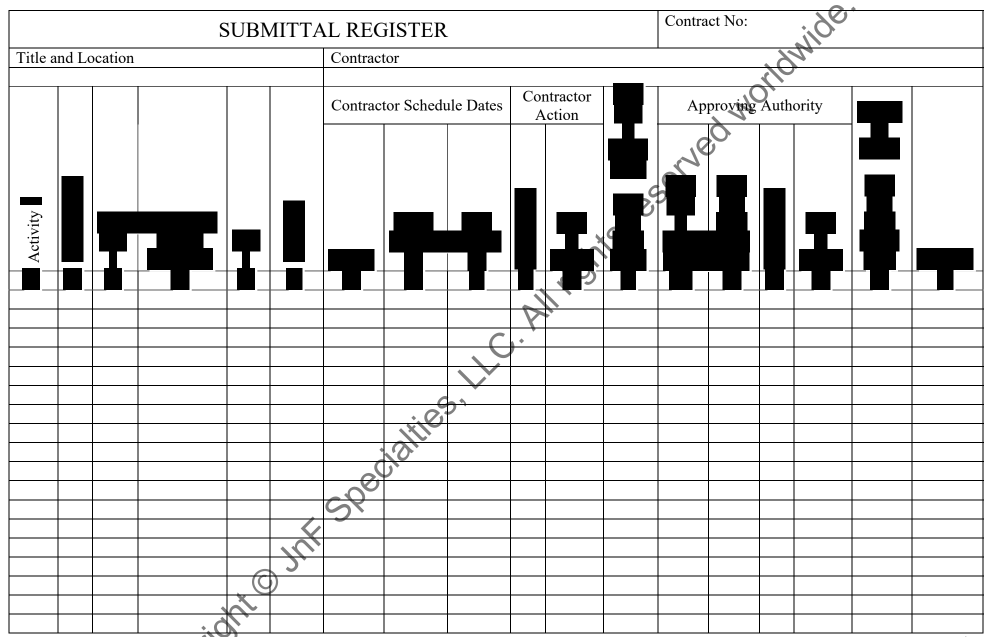


(Your Company Name) Address, City State, Zip Phone, Fax

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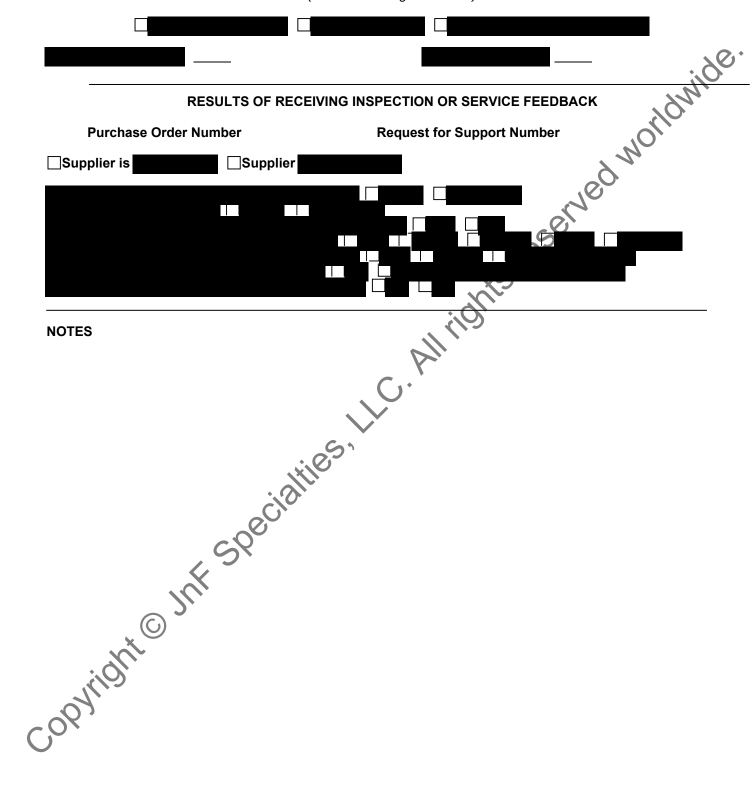
### **Supplier Evaluation**

Supplier:	Commodity:
	If Part I criteria is met, Supplier is approved without further evaluation.
Part I	
	If Part I criteria is NOT met, Supplier must be evaluated under Part II, III and IV.
	the boxes below for each criterion evaluated. Attach evidence where indicated.  Criteria must be checked in Part II for the Supplier to be qualified.
	The supplier to be qualified.
Part III	
Part IV	ali the beautiful of the state
Evaluator: Chec	ck the boxes below to identify if direct or third party review is required for Detailing Subcontractors:
	ck the boxes below to confirm the Detailing Subcontractor identifies the following information on
drawings:	
Evaluator: Whe	n the Company awards a contract in advance of Subcontractor Evaluation, check the boxes below to igation for the following:

### **Supplier Evaluation**

### **RESULTS OF EVALUATION**

(Ref. Purchasing Procedure)



### SUPPLIER PERFORMANCE RATING REPORT

Performance Reporting Dates:

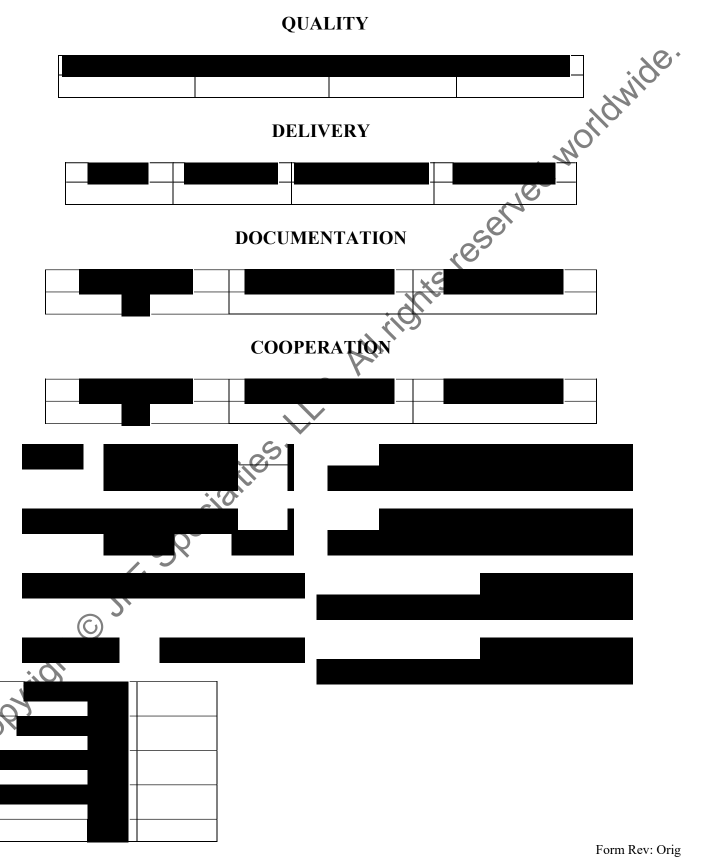
Job #:

Supplier:	
	OVERALL PERFORMANCE RATING 100  Excellent Good Improvement Expected Improvement Required  Weight %
Points (100 Max)	Weight %
Quality	100
Delivery	100
Documentation	100
Cooperation	100
Quality: The number of items accept received times 100.  Delivery: The grace period is	oted divided by the number of items that should have been
completeness of	assessment of the Suppliers willingness to cooperate,
Purchasing Agent	Date Form Rev: Orig

### SUPPLIER RATING WORKSHEET

Supplier: P/N:

### **QUALITY**



Supplier Overall Performance Rating Overall Performance Rating Supplier: Month: Supplier Monthly Rating Report

Rating Monthly and Average Percentage Rating
J F M A M J J A S O N Supplier Copyright Specialties. Date:

Your Logo Your Company Name

# SUPPLIER QUALITY REQUIREMEN Origination Date: XXXX Document

Document Identifier:	Supplier Quality Requirements
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 flowdown requirements for Suppliers.

Your logo	Vour Company Nama	Supplier Quality Requirements
CAGE: xxxxx	Your Company Name	Rev: Orig

### **REVISION LOG**

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### DOCUMENT CHANGE RECORD

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### Page 2 of 7

Your logo	y c y	Supplier Quality Requirements
CAGE: xxxxx	Your Company Name	Rev: Orig
☐PURPOSE and SCOPE		
•	s for supplier Quality Systems necessary to nirements of the Contract. Procedures used pproval upon request.	
APPLICABILITY		July
These requirements shall apply to all su thereto.	applies and services when referenced on the	e Purchase Order and amendments
commitment for an Inspection System s	Seller's Inspection System Level I, as a reconstall be defined by all paragraphs of this spont then the Seller's contractual commitments specification which are checked-off.	pecification. When Buyer's Purchase t for an Inspection System shall be
DEFINITIONS and ABBREVI	ATIONS	reserv
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D.	M, GENERAL	
SELLER'S QUALITY SYSTE	M, GENERAL	
	Quality System planned and developed in co	onjunction with his other functions to
The System shall provide	. 1975	
Records shall be kept available for		
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□NEGOTIATIONS		
It is not the intent of this specification t	o restrict the Seller in his mode of operatio	n; therefore,
301		
PROPRIETARY INFORMATION	ON	
The Seller must identify in writing the	intended use in performance of the Purchas	
or process with respect to which access	by Buyer or Buyer Customer representative	ves for purpose of Quality Assurance by
inspection, test or process surveillance	is proposed to be restricted. The written id	enumention snail state

i our logo	Vaya Cammany Nama	Supplier Quality Requirements
CAGE: xxxxx	Your Company Name	Rev: Ori
The absence of such written identification	on is a representation by Seller	
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☐PROCESS CONTROL		advide
The Seller shall provide for		
Work instructions for all work affecting	quality shall	
Work instructions for an work affecting	quanty shan	
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The Seller shall develop an Inspection/T The Plan shall	Test Plan specific in nature and related d	lirectly to the hardware produced.
Buyer contracts and resultant facility pla	anning by Seller shall	
All Purchase Orders that apply to Buyer	contracts generated by Seller shall	
When approval or certification of specia	I processes operating personnel specia	al equipment, or procedures is required by
the contract, drawing, or specification, the		requipment, or procedures is required by
_		
Seller MRB is not authorized. Seller sha	all notify Buyer within 48 hours of dete	cted failure. Buyer and/or Buyer
Customer representatives shall		
Formal Failure Analysis and Corrective	Action shall	
A Seller Failure Review Board is		
The Seller shall not change		
60.		
When the Dynalis - Out - Out - D	m accountance of a 1-t Aut 1. d. f.	out fakuigated to the arracif of Decree
When the Purchase Order requires Buye configuration shall	acceptance of a 1st Article, the first pa	art raoricated to the specified Buyer

Supplier Quality Requirements

Your logo

Your logo	Vous Commony Nome	Supplier Quality Requirements
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This check sheet is		
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The 1st Article item and the inspection r	record shall	
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SUBCONTRACTOR CONTRO	DL X	
The Seller shall be responsible for		
Buyer inspection is		
Buyer inspection is	<u></u>	
☐DRAWING and CHANGE CO	NTROL ()	
The Seller shall have a procedure and de		
The procedure shall also provide for		
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RECEIVING INSPECTION		
The Seller shall inspect incoming materi		
Acceptance requirements shall include		
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<b>□STOCK CONTROL</b>		

The Seller shall provide for protection and control of supplies and materials stored for use in deliverable Buyer products.

Your logo	Vana Campany Nama	Supplier Quality Requirements
CAGE: xxxxx	Your Company Name	Rev: Orig
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Control shall cover such items as		
Durandaman familia handling of naman fa	amain a makanial aball	
Procedures for the handling of nonconfo	orming material shall	
Buyer furnished material shall		
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SAMPLING INSPECTION		8
Acceptance sampling procedures, if other	er than ANSI Z 1.4, must have Buyer ap	proval prior to use; sampling to permit
defects is		
☐TOOL, GAGE, and TEST EQU	IIDMENT	10,50
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The Seller shall be responsible for provi equipment to assure supplies conform to		stability of tools, gages, and test
A written procedure,		
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MATERIAL CONTROL	63	
Nonconforming material shall		
Seller may not		
Sener may not	O	
The Seller shall maintain traceability of	raw material used in the production of d	leliverable products. A correlation shall
be made between		
The Seller shall maintain controls to ass	ure accomplishment of	
The Selici shall maintain controls to ass	are accompnishment or	
the mayisions of ASTM D 2051 masses.	ration make aims making and marking	Unless otherwise specified,
the provisions of ASTM B 3951 preserv		
Direct shipment of your supplies to Buy advance of your expected shipping date.		rs Purchasing Manager ten (10) days in
201		
Co.		
When product is returned by Buyer to the	ne Seller because of failure to comply wi	ith Purchase Order requirements, the
Seller shall		

Your logo	Vous Commony Name	Supplier Quality Requirements
CAGE: xxxxx	Your Company Name	Rev: Orig

TECHNICAL REQUIREMENTS Unless otherwise specified, Buyer is responsible for  Which the Special times in the specia	wide
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(Date)

Quality Manager«AddressBlock»

Re: Supplier Performance Rating Report Performance Reporting Dates: P.O. #

Dear QC Manager:

served worldwide. We have developed a Supplier Report Card that indicates your Quality Performance. Enclosed is a copy of your Quality Performance, which includes

If you have any questions, please call or email us.

Sincerely,

Your Name

ne ecialiles Your Company Name

Your City, State, Zip

يan يdress City, State none: Your# Fax: Your# Email: Your email

Ref:	Your Company Name
Page 1 / of /	SURVEY REPORT
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# Your Production Area Training Certificato

# awarded to Your Employee Name

**S**Your Specification **Your Details** 

**Your Date** 

Training Supervisor

Quality Manager

### TRAINING LOG

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### QMS Procedure Training Matrix for (Your Company)

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C. eQMS Ch. eQMS	^	^	X	^	^	X	X	^	^	X	^	X	ΟX	X
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X = Applicable QMS Procedure record of orientation training for each Employee. The Company must produce a record of orientation for all employees affected by individual QMS procedures to achieve QMS pedigree.

Note -

### ORIENTATION/TRAINING REOUEST

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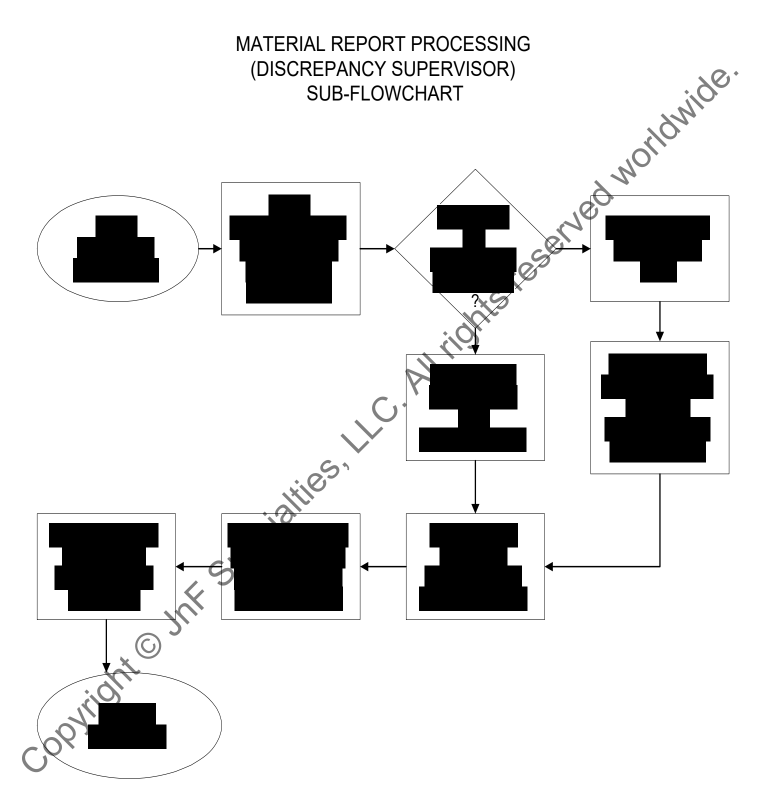
### 1.0 Scope

Document procedures using block diagrams or flowcharts that describe discrete operations in a process. Prepare work instructions to explain details in procedures but only when

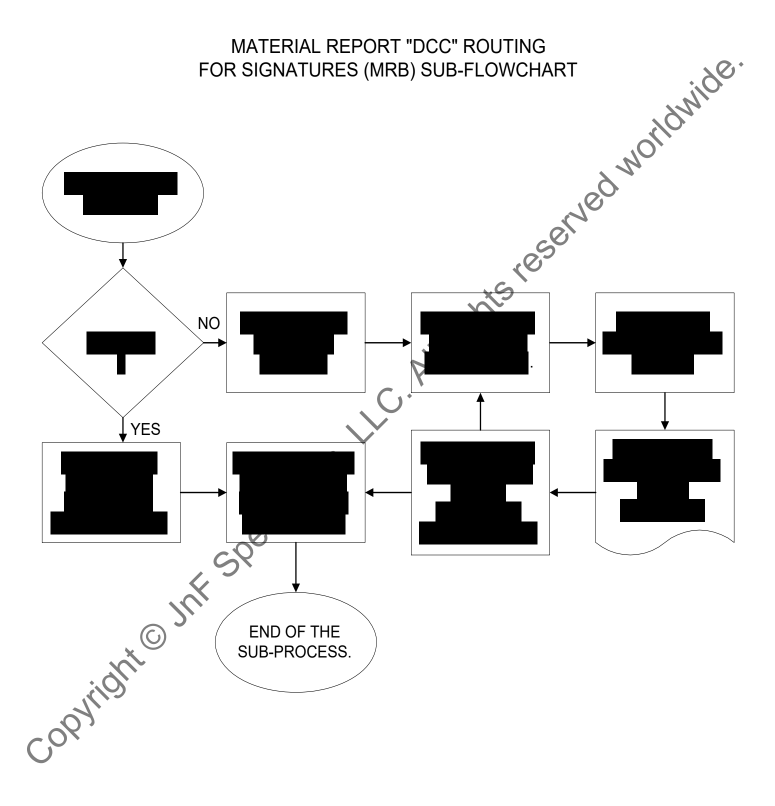
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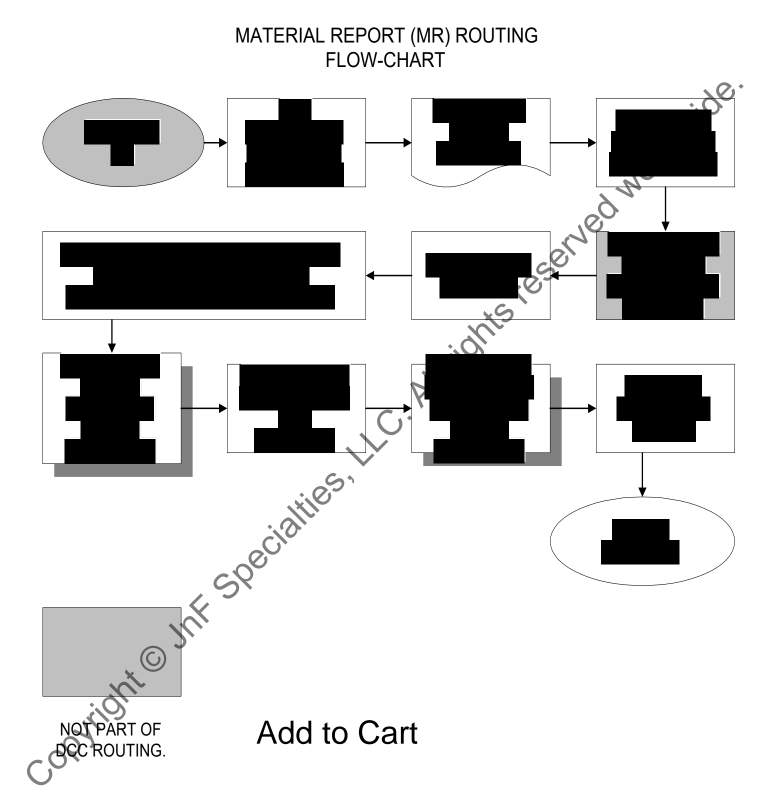
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